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Short Report

Neuregulin 1 affects leptin levels, food intake and weight gain in normal-weight, but not obese, db/db mice

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Abstract

\textbf{Aim.} – Studies in vitro have highlighted the potential involvement of neuregulin 1 (NRG1) in the regulation of energy metabolism. This effect has also been suggested in vivo, as intracerebroventricular injection of NRG1 reduces food intakes and weight gain in rodents. Thus, it was hypothesised that NRG1 might affect serum leptin levels in mice.

\textbf{Methods.} – Weight, food intakes, energy expenditure, spontaneous physical activity and serum leptin levels were evaluated in normal-weight C57BL/6JRJ mice following intraperitoneal administration of NRG1 (50 \textmu g/kg, three times/week) or saline for 8 weeks. Based on the results of this first experiment, leptin-resistant obese db/db mice were then given NRG1 for 8 weeks.

\textbf{Results.} – Leptin serum concentrations were six times higher in C57BL/6JRJ mice treated with NRG1 than in the animals given saline. NRG1 treatment also reduced weight gain by 10\% and food intakes by 15\% compared with saline treatment, while energy expenditure remained unchanged. In db/db mice, serum leptin concentrations, weight gain, food intakes, energy expenditure and spontaneous physical activity were not altered by NRG1 treatment.

\textbf{Conclusion.} – The decrease in food intakes and weight gain associated with NRG1 treatment in C57BL/6JRJ mice may be partly explained by increased leptin levels, whereas db/db mice were not affected by the treatment, suggesting resistance to NRG1 in this pathological state.

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\textbf{Keywords:} NRG1; Obesity; Energy expenditure; Weight loss

\textbf{Abbreviations:} NRG1, Neuregulin 1 ;ErbB, Erythroblastic leukaemia viral oncogene homologue; RQ, Respiratory quotient; VHL, Vehicle (saline).

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1. Introduction

Neuregulin 1 (NRG1) belongs to the epidermal growth factor family. Its biological activity is mediated by erythroblastic leukaemia viral oncogene homologue (ErbB) receptors. NRG1 plays a fundamental role in both developing and mature heart tissue, and NRG1 treatment has positive effects in cardiovascular diseases [1]. Altered NRG1/ErbB signalling has been demonstrated during the development and progression of many tumours [2]. Complex crosstalk between ErbB2 receptors and leptin appears to take place in cancer [3], and ErbB2 induces leptin transcriptional upregulation in human breast epithelial cells [4].
As leptin plays a major role in energy balance regulation, and given the link between leptin and NRG1/ErbB signalling in cancer, it was hypothesized that NRG1 treatment might alter leptin levels.

Therefore, the effect of chronic (8-week) intraperitoneal (i.p.) NRG1 treatment on leptin, weight status, fat mass, food intakes, physical activity and energy expenditure was assessed in normal-weight mice and, based on the results of this first experiment, in leptin-resistant db/db mice also.

2. Methods

2.1. Animals

A total of 16 C57BL/6JRJ and 16 BKS(D)-Leprdb/JOrlRj mice (10 weeks old), provided by CERJ Janvier (Le Genest-Saint-Isle, France), were kept in temperature-controlled (20–22 °C) cages (single-housed) with free access to water and food (A04 diet; Scientific Animal Food and Engineering, France). All animal husbandry and experimental procedures were in accordance with the current legislation on animal experimentation, and were approved by the local ethics committee (CEMEA Auvergne, CE 97-12).

2.2. NRG1 treatment

Recombinant NRG1 (beta-2a; Zensun Sci and Tech, Shanghai, China) [5] at 50 μg/kg body weight or an equivalent volume of 0.9% NaCl (sodium chloride) was administered by i.p. injection three times a week for 8 weeks (n = 8 for each group). Mice were euthanised 3 days after the end of treatment.

2.3. Food intakes, body mass and body composition

Body weight and food intakes were recorded every week during the treatment period. One week before sacrifice, body composition was evaluated by quantitative nuclear magnetic resonance imaging (EchoMRI; Echo Medical Systems, Houston, TX, USA).

2.4. Daily energy expenditure and physical activity

VO₂, VCO₂ and physical activity were measured in mice using a PhenoMaster/LabMaster home cage system (TSE Systems, Bad Homburg, Germany). Energy expenditure was calculated using the Weir equation [6] from measurements of gas exchanges computed for each cage from data sampled every 5 min. Respiratory quotient (RQ) was determined from VCO₂/VO₂. Spontaneous activity was measured using three-dimensional meshes of light beams. For adaptation, mice were placed in individual calorimetric cages (22 °C) for 24 h prior to data collection. Daily energy expenditure and physical activity were computed for a 24-h period.

2.5. Leptin measurement

Serum leptin was evaluated with an ELISA kit (Abcam, Cambridge, MA, USA), and samples were read at 450 nm using a PowerWave spectrophotometer (US BioTek, North Shoreline, WA, USA).

2.6. Statistical analysis

SPSS Advanced Statistics software was used for the statistical analysis (IBM, Armonk, NY, USA). Data are presented as the mean ± SEM. One-way analysis of variance (ANOVA) with repeated measures was performed to assess weight and food intake changes following NRG1 or saline treatment. Unpaired Student’s t-test was used where appropriate. Statistical significance was set at P < 0.05.

