Sugar- and lipid-derived aldehydes are reactive carbonyl species (RCS) frequently used as surrogate markers of oxidative stress in obesity. A pathogenic role for RCS in metabolic diseases of obesity remains controversial, however, partly because of their highly diffuse and broad reactivity and the lack of specific RCS-scavenging therapies. Naturally occurring histidine dipeptides (e.g., anserine and carnosine) show RCS reactivity, but their therapeutic potential in humans is limited by serum carnosinases. Here, we present the rational design, characterization, and pharmacological evaluation of carnosinol, i.e., (2S)-2-(3-amino propanoylamino)-3-(1H-imidazol-5-yl)propanol, a derivative of carnosine with high oral bioavailability that is resistant to carnosinases. Carnosinol displayed a suitable ADMET (absorption, distribution, metabolism, excretion, and toxicity) profile and was determined to have the greatest potency and selectivity toward $\alpha,\beta$-unsaturated aldehydes (e.g., 4-hydroxynonenal, HNE, ACR) among all others reported thus far. In rodent models of diet-induced obesity and metabolic syndrome, carnosinol dose-dependently attenuated HNE adduct formation in liver and skeletal muscle, while simultaneously mitigating inflammation, dyslipidemia, insulin resistance, and steatohepatitis. These improvements in metabolic parameters with carnosinol were not due to changes in energy expenditure, physical activity, adiposity, or body weight. Collectively, our findings illustrate a pathogenic role for RCS in obesity-related metabolic disorders and provide validation for a promising new class of carbonyl-scavenging therapeutic compounds rationally derived from carnosine.
A carnosine analog mitigates metabolic disorders of obesity by reducing carbonyl stress

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Sugar- and lipid-derived aldehydes are reactive carbonyl species (RCS) frequently used as surrogate markers of oxidative stress in obesity. A pathogenic role for RCS in metabolic diseases of obesity remains controversial, however, partly because of their highly diffuse and broad reactivity and the lack of specific RCS-scavenging therapies. Naturally occurring histidine dipeptides (e.g., anserine and carnosine) show RCS reactivity, but their therapeutic potential in humans is limited by serum carnosinases. Here, we present the rational design, characterization, and pharmacological evaluation of carnosinol, i.e., (2S)-2-(3-amino propanoylamino)-3-(1H-imidazol-5-yl)propanol, a derivative of carnosine with high oral bioavailability that is resistant to carnosinases. Carnosinol displayed a suitable ADMET (absorption, distribution, metabolism, excretion, and toxicity) profile and was determined to have the greatest potency and selectivity toward α,β-unsaturated aldehydes (e.g., 4-hydroxynonenal, HNE, ACR) among all others reported thus far. In rodent models of diet-induced obesity and metabolic syndrome, carnosinol dose-dependently attenuated HNE adduct formation in liver and skeletal muscle, while simultaneously mitigating inflammation, dyslipidemia, insulin resistance, and steatohepatitis. These improvements in metabolic parameters with carnosinol were not due to changes in energy expenditure, physical activity, adiposity, or body weight. Collectively, our findings illustrate a pathogenic role for RCS in obesity-related metabolic disorders and provide validation for a promising new class of carbonyl-scavenging therapeutic compounds rationally derived from carnosine.

Introduction

Overnutrition from fatty acids and complex carbohydrates is known to cause oxidative stress from multiple enzymatic and non-enzymatic sources due to the high caloric content and the prevalence of these macronutrients in the Western diet. A pathological role for oxidative stress in obesity has been clearly established through extensive clinical and experimental studies. Reactive sugar- and lipid-derived aldehydes (reactive carbonyl species [RCS]) are spontaneously formed during the preparation of high-fat/high-sugar-containing foods under high heat (1) and are formed in vivo as a byproduct of oxidative stress. These RCS alter proteins via the covalent modifications of cysteine, arginine, lysine, and histidine and form adducts with phospholipids and DNA. Glucose-derived oxoaldehydes such as methylglyoxal (MGO) accumulate in oxidative tissues and react with protein functional groups, forming advanced glycation end-products (AGEs) (2, 3). AGEs are a major cause of chronic inflammation, cardiovascular disease, diabetes, and even some cancers, all of which are associated with obesity (4–8). In addition to reducing sugars and sugar-derived breakdown products, lipid-derived aldehydes are another significant source of carbonyl stress in vivo. Individuals who consume diets rich in vegetable and corn oil have very high endogenous levels of n-6 polyunsaturated fatty acids (PUFAs). The α,β-unsaturated carboxyls derived from n-6 PUFA oxidation have particularly diverse biological effects. Of these, 4-hydroxynonenal (HNE) and acrolein (ACR) have been the most widely studied (9). These aldehydes rapidly form adducts with proteins and phospholipids (10), and chronic exposure to high levels of these aldehydes is toxic (11). The accumulation of HNE has been extensively documented in blood and tissue samples from obese/diabetic patients, typically as a marker of oxidative stress (12–16). However, emerging studies suggest that these reactive aldehydes are more than simply by-products of oxidative stress. Rather, the carbonyl-modifying activity imposed by these reactive species may have a distinct pathogenic role in obesity-related disorders such as insulin resistance, chronic inflammation and fibrosis, dyslipidemia, and liver disease (15, 17–23). Thus, novel compounds that mitigate the production or enhance the removal of RCS remain compelling therapies for cardiovascular and metabolic diseases associated with obesity.
l-carnosine is a naturally occurring dipeptide (β-alanyl-histidine), which, along with its analogs, is a potent endogenous scavenger of RCS and highly concentrated (mM) in muscle and nervous tissues (24). The utility of l-carnosine as a pharmacological agent has been demonstrated in rodent models of metabolic syndrome and cardiovascular disease (25–27) and by its widespread use as eyedrop therapy in patients with ocular diseases (28). Major obstacles exist with respect to the clinical applicability of l-carnosine as an oral drug therapy, however. The largest of these is that high serum and tissue carnosinase activity in humans abrogates the bioavailability of circulating carnosine by rapid hydrolysis of the peptide bond (24).

