Leptin receptors in human skeletal muscle

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Submitted 20 November 2006; accepted in final form 10 January 2007

LEPTIN IS A 16-KDA HORMONE structurally related to cytokines (66) that plays a crucial role in the regulation of appetite and fat deposition (20, 38). This hormone is primarily released by white adipose tissue and acts on brain and peripheral receptors (19, 24, 45) that belong to the class I type cytokine receptor family (61, 65). There are at least six isoforms of leptin (19, 24), while reducing the body fat stores through regular exercise and/or dieting results in lower plasma leptin concentrations (28, 46, 62). Human obesity is characterized by a high concentration of leptin in plasma associated with leptin resistance (8, 60). Obesity also causes insulin resistance in humans (30, 44), which has been associated with raised plasma leptin concentrations, independent of body fat mass (50, 56). Leptin resistance could be caused by a downregulation and/or desensitization of OB-Rs, among other mechanisms.

In this study, we planned to test two hypotheses: first, that leptin receptors are expressed at the protein level in human skeletal muscle; and second, that the amount of OB-R protein expression in skeletal muscles depends on the basal concentration of leptin. To test these hypotheses, we carried out Western blot analysis in protein extracts obtained from human skeletal muscle biopsies and from a human hypothalamus. The hypothalamus protein extract was used as a control to verify that any band identified as a potential OB-R in muscle is also present in the hypothalamic protein extract, since the hypothalamus is rich in OB-R protein content (53). To test the second hypothesis, we determined whether plasma leptin concentration correlates with the protein expression of OB-R in skeletal muscle.

MATERIALS AND METHODS

Materials. The Complete protease inhibitor cocktail was obtained from Roche Diagnostics (Mannheim, Germany). The polyclonal rab-

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bit anti-human leptin receptor that recognizes the extracellular domain of human leptin receptor was obtained from Linco Research (St. Charles, MO). The recombinant human (RH) leptin R/Fc chimera, generated from DNA containing the extracellular domain of OB-R (amino acid residues 1-839) fused to the Fc region of human IgG1, was obtained from R&D Systems (McKinley Place). The monoclonal mouse anti-α-tubulin antibody was obtained from Biosigma (Madrid, Spain). The secondary horseradish peroxidase (HRP)-conjugated goat anti-rabbit and donkey anti-mouse antibodies were from Jackson Immunoresearch (West Grove, PA). The Hybrid-P Transfer membranes, Hyperfilm enhanced chemiluminescence (ECL), and the ECL ImmunoResearch (West Grove, PA). The Hybond-P transfer membranes, 1.9%) participated in this investigation. Written, informed consent was obtained, as reported above. Then, in the same gel, skeletal muscle protein extracts (50 μg) were run together with subcutaneous adipose tissue protein extracts (50 μg) were run together with subcutaneous adipose tissue protein extract samples containing 1, 2, 3, 4, or 5 μg of protein (Fig. 1A). Then Western blots were performed using a polyclonal rabbit anti-perilipin A antibody as described above. From the band densities obtained for perilipin, a standard curve was calculated by linear regression (all curves had a r² value ≥0.98) (Fig. 1B). The corresponding equation was used to calculate the maximal amount of fat that could be present in each muscle biopsy, assuming that skeletal muscle fibers have undetectable amounts of perilipin A (26, 47). To evaluate the specificity of the anti-OB-R antibody used in this investigation, competitive assays were performed with increasing amounts of RH leptin R/Fc (RH OB-R) chimera (0, 10, 100, 500 ng) preincubated with anti-OB-R antibody (diluted 1:1,500 in BSA blocking buffer) were performed for 1 h at room temperature. Antibody-specific labeling was revealed by incubation with a HRP-conjugated goat anti-rabbit antibody (1:20,000) or a HRP-conjugated donkey anti-mouse (1:10,000) antibody, both diluted in blotto blocking buffer and visualized with the ECL kit (Amersham Biosciences). Specific bands were scanned with the GS-800 Calibrated Densitometer and analyzed with the image analysis program Quantity One (Bio-Rad Laboratories, Hercules, CA). Data are reported as band intensity of immunostaining values (arbitrary units) obtained for OB-R relative to those obtained for α-tubulin. α-Tubulin content in the muscle biopsies was similar in all of the subjects analyzed (3.54 ± 0.22 arbitrary units of band density of immunostaining).