3. Results

3.1. NRG1 increases serum leptin and decreases weight gain, food intakes and spontaneous physical activity in C57BL/6JRJ mice

On evaluating NRG1 effects in C57BL/6JRJ mice (Fig. 1), serum leptin was strongly increased compared with the saline vehicle (VHL)-treated animals (183.4 ± 46.6 pg.mL⁻¹ vs. 29.5 ± 5.1 pg.mL⁻¹, respectively; P < 0.05; Fig. 1A). The VHL group also gained 9.6% in body mass (from 25.1 ± 0.95 g to 27.6 ± 0.85 g) whereas, in NRG1-treated mice, weight remained stable (from 26.6 ± 0.52 g to 26.1 ± 0.43 g) with treatment (Fig. 1B). Similarly, food intakes increased by 15.3% in the VHL group (from 27.1 ± 0.59 g.week⁻¹ to 31.5 ± 1.5 g.week⁻¹; P < 0.05; Fig. 1C), but did not change in NRG1-treated mice (from 29.3 ± 0.57 g.week⁻¹ to 29.3 ± 0.69 g.wk⁻¹; Fig. 1C). At the end of the treatment period, body fat percentages were lower in the NRG1 than in the VHL group (12.5 ± 0.67% vs. 16.3 ± 0.91%, respectively; P < 0.05; Fig. 1D). Daily energy expenditure (Fig. 1E) and RQ (not shown) did not differ between the groups. Spontaneous activity over 24 h and at night was lower in the NRG1 than in the VHL group (288.2 ± 31.5 m vs. 502.6 ± 102.3 m and 227.7 ± 24.9 vs. 373.6 ± 56.1 m, respectively; P < 0.05), whereas diurnal activity was similar in both groups (Fig. 1F).

3.2. NRG1 has no effect on serum leptin, weight gain, food intakes or spontaneous physical activity in db/db mice

As NRG1 treatment increased serum leptin levels in C57BL/6JRJ mice, it was postulated that NRG1 effects might be blunted in leptin-resistant db/db mice. Serum leptin concentrations were comparable in both NRG1- and VHL-treated db/db mice (Fig. 2A). Similarly, weight (Fig. 2B) and food intakes (Fig. 2C) remained stable in both groups of db/db mice, and the percentage of body fat was also comparable in both groups at the end of treatment (Fig. 2D). Energy expenditure (Fig. 2E),
RQ (not shown) and spontaneous physical activity (Fig. 2F) did not significantly differ between groups.

4. Discussion

Our results show that, in C57BL/6JRJ mice, NRG1 treatment increases serum leptin concentrations, prevents weight gain and lowers food intakes compared with saline-treated controls. In contrast, NRG1 had no significant effects in db/db mice.

The strong leptin increase in C57BL/JJRJ mice after 8 weeks of NRG1 treatment might explain the decreases in food intakes, weight gain and body fat percentages, as shown previously with leptin treatment [7]. The cause of the leptin increase, however, is unknown. Leptin production is indirectly under insulin control via the PI3 K/AKT signalling pathway [8], and NRG1 stimulates this pathway in muscle cells [9]. ErbB2 activation upregulates leptin in human breast epithelial cells [4]. However, it is not known whether NRG1 administration can affect white adipose tissue, the main leptin producer [8]. The fact that ErbB receptors are expressed in human preadipocytes [10] allows the assumption that adipose tissue may be involved in the NRG1-mediated increase in leptin.

Alternatively, leptin levels might be regulated centrally, as the brain has been shown to express leptin and to release it into
Indeed, intracerebroventricular injection of NRG1 in hamsters had effects on food intakes and weight gain similar to those reported in our study [12], suggesting that the brain may be participating in the NRG1-mediated increase in circulating leptin, although it cannot be excluded that NRG1 effects may also be independent of leptin. Also, there is evidence to suggest that NRG1 could also have leptin-independent effects. Spontaneous physical activity was markedly decreased in normal-weight mice, as previously reported [12], while total energy expenditure was not affected by NRG1 treatment.

These results suggest that the resting metabolic rate, which represents a major component of total energy expenditure (about 86% in mice [13]), could compensate for the decreased energy expenditure from physical activity, resulting in unchanged total daily energy expenditure. Nevertheless, leptin is known to increase both spontaneous physical activity and energy expenditure [14]. Therefore, our present results suggest a complex action of NRG1 whereby food intakes and weight gain are both decreased possibly through an increase in leptin, while physical activity and energy expenditure are limited through unknown pathways. However, the possibility of a side-effect induced by NRG1 treatment that might promote a decrease in mouse locomotion cannot be ruled out, although human studies using comparable dose ranges have shown no such significant side-effect induced by chronic NRG1 treatment [15,16]. More studies are now needed to elucidate the origin of the
NRG1-induced increase in circulating leptin and the contribution of increased circulating leptin to altered food intakes and weight gain.

Unexpectedly, none of the effects observed in C57BL/6JRJ mice following NRG1 treatment were seen in the leptin-resistant obese db/db mice, and serum leptin concentrations did not increase. This result suggests that the NRG1/ErbB signalling pathway might be impaired in these mice. As NRG1 activates the PI3K/Akt pathway [9], insulin resistance in db/db mice could have inhibited leptin production, which is triggered by these kinases [10]. Moreover, palmitate impairs NRG1 signalling in rat cardiac myocytes [17]. This suggests that the dyslipidaemia commonly seen in metabolic diseases, and particularly in db/db mice, might perturb NRG1 signalling. Such an NRG1-resistant state could, in turn, result in silenced leptin upregulation.

In conclusion, our present study has demonstrated that chronic NRG1 administration increases serum leptin concentration in normal-weight mice, and this increase might partly explain the reduction in food intakes, body weight gain and body fat percentage in these mice. On the other hand, NGR1 treatment appeared to have no such effects in obese db/db mice, suggesting ‘NRG1 resistance’ in this pathological state.

Disclosure of interest

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