Here, we present the rational design, characterization, and pharmacological evaluation of carnosinol, i.e., (2S)-2-(3-aminopropanoylamino)-3-(1H-imidazol-5-yl)propanol, a reduced derivative of l-carnosine that is impervious to metabolism by carnosinase. Carnosinol displayed selectivity for reaction with RCS in vitro and in vivo, oral bioavailability and long duration in vivo, and negligible toxicity in human cell cultures and animal models. In rodent models of diet-induced obesity, carnosinol dose-dependently reduced systemic carbonyl stress, normalized glycemic control and many inflammatory parameters, and mitigated steatohepatitis. Collectively, these findings illustrate a distinct pathological role of RCS in metabolic diseases of obesity and validate the use of a novel RCS-scavenging l-carnosine derivative to treat these diseases.

Results

Rational design of carnosinol

In addition to the common properties that characterize a drug-like molecule (i.e., chemical and metabolic stability, bioavailability, and safety), our a priori rationale was that an RCS-sequestering therapeutic compound should: (a) be stable in plasma; (b) effectively scavenge (i.e., trap) circulating RCS; and (c) be highly reactive and selective toward damaging RCS (29). l-carnosine fulfills some of the above-mentioned requirements (30). However, it lacks an important basic requirement to be a suitable RCS-sequestering agent, because it is unstable in the circulation as a result of the hydrolytic action of carnosinases.

Hence, the rational design of improved l-carnosine derivatives should be focused on molecules that, besides maintaining or even enhancing quenching activity and selectivity, are endowed with plasma stability and oral bioavailability. To this end, the ideal derivative should: (a) maintain or better optimize its quenching activity, at least toward HNE, while preserving its selectivity; (b) maintain the active transport by human H+/peptide cotransporter-1 (hPepT1); and (c) eliminate recognition by human serum carnosinase (30, 31). As depicted in Figure 1A, this can be pursued by modifying the carboxyl group, which (a) is not involved in the quenching mechanism, even though its complete deletion has a detrimental effect, as seen in carcinine; (b) has a crucial role in carnosinase-1 (CN1) binding (Figure 1B); and (c) has a marginal role in hPepT1 transport (Figure 1C). Reduction of the carboxyl group has only a modest impact on the PepT1 interaction (Figure 1C), since both the carboxyl and hydroxyl functions elicit comparable and (weak) H-bonds with surrounding backbone atoms, as also confirmed by very similar interaction energies (–19.056 vs. –17.934 kcal/mol). In contrast, the carnosine carboxyl group is engaged in a pivotal ion pair with Arg350 in CN1 (Figure 1B), and its reduction to the hydroxyl group has a dramatic impact on the stability of the CN1-carnosinol complex, as confirmed by the reported drop in the interaction energy (–22.770 vs. –14.862 kcal/mol).

Given these modeling results, the carboxyl reduction to yield carnosinol should meet all the above requirements and should
even enhance the quenching activity, since it optimizes some physicochemical parameters that influence the quenching mechanism (e.g., flexibility and lipophilicity). Besides a greater flexibility, physicochemical analysis reveals that carnosinol is also more lipophilic than carnosine, as seen in experimental log P and log D values, a property that would benefit the molecular engagement by lipophilic RCS (Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/JCI94307DS1). Furthermore, carboxyl reduction did not significantly affect the ionization constant or the chelating activity by lipophilic RCS (Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/JCI94307DS1). Quantum mechanical simulations showed that carnosinol also possesses favorable stereoelectronic parameters (Supplemental Table 1).

### Carnosinol is a selective and potent RCS-sequestering agent

We first performed in vitro testing of the sequestering activity of carnosinol toward the most widely studied RCS involved in oxidative-based diseases and belonging to the chemical classes of α,β-unsaturated aldehydes, i.e., 4-hydroxynonenal (HNE), ACR dialdehydes, i.e., malondialdehyde (MDA) and glyoxal (GO), and ketoaldehydes, i.e., MGO.

#### Sequestering of HNE

The sequestering activity of carnosinol toward HNE and ACR was first evaluated and compared with L-carnosine and for other well-known RCS-sequestering agents such as aminoguanidine and pyridoxal. The increased potency of carnosinol compared with that of L-carnosine can be explained by the favorable stereoelectronic parameters as predicted by quantum mechanical simulations (Supplemental Table 1) and also by the ability of the hydroxyl group to form a hemiacetal intermediate with HNE, which, in addition to the amino group of β-alanine, further catalyzes the formation of the Michael adduct.

The mechanism of carnosinol reaction toward HNE was fully elucidated on the basis of the reaction products identified and characterized by mass spectrometry (MS) experiments (Supplemental Figure 1). The proposed reaction mechanism (Figure 2A) is similar to that already clarified for L-carnosine and based on the formation of an imine derivative (CI) that catalyzes the Michael adduct (CII) (Figure 2B).