Competitive assays for OB-R. To evaluate the specificity of the anti-OB-R antibody used in this investigation, competitive assays were performed with increasing amounts of RH leptin R/Fc (RH OB-R) chimera (0, 10, 100, 500 ng) preincubated with anti-OB-R antibody (diluted 1:1,500 in BSA blocking buffer) were performed for 1 h at room temperature. Antibody-specific labeling was revealed by incubation with a HRP-conjugated goat anti-rabbit antibody (1:20,000) or a HRP-conjugated donkey anti-mouse (1:10,000) antibody, both diluted in blotto blocking buffer and visualized with the ECL kit (Amersham Biosciences). Specific bands were scanned with the GS-800 Calibrated Densitometer and analyzed with the image analysis program Quantity One (Bio-Rad Laboratories, Hercules, CA). Data are reported as band intensity of immunostaining values (arbitrary units) obtained for OB-R relative to those obtained for α-tubulin. α-Tubulin content in the muscle biopsies was similar in all of the subjects analyzed (3.54 ± 0.22 arbitrary units of band density of immunostaining).

Potential contamination by whole blood or subcutaneous adipose tissue. To assess if a small contamination by blood could influence the OB-R immunostainings, whole human blood protein extracts were obtained from two healthy subjects and processed for Western blot analysis as described above. Skeletal muscle biopsies may be contaminated by a small amount of adipose tissue, which may come from the adipose tissue accumulated between the muscle bundles and/or by subcutaneous fat tissue. Although the latter possibility was minimized by avoiding the use of suction, the amount of protein material coming from adipose tissue was always assessed in all muscle samples. For this purpose, a protein extract from subcutaneous adipose tissue was first obtained, as reported above. Then, in the same gel, skeletal muscle protein extracts (50 μg) were run together with subcutaneous adipose tissue protein extract samples containing 1, 2, 3, 4, or 5 μg of protein (Fig. 1A). Then Western blots were performed using a polyclonal rabbit anti-perilipin A antibody as described above. From the band densities obtained for perilipin, a standard curve was calculated by linear regression (all curves had a r² value ≥0.98) (Fig. 1B). The corresponding equation was used to calculate the maximal amount of fat that could be present in each muscle biopsy, assuming that skeletal muscle fibers have undetectable amounts of perilipin A (26, 47). To
The protein expression of the OB-R in human skeletal muscle was determined by Western blot analysis. The levels of OB-R were assessed in protein extracts from muscle and subcutaneous adipose tissue (SAT) and adipose tissue infiltration in human skeletal muscle biopsies. Protein extracts were prepared from muscle, SAT, and thigh subcutaneous adipose tissue (Fig. 2, A and D). The 170-kDa band was detected in skeletal muscle and hypothalamus protein extracts (Fig. 2A). The other two bands migrating at 128 and 98 kDa were identified in skeletal muscle and thigh subcutaneous adipose tissue protein extracts (Fig. 2A). No immunosignal was observed in the absence of primary antibody (data not shown). The 128-kDa isoform was not detected in four subjects, implying that the level of expression was undetectable or that the 128-kDa isoform was not expressed at all in the skeletal muscle (Fig. 2A). In the latter case, the 128-kDa observed in some subjects indicated contamination by surrounding adipose tissue or adipocytes located in between the muscle fibers. In contrast, the 170- and the 98-kDa isoforms were detected in all 14 subjects (Fig. 2A).

