

# Discovery and uses of pegvisomant: a growth hormone antagonist

Odkrycie i zastosowanie pegwisomantu: antagonisty hormonu wzrostu

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#### Abstract

Growth hormone (GH) is a well established participant in several complex physiological processes including growth, differentiation, and metabolism. Recombinant human GH is a drug that has been approved for use for several clinical conditions where the action of GH is diminished or completely lacking. Thus there is considerable interest in developing novel drugs that modify the function of GH. Only in the last several decades have the detailed structural features of GH along with its interaction with its receptor been elucidated. In this review we summarise the basic structural and functional properties of GH, its receptor and their interaction. In addition, we discuss the discovery and development of an effective GH receptor antagonist, pegvisomant, and summarise potential therapeutic uses of this drug.

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Key words: growth hormone, growth hormone receptor, growth hormone receptor antagonist

#### Streszczenie

Hormon wzrostu (GH, *growth hormone*) uczestniczy w wielu fizjologicznych procesach dotyczących wzrastania, różnicowania i metabolizmu. Leczenie rekombinowanym ludzkim GH jest akceptowane w wielu schorzeniach wiążących się z całkowitym brakiem lub zmniejszeniem działania GH. Wynika stąd znaczne zainteresowanie rozwojem nowych leków mogących modyfikować czynność GH. Dopiero niedawno wyjaśniono dokładną strukturę GH i jego interakcje z receptorem. W niniejszej pracy autorzy podsumowują wiedzę dotyczącą podstawowej budowy GH, jego receptora i interakcji między nimi. Ponadto, omówiono odkrycie i rozwój skutecznego antagonisty receptora GH, pegvisomantu i przedstawiono potencjalne możliwości zastosowania terapeutycznego tego leku.

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Słowa kluczowe: hormon wzrostu, receptor hormonu wzrostu, antagonista receptora hormonu wzrostu

# Introduction

The functions of growth hormone (GH) are pervasive, having a direct or indirect impact on most tissues in the body. To exert its biological effect, GH interacts with specific GH receptors (GHRs) on the surface of target

Prof. John J. Kopchick Ohio University, Athens, OH tissues. GHRs have been detected in a variety of tissues, including liver, adipose tissue, muscle, lymphocytes, prostate, kidney, placenta, heart, brain and mammary gland [1–5]. Binding of GH to GHRs on target tissues activates proteins involved in the Janus kinase/signal transducer and activator of transcription (JAK-STAT) signal transduction pathway, as well as other pathways [6]. In addition to having a direct impact on target tissues, GH stimulates the synthesis and release of insulin-like growth factor-1 (IGF-1). Since IGF-1 has many distinct metabolic effects, GH has the ability to alter tissue function, both directly and indirectly, via IGF-1 production. Thus GH, along with IGF-1, is considered to have dual effects on target tissues [7, 8] with the initiator of this cascade being the interaction of GH with the GHR.

Disorders in growth, either via GH deficiency (GHD) or by production of elevated levels of GH such as in acromegalic individuals, have resulted in a variety of treatment modalities. For deficiency states rhGH has been approved by the FDA for treatment of several growth retardation conditions in children including GHD, Turner syndrome, chronic renal disease, Prader-Willi syndrome and intrauterine growth retardation and for children born small for gestational age (SAGE) or with idiopathic short stature. In adults, rhGH has been approved for GHD associated with a history of hypothalamic and/or pituitary disorders and, more recently, for human immunodeficiency virus (HIV)-associated wasting. The guidelines for rhGH use in children and adults have been thoroughly reviewed elsewhere in more detail [9-11]. Recently, recombinant IGF-1 has been approved for children resistant or insensitive to GH treatment [12]. For conditions of elevated GH (gigantism and acromegaly) drugs that lower GH secretion (somatostatin analogues) or inhibit GH activity (GH antagonists) are currently used. Thus by altering the GH/GHR interaction both GH and IGF-1 activities will be affected. In the future the ability to uncouple the action of GH from that of IGF-1 will certainly result in the discovery of new therapeutic targets.