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#### Sequestering of MDA

The sequestering activity of carnosinol toward MDA was also first evaluated and compared with L-carnosine (Supplemental Figure 2). The sequestering activity toward MDA was fully elucidated on the basis of the reaction products identified and characterized by mass spectrometry (MS) experiments (Supplemental Table 1). The sequestering activity toward MDA was found superimposable with respect to carnosine, and the N-prope-

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**Table 1. Comparison of carnosinol reactivity with HNE and the biogenic aldehyde pyridoxal, reported as consumption percentages (Q% ± SD)**

<table>
<thead>
<tr>
<th>Compound</th>
<th>HPLC (Q% ± SD) HNE</th>
<th>MS (Q% ± SD) HNE</th>
<th>Pyridoxal HPLC (Q% ± SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>L-carnosine</td>
<td>37.3 ± 2.3</td>
<td>25.4 ± 3.7</td>
<td>0.1 ± 0.9</td>
</tr>
<tr>
<td>Carnosinol</td>
<td>61.5 ± 1.9</td>
<td>38.1 ± 4.0</td>
<td>0.1 ± 0.1</td>
</tr>
<tr>
<td>Pyridoxamine</td>
<td>2.1 ± 0.5</td>
<td>0.0 ± 3.4</td>
<td>2.1 ± 0.1</td>
</tr>
<tr>
<td>Hydralazine</td>
<td>87.2 ± 3.2</td>
<td>15.2 ± 3.3</td>
<td>100.0</td>
</tr>
<tr>
<td>Aminoguanidine</td>
<td>2.3 ± 0.7</td>
<td>3.9 ± 3.2</td>
<td>12.8 ± 0.1</td>
</tr>
</tbody>
</table>

HPLC Q% values express the percentages of the reacted target aldehyde in the presence of the tested quenchers after a 24-hour incubation, with a molar ratio of quencher/aldehyde equal to 1:1 (HNE) or 10:1 (pyridoxal). MS Q% values express the percentage of inhibition of carbonylated ubiquitin induced by HNE after a 24-hour incubation, with a molar ratio of quencher/aldehyde equal to 1:1 (HNE).

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amine. We found carnosinol to be less effective only by comparison with hydralazine, a commonly prescribed antihypertensive medication with non-selective RCS reactivity, as shown by its effects with pyridoxal. In contrast, we found that carnosinol did not react with pyridoxal, and hence it can be considered a selective RCS-sequestering agent, like the parent compound L-carnosine.
reduces atherogenesis (27, 34, 35), dyslipidemia, and renal dysfunction (25) in rodent models of cardiometabolic disease. To determine whether carnosinol retained or surpassed the efficacy of L-carnosine, we tested the effect of low-dose (10 mg/kg/day) and high-dose (45 mg/kg/day in the drinking water) carnosinol on cardiometabolic and inflammatory parameters using a short-term intervention (3 weeks) in rats fed a high-fructose (HF) diet (60% in the food pellet) (Figure 4A). An additional group of rats was treated for an equivalent period with a clinically relevant dose of rosiglitazone, the PPAR agonist widely prescribed for the treatment of metabolic syndrome and type 2 diabetes and which we recently determined to be a very effective therapy in this model (36). HF feeding led to significant gains in body weight (Figure 4B and Supplemental Table 3) compared with control-fed rats, and drug treatment did not cause significant differences in body weight gains. Total food and water intake was similar between the groups. Systolic blood pressure was substantially higher in fructose-fed rats, and this was dose-dependently reduced by carnosinol to levels comparable to those seen with rosiglitazone (Supplemental Table 3). This improvement in blood pressure with carnosinol mirrors the effect of L-carnosine supplementation on these parameters in similar models (25, 26).

Next, we sought to examine the effect of carnosinol on markers of systemic oxidative and inflammatory stress. Notably, carnosinol dose-dependently reduced serum AGEs (Figure 4C), urinary 8-isoprostanes (Table 3), and proinflammatory cytokines (Figure 4, D–F) in fructose-fed rats. Importantly, the beneficial effects of carnosinol treatment on oxidative and inflammatory stress were paralleled by improvements in metabolic parameters. Carnosinol treatment led to dose-dependent reductions in serum tri-

nal derivative of carnosinol was identified as the reaction product (Supplemental Figure 3).

Taken together, the data indicate that carnosinol is selective and very reactive toward biogenic sugar- and lipid-derived aldehydes and the most potent RCS-sequestering agent among those commercially available.

ADMET studies
Carnosinol was evaluated in in vitro ADMET studies as well as in exploratory pharmacokinetic (PK) and toxicological studies (additional information is available in the supplemental material). Carnosinol was found to be stable in human serum for at least 60 minutes, as well as in rat and human liver preparations, thus indicating that it is not hydrolyzed by carnosinases. In vitro experiments indicated a good safety profile of the compound (no inhibition of the most important isoenzymes of cytochrome P450, no interaction with hERG K+ channel, no cell toxicity up to 100 μM concentration in a human hepatoma cell line). We then confirmed the safety profile of carnosinol in rats in both single- and repeat-dose oral toxicity studies. PK studies in rats showed a moderately good oral bioavailability that gave a significant exposure in plasma after oral administration at a dose of 45 mg/kg.

The data reported above demonstrate that carnosinol fulfills the optimization requirements we took into account in the rational design.

Dose-dependent effects of carnosinol on systemic inflammation and metabolic derangements in fructose-fed rats
Prior studies by our group and others have demonstrated that oral administration of L-carnosine and its derivatives effectively reduces atherogenesis (27, 34, 35), dyslipidemia, and renal dysfunction (25) in rodent models of cardiometabolic disease. To determine whether carnosinol retained or surpassed the efficacy of L-carnosine, we tested the effect of low-dose (10 mg/kg/day) and high-dose (45 mg/kg/day in the drinking water) carnosinol on cardiometabolic and inflammatory parameters using a short-term intervention (3 weeks) in rats fed a high-fructose (HF) diet (60% in the food pellet) (Figure 4A). An additional group of rats was treated for an equivalent period with a clinically relevant dose of rosiglitazone, the PPAR agonist widely prescribed for the treatment of metabolic syndrome and type 2 diabetes and which we recently determined to be a very effective therapy in this model (36). HF feeding led to significant gains in body weight (Figure 4B and Supplemental Table 3) compared with control-fed rats, and drug treatment did not cause significant differences in body weight gains. Total food and water intake was similar between the groups. Systolic blood pressure was substantially higher in fructose-fed rats, and this was dose-dependently reduced by carnosinol to levels comparable to those seen with rosiglitazone (Supplemental Table 3). This improvement in blood pressure with carnosinol mirrors the effect of L-carnosine supplementation on these parameters in similar models (25, 26).