The densities of the 98-kDa and the 128-kDa OB-R bands \((r = 0.76, P < 0.01)\) and the 170-kDa and the 98-kDa OB-R bands \((r = 0.74, P < 0.01)\) were correlated. No relationship

\[ r^2 = 0.99 \]

against its extracellular domain. This antibody revealed the presence of three bands in which molecular mass was ~170, 128, and 98 kDa (Fig. 2, A and D). The 170-kDa band was detected in skeletal muscle and hypothalamus protein extracts (Fig. 2A). The other two bands migrating at 128 and 98 kDa were identified in skeletal muscle and thigh subcutaneous adipose tissue protein extracts (Fig. 2A). The 98-kDa band was also found in hypothalamus protein extracts (Fig. 2A). No immunosignal was observed in the absence of primary antibody (data not shown). The 128-kDa isoform was not detected in four subjects, implying that the level of expression was undetectable or that the 128-kDa isoform was not expressed at all in the skeletal muscle (Fig. 2A). In the latter case, the 128-kDa observed in some subjects indicated contamination by surrounding adipose tissue or adipocytes located in between the muscle fibers. In contrast, the 170- and the 98-kDa isoforms were detected in all 14 subjects (Fig. 2A).

The densities of the 98-kDa and the 128-kDa OB-R bands \((r = 0.76, P < 0.01)\) and the 170-kDa and the 98-kDa OB-R bands \((r = 0.74, P < 0.01)\) were correlated. No relationship
was observed between the densities of the 170-kDa and 128-kDa OB-R bands.

Specificity of the antibody: competitive assays. Competitive assays with RH leptin R/Fc chimera as a competitive blocker of the antigen-antibody interaction showed that the antibody was able to bind specifically to the three bands detected (Fig. 3). This implies that the OB-R bands share a common epitope with the RH leptin R/Fc chimera, which contains an amino acid sequence of the extracellular domain of the human OB-R.

Contribution of adipose tissue to the OB-R band density in skeletal muscle tissue. Assuming that muscle fibers do not express perilipin A (47), the band density of perilipin A measured by immunoblotting (Fig. 1) and normalized by expressing perilipin A (47), the band density of perilipin A measured for a control (0 ng of RH OB-R). *P < 0.01 vs. 0 ng of RH OB-R.

Fig. 3. The anti-OB-R antibody recognized specifically the three OB-R bands detected in the muscle protein extracts. Increasing amounts of recombinant human (RH) leptin R/Fc (RH OB-R) chimera (0, 10, 100, 500 ng) were preincubated with anti-OB-R antibody (1:2,000). OB-R protein expression from muscle extracts was analyzed by immunoblotting using the preincubation solution. A: representative Western blot analysis with different preincubation solutions in the same muscle protein extract (50 μg). B: representative immunoblot with the α-tubulin antibody as a loading control. C: densitometric percentage of OB-R immunostaining values (band quenching) in presence of increasing amounts (10, 100, 500 ng) of RH OB-R relative to those observed for a control (0 ng of RH OB-R). *P < 0.01 vs. 0 ng of RH OB-R.

Fig. 4. Assessment of the contribution of different amounts of adipose tissue on the optical density of the three OB-R bands and determination of the minimal amount of muscle protein extract needed to detect the presence of OB-R. Protein extracts were obtained from human muscle and SAT, and OB-R protein expression was determined by Western blot using a polyclonal rabbit anti-OB-R antibody. A: representative immunoblot performed with increasing amounts of protein extracts (1, 2, and 4 μg) from SAT added to 50 μg of muscle protein extract. B: representative Western blot performed with increasing amounts of muscle protein extracts (0 to 50 μg) added to 10 μg of protein extract from SAT.