# GH and GHR structure

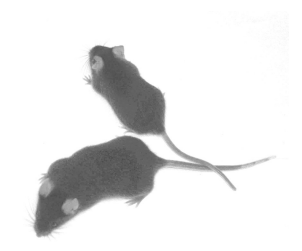
The primary sequence of GH from many species, as well as the crystal structure of both porcine (p) and human (h) GH, has provided significant insight as to the structurally significant regions of this hormone. Although somewhat variable according to the species, the main secreted form of GH is composed of ~191 amino acids. Analysis of the three-dimensional structure of pGH [13] and hGH [14] has revealed that both are globular proteins which contain four highly conserved cysteine residues. These cysteine residues form both a large and a small disulfide bridge with the large bridge being important for GH activity [15]. Approximately one half of the amino acid residues in GH reside in four distinct alpha helices. These four anti-parallel helices connect in an "up-up-down-down" pattern with the core of the four-helical bundle consisting of mostly hydrophobic residues, which presumably function to hold the helices in a specific packed configuration [14]. Relevant to this review, a tryptic peptide of GH containing helix 3 was previously shown to have significant growth-promoting activity [16], although this was not documented to be critical for GHR recognition.

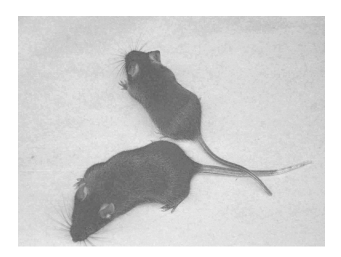
The GHR belongs to the cytokine receptor superfamily, which also includes the receptors for granulocyte-colony stimulating factor, leptin and prolactin as well as other cytokines [17]. There are common features and motifs among this receptor family. In particular, receptor family members contain several specific disulfide bonded Cys residues and a distinct WSXWS-like (Trp, Ser, any amino acid, Trp and Ser) motif near the cell membrane. The GHR is composed of approximately 620 amino acids. The N-terminus contains the extracellular hormone-binding region (~245 amino acids), followed by the 24 amino acid hydrophobic transmembrane region and the C-terminal domain (~350 amino acids), which contains motifs important in intracellular signalling [18]. On the basis of analysis of the crystal structure of human GHR, the extracellular region contains two distinct yet similarly designed domains (termed 1 and 2), each composed of seven beta strands divided into two anti-parallel beta sheets [14].

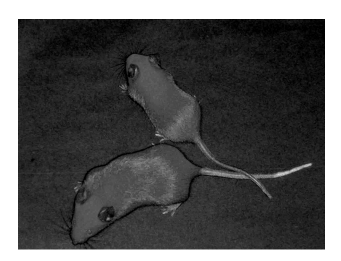
### Interaction of GH with the GHR

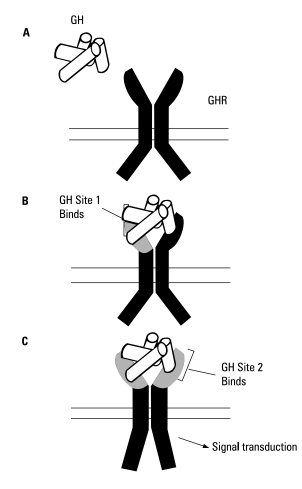
The sensitivity of a tissue to GH is partly dependent on the number of cell-surface GHRs in that tissue. The stoichometry of the ligand:receptor complex is 1:2, based on a number of biophysical methods and later confirmed through X-ray crystallography data [14, 19]. Thus a single GH molecule interacts with a homodimer of the GHR. Several studies have revealed that GHR exists as a preformed homodimer [20–22], which undergoes a conformational change in the intracellular signalling region of the receptor, initiating the signalling cascade [23]. This account of the heterotrimeric GH:GHR interaction is depicted in Figure 1.