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fat/high-sucrose (HFHS) diet (15). These cardiometa-
abolic disorders in GPx4+/– mice were accompanied by
extensive carbonyl stress in liver and heart, and cardiac
mitochondria in obese GPx4+/– mice had decreased fatty
acid–supported respiration and increased ROS produc-
tion compared with obese WT mice. These results are
consistent with mitochondrial localization of GPx4 and
its known role in protecting mitochondria from oxidative
stress (40). Importantly, diabetic patients have higher
levels of HNE adducts and lower GPx4 enzyme levels
in their myocardial tissue compared with nondiabetic
patients (15), a finding that corroborates previous obser-
vations concerning HNE adducts in diabetic patients
(12–14, 16). Thus, GPx4+/– mice are ideal for the pharma-
cological evaluation of carnosinol, because much of the
underlying stress-induced pathology of these mice is due
to oxylipid-derived aldehydes.

To determine whether carnosinol is effective at
mitigating obesity-related metabolic disorders, we
administered the compound at the high dose (45 mg/kg/day in
the drinking water) in a cohort of HFHS diet–induced obese WT
mice. Given our previous findings in GPx4+/– mice on a HFHS
diet, a cohort of GPx4+/– mice was used in parallel with WT mice
to allow for assessment of carnosinol in a clinically relevant,
translational model of severe carbonyl stress. Drug was admin-
glycerides and cholesterol (Figure 4, G and H), improved glycemic
control (Figure 4, I and J), and mitigated liver toxicity (Figure 4K)
and steatosis (Table 3) induced by fructose feeding. All improve-
ments in systemic metabolic and inflammatory parameters with
carnosinol treatment were paralleled by significant reductions
in plasma (Figure 4, L and M), kidney, and liver of HNE adducts
(Supplemental Figure 4). Liver and renal fibrosis were not
significantly affected by fructose feeding or drug treatment
(data not shown), although the urinary creatinine clearance
rate decreased with fructose feeding (Table 3), and this effect
was blunted with carnosinol and rosiglitazone treatment.

Carnosinol improves glycemic control and muscle insulin
sensitivity in mouse models of severe carbonyl stress and
diet-induced obesity
Glutathione peroxidase 4 (GPx4), also named phospholip-
id hydroperoxide glutathione peroxidase, is the only known
selenoenzyme that exclusively neutralizes lipid peroxides in
membranes and lipoproteins (37). Homozygous null GPx4
mice die around gestational day 7, underscoring the critical
role for this enzyme in development. GPx4-haploinsufficient
(GPx4+/–) mice have approximately 40% of the WT GPx4
enzyme levels in oxidative tissues and are highly susceptible
to environmental stressors due to enhanced lipid peroxidation
and protein carbonylation (38, 39). Recently, we observed that
GPx4+/– mice displayed exacerbated metabolic derangements
and cardiomyopathy when fed an n-6 PUFA–enriched, high-

<table>
<thead>
<tr>
<th>Compound</th>
<th>HPLC (%) ± SD</th>
<th>MS (%) ± SD</th>
<th>MDA (%) ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>l-carnosine</td>
<td>12.8 ± 1.6</td>
<td>41 ± 0.6</td>
<td>4 ± 0.4</td>
</tr>
<tr>
<td>Carnosinol</td>
<td>36.4 ± 1.6</td>
<td>13.6 ± 2.1</td>
<td>3.8 ± 0.3</td>
</tr>
<tr>
<td>Pyridoxamine</td>
<td>2.5 ± 1.4</td>
<td>0.1 ± 0.1</td>
<td>1.3 ± 0.1</td>
</tr>
<tr>
<td>Hydralazine</td>
<td>48.6 ± 4.2</td>
<td>14.2 ± 2.0</td>
<td>3.1 ± 0.5</td>
</tr>
<tr>
<td>Aminoguanidine</td>
<td>34.7 ± 2.7</td>
<td>11.3 ± 2.0</td>
<td>9.4 ± 0.7</td>
</tr>
</tbody>
</table>

HPLC %, fraction (percentage of total) of aldehyde quenched after a 24-hour
incubation; MS %, percentage of carbonylated ubiquitin formation inhibited
by the tested compounds; UC50, concentration required to inhibit MDA-induced
ubiquitin carbonylation by 50%. Carnosinol activity was compared with that
of carnosine and of well-known RCS-sequestering agents. MGO activity was
evaluated by HPLC and MS assays and the dose-dependent activity toward MDA
by MS. Data for the quenching activity of l-carnosine, pyridoxamine, hydralazine,
and aminoguanidine toward MDA and MGO are from ref. 30.
istered to mice starting after 8 weeks on the HFHS diet, and a control group was fed normal chow for the duration of the study (Supplemental Figure 5A). As expected, the HFHS diet increased body weights and adiposity in both WT and GPx4 +/– mice (Supplemental Figure 5, B and C), and carnosinol had no effect on these parameters, although fasting serum triglyceride and cholesterol levels were decreased in the carnosinol-treated mice (Supplemental Table 4). We detected no effect of carnosinol on whole-body energy expenditure (Supplemental Figure 6) or food or water intake (data not shown) in mice compared with those on a HFHS diet alone. Carnosinol treatment led to enhanced glucose disposal following oral glucose challenge in obese WT, but not GPx4 +/–, mice (Figure 5, A–C). This improved glucose disposal may be attributed in part to increased skeletal muscle insulin sensitivity, as carnosinol normalized insulin-stimulated 2-deoxyglucose (2-DOG) uptake in extensor digitorum longus (EDL) to levels similar to those in control-fed lean mice (Figure 5D), although this effect was not seen in soleus (Figure 5E). We observed no improvement in insulin sensitivity with carnosinol in soleus tissue from obese WT mice, although we observed a modest improvement in obese GPx4 +/– soleus (Figure 5F).