- **A**
  - SAT protein extracts (50 μg) + increasing SAT protein extracts (μg)
  - kDa (10 μg)
  - 0 1 2 4 0 1 2 4
  - 170 128 98

- **B**
  - SAT protein extracts (10 μg)
  - kDa (10 μg)
  - 0 1 2 5 10 20 30 40 50
  - 170 128 98

**DISCUSSION**

In the present study, we hypothesized that the OB-R protein should be expressed in human skeletal muscle. This was based
on previous studies revealing the presence of OB-R mRNA in human skeletal muscle (17) and cultures of primary skeletal muscle cells (55), and also on the fact that primary skeletal muscle cells in culture respond to leptin by increasing ERK activity (55) and/or AMP-activated protein kinase activity and fatty acid oxidation (41, 59). This study confirms this hypothesis and describes a Western blot-based procedure to assess OB-R protein. This immunoblotting analysis was carried out using a polyclonal rabbit anti-human OB-R antibody in protein extracts obtained from muscle biopsies and revealed the presence of a dense band with a molecular mass close to 98 kDa and another two less intense bands, with molecular masses of 128 and 170 kDa. The 128 and 98 kDa bands were in agreement with the molecular mass of the short and long isoforms of OB-R (OB-Ra and OB-Rb, respectively), detected in other human tissues including brain, liver, digestive tract, umbilical cord, and fetal membranes (2, 3, 14, 22, 39). Furthermore, the 170-kDa band was compatible with the molecular mass observed for OB-Rb in human umbilical venous endothelial cells (13). Our results also demonstrate that the density of these three bands was reduced in competitive Western blot assays performed with increasing concentrations of RH leptin R/Fc chimera, which contains the extracellular domain (aa residues 1-839) of OB-R. These data suggest that the antibody used in this study recognized specifically the three OB-R bands detected in skeletal muscle and that muscular tissue OB-R proteins detected with this antibody contain the extracellular domain of the human OB-R.

These results implied that human skeletal muscle expresses the long and short isoforms of the leptin receptor. However, skeletal muscle is a complex tissue, and some adipose tissue (or adipocytes) may be present in between or around the muscle fibers and/or bundles (25, 29). Only the intermuscular adipose tissue (IMAT) that was visible could be removed during the manipulation of the muscle biopsies. This means that, in any muscle biopsy, there is always the potential for contamination by IMAT, which may be irrelevant for many purposes, but critical in this study. Whole body IMAT has been measured using multislice MRI (25, 29). The IMAT compartment includes IMAT that is located between muscle groups and beneath the muscle fascia and IMAT that is distributed within individual muscles visible on MRI images. IMAT mean values of 1.7, 2.2, and 2.5% have been reported in men having a mean percentage of body fat of 10.8, 25.3, and 20.2%, respectively (25, 29). Using a different approach that allows a physical separation of adipocytes from the muscle fibers in surgical muscle biopsies, Mingrone et al. (40) reported that intermuscular triglycerides represented 3.1 and 15.9% of the muscle mass in lean and obese subjects, respectively, which is equivalent to 4-20% in mass of adipose tissue, assuming that triglycerides represent ~80% of the adipocyte composition. In the present investigation, we observed that 2.4% of the proteins extracted from the muscle biopsies were from IMAT. This implies that IMAT mass in our muscle biopsies should have attained a higher value, which could only have been ascertained by knowing the protein composition of the muscle and adipose tissue in this location. However, the important point to bear in mind is that even a “clean” skeletal muscle biopsy always contains a significant amount of adipose tissue, a fact that has been often overlooked in other studies examining the expression of OB-R mRNA (17, 34, 48).

Solberg et al. (55) reported the existence of a functional long isoform of the OB-R in primary skeletal muscle cells derived from human skeletal muscle biopsies. To obtain these cells, the authors first separated the satellite cells by dissection and successive incubations with trypsin/EDTA. Then the satellite cells were grown in culture wells where they differentiated into myoblasts and fused together, leading to the formation of myotubes. When these myotubes were exposed to leptin, they responded by ERK activation, with a small increase in fatty acid oxidation. A similar stimulation of fat oxidation by leptin has also been reported in cultured myotubes derived from lean but not obese humans (59). However, it should be considered that myotubes may express different proteins from adult muscle fibers in vivo and that, during the process of in vitro differentiation, some satellite cells could have differentiated into adipocytes (52). Using an isolated rectus abdominis muscle preparation from lean and obese humans, Steinberg et al. (60) observed that leptin promotes fat oxidation only in lean subjects, when stimulated at high nonphysiological leptin concentrations (in the absence of insulin and other hormonal factors). Although these findings indirectly suggest the presence of a functional leptin receptor in human skeletal muscle, this in vitro preparation would likely contain a considerable amount of IMAT and other cell types, which could account differentially for the effects reported in fat oxidation.