Reactive aldehyde derivatives of lipid peroxidation have recently come into focus as novel redox signaling agents that, paradoxically, have beneficial and pathological roles, depending on the concentration and tissues and cells affected (11, 41). Protein-HNE adducts are known to be increased in skeletal muscle of type 2 diabetic and obese/insulin-resistant patients, and these adducts are associated with the severity of insulin resistance (42). Experimental models have reported that lipid peroxidation in skeletal muscle blunts insulin signaling and glucose uptake in skeletal muscle.
via HNE adduct formation (43, 44). Lipid peroxidation and HNE adduct formation have also emerged as potential causal factors in hyperinsulinemia and the eventual loss of pancreatic β cell function in models of obesity/overnutrition (45-47), which is further evidence of a system-wide pathological role for reactive aldehydes in metabolic syndrome. Here, HNE adduct formation in mixed gastrocnemius skeletal muscle (Figure 5, F and G) and pancreas (Figure 5, H and I) increased with a HFHS diet, particularly in GPx4+/– mice. Carnosinol effectively mitigated the accumulation of these adducts in both WT and GPx4+/– mice on a HFHS diet, suggesting that this may be one aspect of the mechanism by which carnosinol improves systemic glycemic control in obese mice.

Liver inflammation and fibrosis are mitigated by carnosinol in mouse models of severe carbonyl stress– and diet-induced obesity
Nonalcoholic steatohepatitis (NASH) is among a cluster of obesity-related pathologies and is closely linked with insulin resistance. NASH is distinct from fatty liver disease, in that intralobular inflammation and fibrosis are present in addition to the steatosis (48). The fibrosis component of this disease has been specifically identified by numerous studies to be the most likely to predict adverse outcomes in patients (49). Thus, therapeutic strategies that specifically target liver inflammation and fibrosis in obese patients will be highly valued by clinicians (50). A number of studies have implicated RCS as having a causal role in NASH, due in large part to the known effect of RCS on activation of the proinflammatory receptor for AGEs (RAGE) pathway (22, 51). In the recent study from our group, obese GPx4+/– mice were found to have elevated RCS (not shown) mRNA levels. In the present study, RAGE expression in liver was unchanged in WT and GPx4+/– mice on a HFHS diet, but carnosinol significantly decreased RAGE expression in both groups (Figure 6A). Expression of the proinflammatory cytokines TNF-α and IL-6 was significantly higher in the livers of HFHS-fed mice, and carnosinol mitigated the expression of TNF-α, but not IL-6, in this tissue (Figure 6, B and C).

To further examine the effect of carnosinol on liver pathology in obesity, sections of liver tissue were fixed and stained with oil red O and Picrosirius red to label triglycerides and collagen, respectively. As in the previous study, we observed that the HFHS diet increased liver triglyceride content in WT and GPx4+/– mice, with substantially greater lipid deposition occurring in the obese GPx4+/– mice (Figure 6, D and E). Interestingly, we found that the increase in liver triglyceride deposition with a HFHS diet was not accompanied by changes in serum triglycerides (Supplemental Table 4), although this could be attributable to necropsy being performed while the mice were in a fasted state. Although total triglyceride content in liver did not significantly change with carnosinol (data not shown), we observed a shift from predominantly macro- to microvesicular steatosis with carnosinol treatment in obese WT and GPx4+/– mice (Figure 6, D and E). Interestingly, we found that the decrease in liver triglyceride deposition with a HFHS diet was not accompanied by changes in serum triglycerides (Supplemental Table 4), although this could be attributable to necropsy being performed while the mice were in a fasted state. Although total triglyceride content in liver did not significantly change with carnosinol (data not shown), we observed a shift from predominantly macro- to microvesicular steatosis with carnosinol treatment in obese WT and GPx4+/– mice (Figure 6, D and E).

In order to determine whether a HFHS diet and/or carnosinol treatment alters collagen cross-linking in the tissue, we measured the ratio of insoluble to soluble hydroxyproline quantity in liver extracts. A HFHS diet led to an increase in the insoluble hydroxyproline fraction in WT and GPx4+/– livers (Figure 6G), and carnosinol normalized these levels. Indeed, carnosinol treatment was associated with significantly lower levels of collagen overall, including soluble hydroxyproline content (Figure 6H), regardless of diet or genotype. This decrease in collagen content caused by carnosinol treatment in obese WT and GPx4+/– mice does not appear to be due to decreased collagen gene expression, as we detected no effect of carnosinol on collagen-1 (Figure 6I) or collagen-4 (not shown) mRNA levels.

Pharmacodynamics and reactivity of carnosinol in mouse models of obesity and in human serum
In order to ascribe the beneficial effects of carnosinol in our obese mouse models to its aldehyde-scavenging capacity in vivo, we developed a high-resolution liquid chromatography–MS–based (LC-MS-based) approach to measure free carnosinol and carnosi- nal-aldehyde adducts in biological material. Pooled samples of tissue from skeletal muscle, pancreas, liver, kidney, and adipose tissue were analyzed using this method, and free carnosinol was found to be detectable only in liver, kidney, and adipose tissue (Supplemen-
The tandem mass spectra of the signal at 269.16 from traces in Figure 7A were able to confirm the structure of the putative adduct (Supplemental Figure 7). No signal of adducts with other reactive carbonyl species (e.g., HNE, HHE, malondialdehyde) were detectable in control or carnosine-supplemented animals. This may be due to the instability of carnosinol-HNE adducts in liver homogenate, which we determined in separate experiments. Only 63.24% ± 3.47% of an initial amount of 5 μM carnosinol-HNE conjugate was detectable after a 2-hour incubation in liver homogenate, with an estimated half-life of 3 hours according to a first-order decay model. We did not observe this instability with the carnosinol-ACR adduct in liver homogenate.

To further characterize carnosinol reactivity and the stability of aldehyde adducts, we tested the HNE-scavenging capacity of carnosinol in human serum by spiking the serum with a known Table 5). We found that carnosinol levels in liver and kidney were lower in the HFHS-fed GPx4+/– mice than in the WT mice. Importantly, the lower concentration of free carnosinol in these mice corresponded with a higher concentration of carnosinol-ACR adducts in the liver as compared with concentrations in HFHS-fed WT mice treated with carnosinol (Figure 7A). Specifically, a signal at 269.16082 (i.e., the expected m/z value for the carnosinol-ACR Michael adduct) was detectable at the same retention time observed in a reference sample prepared by spiking an aliquot of carnosinol-ACR adduct in rat liver homogenate at a final concentration of 0.5 μM (Figure 7A, bottom). The narrow mass range considered for extracting the chromatograms in Figure 7A (i.e., 1 ppm mass tolerance) and the reproducible retention time if compared with a spiked sample are conclusive evidence of the formation of a carnosinol metabolite after supplementation.

Figure 5. Therapeutic effect of carnosinol on metabolic homeostasis and carbonyl stress in mouse models of diet-induced obesity. The effect of a HFHS diet with and without carnosinol treatment is shown for glucose tolerance (A–C, n = 8) and insulin sensitivity of EDL (D) and soleus muscle (E) tissue upon termination of the study for each group (n = 6). Representative immunoblots for HNE adducts in whole-tissue homogenates prepared from mixed gastrocnemius skeletal muscle (F) and pancreas (H) tissue (n = 3 mice per group), along with the corresponding densitometric analysis (G and I). †P < 0.01 versus control diet within each respective genotype. A 2-way ANOVA followed by Tukey’s multiple comparisons test was used to test for the main effect of the treatment within each genotype. Tandem mass spectra of the signal at 269.16 from traces in Figure 7A were able to confirm the structure of the putative adduct (Supplemental Figure 7). No signal of adducts with other reactive carbonyl species (e.g., HNE, HHE, malondialdehyde) were detectable in control or carnosine-supplemented animals. This may be due to the instability of carnosinol-HNE adducts in liver homogenate, which we determined in separate experiments. Only 63.24% ± 3.47% of an initial amount of 5 μM carnosinol-HNE conjugate was detectable after a 2-hour incubation in liver homogenate, with an estimated half-life of 3 hours according to a first-order decay model. We did not observe this instability with the carnosinol-ACR adduct in liver homogenate.

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The extent to which carbon-yl stress plays a causal role in the metabolic disorders of obesity has remained unclear, however. In the present study, we sought to investigate the pathological role of RCS in obesity with a ratio-nal design and pharmacological evaluation of carnosinol, a chem-ical analog of l-carnosine with high oral bioavailability and a RCS-scavenging capacity. Following the design and characteriza-tion of carnosinol, we performed a comprehensive set of in vitro and in vivo studies profiling its effects across a range of experi-mental models, including rodent models of diet-induced obesity and metabolic syndrome. Our findings support the hypothesis that sugar- and lipid-derived RCS have a causal role in metabolic dis-orders associated with obesity. Moreover, we show that carnosinol concentration of preformed carnosinol and HNE as well as with carnosinol alone. Using our high-resolution LC-MS approach, we found that not only was carnosinol-HNE detectable, but unlike in tissues, the adduct was highly stable in human serum, with more than 85% of the carnosinol-HNE adduct remaining after a 2-hour incubation (Figure 7B and Supplemental Figure 8). Moreover, carnosinol can form adducts with trace amounts of HNE that are already present in human serum.

**Discussion**

An association between sugar- and lipid-derived RCS and obesity has been known for many years. The extent to which carbon-yl stress plays a causal role in the metabolic disorders of obesity has remained unclear, however. In the present study, we sought to investigate the pathological role of RCS in obesity with a ratio-nal design and pharmacological evaluation of carnosinol, a chem-ical analog of l-carnosine with high oral bioavailability and a RCS-scavenging capacity. Following the design and characteriza-tion of carnosinol, we performed a comprehensive set of in vitro and in vivo studies profiling its effects across a range of experi-mental models, including rodent models of diet-induced obesity and metabolic syndrome. Our findings support the hypothesis that sugar- and lipid-derived RCS have a causal role in metabolic dis-orders associated with obesity. Moreover, we show that carnosinol...
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RESEARCH ARTICLE

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jci.org   Volume 128   Number 12   December 2018

53). Thus, the effect of carnosinol on RCS in adipose tissue, and the resulting impact on metabolic parameters, may have played a significant role in the outcomes of the present study.

Though it is clear that lipid peroxidation and RCS do indeed have deleterious effects in the context of obesity, there are also complex time-, concentration-, and tissue-dependent factors to consider. Studies in pancreatic β cells have shown that short-term exposure to low levels of lipid peroxidation and subsequent 4-hydroxyalkenal formation stimulates an adaptive response mediated by PPAR δ, which causes increased glucose-stimulated insulin secretion (46, 47). In a previous study, we found that numerous enzymes involved in redox buffering and fatty acid metabolism are enhanced in rat myocardium in parallel with lipid peroxidation following 12 weeks of a HFHS diet (54). Such a “hormetic” effect of oxidative stress has been documented by other groups using similar obese models (55). Certainly, much remains to be determined about the precise mechanisms and factors involved in lipid peroxidation and subsequent RCS in obesity.