The present investigation clearly shows that the 170-kDa OB-R isoform is only present in the muscle fibers and is not detectable in adipose tissue. However, both the 98- and 128-kDa bands could originate from the IMAT. This is further demonstrated by the fact that loading the gels with increasing amounts of protein extracts from subcutaneous adipose tissue increased the staining intensity corresponding to the 98- and 128-kDa bands, without any effect on the 170-kDa band. Knowing the amount of protein from adipose tissue present in each biopsy and the amount of 98- and 128-kDa OB-R density present in the subcutaneous adipose tissue, we have calculated that IMAT is able to explain all of the 128-kDa OB-R band density and 89% of the 98-kDa OB-R band density. The lack of antibodies specific for the 170- and 98-kDa isoforms impedes our ability, using immunohistochemical techniques, to resolve whether the 98 kDa is really present at the protein level in the muscle fibers.

Although a circulating form of the leptin receptor (OB-Re) lacking the transmembrane and intracellular domains (24) may contaminate the skeletal muscle samples, this isoform was not recognized by the anti-OB-R antibody used in this investigation, since Western blot analysis loading up to 100 μg of protein extract from blood leucocyte fraction was negative for OB-R (data not shown). This is likely due to structural and/or compositional differences between the extracellular domain of the OB-Re and that of the OB-Ra, OB-Rb, and OB-Rf isoforms (1). Thus we can rule out contamination by blood as source of OB-R immunoreactivity in our muscle samples.

The presence of a long isoform of the leptin receptor in the skeletal muscle fibers might have important implications for the understanding of the metabolic regulation of human energy metabolism and may be critical to unravel the physiopathology of the metabolic syndrome and insulin resistance (57, 59). The 170-kDa band could very well be the main ligand for leptin in skeletal muscle (9, 11, 12, 61). It has also been shown that this isoform phosphorylates in response to leptin binding (8), and...
this phosphorylation has been linked to the activation of intracellular cascades with subsequent effects on fatty acid transport and metabolism (41, 42, 58).

In summary, this study shows that a long isoform of the leptin receptor with a molecular mass close to 170 kDa is expressed at the protein level in human skeletal muscle. The amount of 170-kDa protein appears to be independent of the basal concentration of leptin in serum. In addition, we describe a procedure based on the determination of perilipin A content, a protein exclusive of adipocytes, to determine the degree of adipose tissue infiltration in human muscle biopsies. The latter procedure was critical for the interpretation of our results.

Adipose tissue contamination must be assessed when using rough protein extracts from skeletal muscle, if the aim is to study molecules that may also be present in IMAT. Future studies with longitudinal studies in dieting and/or exercising humans, should be carried out to establish the role of this isoform of the leptin receptor in the regulation of skeletal muscle metabolism.

ACKNOWLEDGMENTS

The authors thank Dr. Andrew S. Greenberg for kindly providing the anti-perilipin A antibody. Special thanks are given to José Navarro y Guerra del Río for excellent technical assistance and to Ana Navarro y Guerra del Río for anti-perilipin A antibody. Special thanks are given to José Navarro de Tuero for excellent technical assistance and to Ana Navarro y Guerra del Río for anti-perilipin A antibody. Special thanks are given to all subjects who volunteered for the study molecules that may also be present in IMAT. Future studies with longitudinal studies in dieting and/or exercising humans, should be carried out to establish the role of this isoform of the leptin receptor in the regulation of skeletal muscle metabolism.

REFERENCES