A major determinant of lipid peroxidation now known to be intimately involved in regulating disease pathology is the expression and activity of the selenoenzyme GPx4. As 1 of only 3 antioxidant enzymes essential for development (56), GPx4 has recently been the target of intense scrutiny by investigators. In particular, a critical role for GPx4 in regulating ferroptosis and subsequent organ failure has been reported (57–61). Furthermore, genetic variants of gpx4 that result in diminished activity and/or enzyme content are associated with obesity and cardiovascular disease in humans (62–64). We previously reported that GPx4-deficient (GPx4+/–) male mice acquire severe insulin resistance, steatohepatitis, and cardiomyopathy on a HFHS diet and that diabetic patients have diminished GPx4 content and elevated HNE adduct levels in their heart tissue compared with levels in age-matched nondiabet-

Figure 7. Pharmacodynamics and aldehyde-scavenging capacity of carnosinol. LC-MS analyses were performed on pooled, prepared liver homogenates from WT and GPx4+/– mice from each of the treatment groups. (A) Single-ion chromatograms of the carnosinol-ACR adduct (269.16082 m/z, ± 1 ppm). The bottom trace in A is the reference standard used in this analysis and was prepared by spiking rat liver homogenate with 0.5 μM carnosinol-ACR adduct. (B) LC-MS analysis showing the single-ion chromatogram of carnosinol-HNE adducts in human serum spiked with carnosinol and HNE (top), carnosinol only (middle), and serum only. Values shown in the green boxes denote the corresponding concentration of these adducts.
ic patients (15). In the present study, we used male GPx4+/– mice as a model to pharmacologically assess carnosinol in a context of enhanced lipid peroxidation–induced by obesity. As shown in Figure 5, GPx4+/– mice had greater levels of HNE adducts in skeletal muscle and pancreas following a HFHS diet compared with levels in WT mice. Carnosinol was effective at mitigating HNE adducts in these organs from both WT and GPx4+/– mice but was only fully effective at mitigating insulin sensitivity and glucose tolerance in WT mice. A reason for this may be that the dose of carnosinol used in this study was insufficient for a complete therapeutic effect in the GPx4+/– mice. Evidence supporting this conclusion is that there was a greater concentration of the carnosinol-ACR adduct in HFHS-fed GPx4+/– livers (Figure 7A), which corresponded with a lower concentration of free carnosinol in the livers and kidneys of HFHS-fed GPx4+/– mice (Supplemental Table 5). These findings are consistent with a greater aldehyde load in the GPx4+/– mice on a HFHS diet and a correspondingly greater demand for carnosinol in these mice as compared with WT mice on a HFHS diet. Taken together, we have uncovered compelling evidence that the therapeutic effect of carnosinol is mediated, at least in part, by its RCS-scavenging capacity.

Other compounds that scavenge RCS have shown significant therapeutic potential in preclinical studies using cardiovascular, renal, and metabolic disease models. Of all those tested to date, aminoguanidine and members of the B vitamin family (pyridoxamine, thiamine) have shown the most promising therapeutic effects (29, 65). However, in randomized, controlled trials these compounds have yielded largely disappointing results (with the exception of pyridoxamine), probably because of their promiscuous reactivity to all aldehydes (e.g., pyridoxal). Such off-target effects, particularly in the case of aminoguanidine (66), have made existing RCS-scavenging therapies unsuitable for mainstream clinical practice (67). These findings underscore the complexity of carbonyl biochemistry in physiological systems and illustrate that, in principle, agents that scavenge or block RCS have clinical potential. Thus, the development of novel RCS-scavenging therapies is warranted.

Recently, interventional studies based on a daily dose of L-carnosine (2 g/day for 12 weeks) in overweight and obese subjects resulted in the preservation of insulin sensitivity and insulin secretion and normalized glucose tolerance compared with the placebo group. Furthermore, L-carnosine supplementation reduced 2-hour insulin levels after oral glucose tolerance tests (OGTTs) in a subgroup of individuals with impaired glucose tolerance (68). Although L-carnosine was not detected in serum, it was easily detectable in the urine as a conjugate with ACR (69). The in vivo RCS-scavenging capacity of L-carnosine was further confirmed by 2 independent studies reporting the presence of covalent adducts between carnosine and lipid peroxidation–by-products (e.g., ACR, HNE, HHE) in the urine of human volunteers (70, 71).

Given the findings of these translational studies, it would seem that orally supplemented L-carnosine, despite its rapid serum degradation by carnosinases, provides a promising therapeutic effect in obese subjects, one that may be linked to a RCS-scavenging mechanism. However, several studies indicate that the beneficial effect of L-carnosine is strictly associated with its serum stability and bioavailability. An allelic variant of CN1 that results in increased enzyme activity (and, consequently, lower serum L-carnosine levels) is linked to diabetic nephropathy (72). Transgenic db/db mice overexpressing human CN1 and having reduced serum carnosine levels exhibit higher fasting plasma glucose and HbA1c levels, to the extent that the glucosuria in these mice causes a significant reduction of body weight (73). It can be inferred from these studies that serum L-carnosine levels are directly linked to glycemic control and that human CN1 presents an attractive drug target for this patient population.

Our group envisioned 2 approaches that could circumvent the challenge posed by endogenous carnosinases: (a) inhibition of these enzymes during simultaneous oral L-carnosine therapy; and (b) design of stable carnosine peptide mimetics resistant to carnosinases. The former approach was not pursued because of a high potential for toxicity, as CN1 is critical for neurotransmitter production, histidine metabolism, and other vital functions. To address the latter approach, we previously synthesized and characterized the carnosinase-resistant enantiomer D-carnosine and found that this compound exhibited a significant RCS-scavenging effect in vitro and in vivo, but had poor intestinal absorption due to a low affinity for and transport by PepT1 (74). Other groups have successfully made carnosine derivatives that are resistant to carnosinase and initially showed therapeutic potential, but failed in preclinical testing largely because of decreased absorption (75, 76). An octylester derivative of D-carnosine showed enhanced intestinal absorption and therapeutic effects in a mouse model of cardiometabolic disease (34), but the translational applicability of this compound is low because of potency issues that likely stem from dosing limitations resulting from diminished absorption. Carnosinol is the most promising L-carnosine derivative that has been synthesized at this point. It is easily transportable by PepT1 and not metabolized by CN1 (Figure 1), it maintains the outstanding safety profile of L-carnosine, and it was determined to be more reactive toward HNE, MGO, and ACR compared with L-carnosine (Tables 1 and 2 and Supplemental Table 2). As such, it represents a promising lead compound for counteracting RCS, particularly lipid peroxidation–derived α,β-unsaturated aldehydes, which have recently come to the forefront as primary driving forces in chronic disease.

Conclusions and translational perspective. The present study supports the hypothesis that sugar- and lipid-derived RCS have a causal role in metabolic disorders associated with obesity and further demonstrates that carnosinol represents a very promising lead compound in a new class of RCS-scavenging agents derived from the histidyl dipeptide L-carnosine.

Methods
A more detailed description of the materials and methods used in this study can be found in the supplemental material.

Materials and reagents. Solvents for HPLC and LC-MS and all analytical-grade chemicals were purchased from Sigma-Aldrich. L-carnosine and carnosinol were gifts of Renato Canevotti and Stefania Gagliardia (Flamma S.p.A.). For molecular and biochemical endpoints, other compounds that scavenge RCS have shown significant therapeutic potential in preclinical studies using cardiovascular, renal, and metabolic disease models. Of all those tested to date, aminoguanidine and members of the B vitamin family (pyridoxamine, thiamine) have shown the most promising therapeutic effects (29, 65). However, in randomized, controlled trials these compounds have yielded largely disappointing results (with the exception of pyridoxamine), probably because of their promiscuous reactivity to all aldehydes (e.g., pyridoxal). Such off-target effects, particularly in the case of aminoguanidine (66), have made existing RCS-scavenging therapies unsuitable for mainstream clinical practice (67). These findings underscore the complexity of carbonyl biochemistry in physiological systems and illustrate that, in principle, agents that scavenge or block RCS have clinical potential. Thus, the development of novel RCS-scavenging therapies is warranted.

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Laboratories. Additional details and descriptions of the ADMET studies are provided in the supplemental material.

Rodent models of obesity and metabolic syndrome. Both rat and mouse models of diet-induced obesity were used in this study. Eight-week-old male Sprague-Dawley rats (Harlan Laboratories) weighing 200±20 g were used for this study. Rats were housed under constant environmental conditions and were fed standard laboratory rat chow or a 60% HF diet (Mucedola S.R.L.) and tap water ad libitum. In particular, the control diet contained 60% corn starch (carbohydrates), 20% casein (protein), 0.3% methionine, 5% lard (fat), 8% cellulose, 5% mineral mixture and 1% vitamin mixture, and zinc carbonate 0.004%. The fructose diet contained all the ingredients except corn starch, which was replaced by an equal quantity of fructose. Animals were acclimatized for a period of at least 7 days before use in the study. Subsequently, rats were randomly divided into 5 groups. The control group received a standard rat chow diet for 6 weeks, whereas the other 4 groups of mice were given a fructose-enriched diet for 6 weeks.

For mouse models of diet-induced obesity, C57BL6/J female mice (The Jackson Laboratory) were crossed with male GPx4+/– mice, and the pups were genotyped by PCR using previously described primers (15). At 8 to 12 weeks of age, WT and GPx4+/– male age-matched littermates were randomly assigned to groups and individually housed. Mice were fed either a control (TD110367) or a HFHS (TD110365) diet from Harlan-Teklad Laboratories ad libitum for 25 weeks. The composition of this diet was a special formulation consisting of mixed saturated and n-6 PUFA (44.6% kcal/g fat), with a high-sucrose (34% kcal/g) content (54). After 8 weeks of the HFHS diet, half of the mice in the HFHS diet cohort (WT and GPx4+/–) were administered carnosinol at two doses: 10 and 45 mg/kg/day. An additional group was treated with 10 mg/kg/day rosiglitazone (GlaxoSmithKlein), and the fifth group continued with the fructose-enriched diet alone (HF group). Rosiglitazone was orally administered to rats by gastric gavage during the last 3 weeks of the study, while carnosinol was dissolved in the water. During all the experiments, rats had ad libitum access to food and water.

Three weeks after starting the fructose diet, two groups were treated with carnosinol at two doses: 10 and 45 mg/kg/day. An additional group was treated with 10 mg/kg/day rosiglitazone (GlaxoSmithKlein), and the fifth group continued with the fructose-enriched diet alone (HF group). Rosiglitazone was orally administered to rats by gastric gavage during the last 3 weeks of the study, while carnosinol was dissolved in the water. During all the experiments, rats had ad libitum access to food and water.

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Acknowledgments
Carnosinol was provided by Flamma S.p.A. This project was supported by funds from the Ministry of Education, Universities, and Research of Italy (MIUR) (PRIN 2009); the Regione Lombardia – MIUR (L.297–Art. 12/BioTech DM27909 and decree no. 6737 July, 2, 2009); and the NIH (R01HL122863 and R21AG057006, to EJA). The authors would like to thank Cherese Beatty (East Carolina University), who assisted with animal experiments and tissue histology. GV gratefully acknowledges Concetta De Stefano (University of Messina, Messina, Italy) for assistance in determining the chelation constants.

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Author contributions
GV, GA, and EJA made equivalent intellectual contributions to this manuscript and are the guarantors of all the data contained herein. GV (in silico modeling) and GA (bioanalysis and testing) were principally responsible for the design of carnosinol. EJA was principally responsible for the overall study design, experimental model development, and pharmacological testing of carnosinol and for writing the manuscript. LAK, KF, TBM, LR, EG, LC, M. Colzani, DDM, GR, and M. Carini designed and conducted experiments, analyzed data, and assisted with manuscript preparation. LR and EG designed and optimized the assay for LC-MS carnosinol-aldehyde adduct determination. RC and SG contributed to carnosinol synthesis.

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