The regions of GH responsible for receptor binding have been exhaustively studied [24-29]. These studies identified a patch of three regions of GH that come into close proximity in the three-dimensional structure, which are responsible for the high affinity binding of GH to its receptor [24, 25]. These three regions of GH, collectively referred to as Site 1, include the N-terminal portion of helix 1, a portion of the connection between helices 1 and 2, and the C terminal portion of helix 4. Yet, GH forms a hGH:GHR2 complex and is an asymmetric protein, suggesting that an additional site within GH (first suggested by Chen et al [30]) was responsible for binding the second GHR monomer. This additional site was later found in helix 3 of GH and is called Site 2. Because two physically separate sites of a single GH protein are responsible for binding to the GHR, it may not be surprising that Site 1 of GH interacts with higher affinity to GHR than Site 2 [14]. Described sequentially, Site 1 of GH is thought to interact with higher









**Figure 1.** Model for the formation and signal transduction of the heterotrimeric complex between GH and 2GHR. **A**. The preformed dimer of GHR is shown embedded in the lipid bilayer and GH is in the extracellular space. **B**. Site 1 (within GH) binds with high affinity to a monomer of the preformed GHR dimer. **C**. Site 2 (within GH) subsequently binds the second GHR, resulting in signal transduction. (Reprinted with permission from Cold Spring Laboratory Press [91]

Rycina 1. Schemat powstawania i przekazywania sygnału w obrębie kompleksu heterotrimerycznego między cząsteczkami GH i 2GHR. A. Wbudowany w dwuwarstwę lipidową preformowany dimer GHR; GH znajduje się w przestrzeni zewnątrzkomórkowej. B. Miejsce 1. (w obrębie GH) wiąże się z dużym powinowactwem do jednego z monomerów proformowanego dimeru GHR. C. Miejsce 2. (w obrębie GH) wiąże się następnie z drugim monomerem dimeru GHR, co powoduje przewodzenie sygnału

affinity to the first GHR monomer, followed by the binding of Site 2 of GH to the second GHR monomer with lower affinity.

#### Discovery of the GH antagonist

Detailed focus was placed on the  $3^{rd} \alpha$ -helix of GH due to the growth-promoting abilities [16, 30, 31] observed to be specific to that helix. Mutations at 3 amino acid



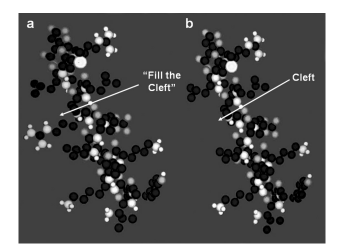
Darlene E. Berryman et al.

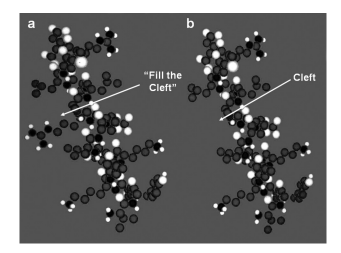
**Figure 2.** *Photograph showing size comparison of control (bottom) and GH antagonist transgenic mice (top). Shown are 6-week old male mice* 

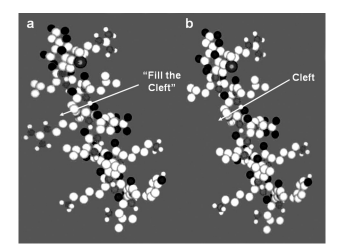
**Rycina 2.** Na fotografii przedstawiono różnice w wielkości między kontrolną (u dołu) i transgeniczną myszą po zadziałaniu antagonistów GH (u góry). Przedstawione myszy to 6-tygodniowe osobniki męskie

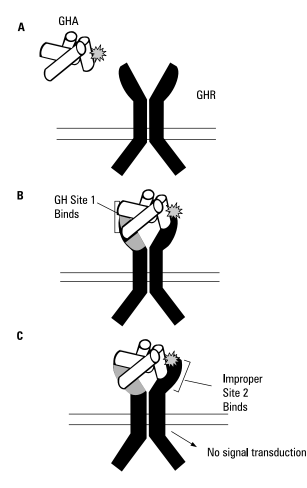
sites within bovine (b) GH helix 3 were engineered to provide an amphipathic formation hypothesised to further enhance the growth-promoting activity of this helix. Specifically, the substitutions were Glu-117 to Leu, Gly-119 to Arg, and Ala-122 to Asp in bGH. This GH analogue bound to the GHR's with the same affinity as wild-type GH [30]. Surprisingly, this GH analogue antagonised the action of wild-type GH in transgenic mice, resulting in a dwarf phenotype [30-32]. This result represents the first discovery of a GH antagonist. Further investigation of each individual substitution revealed that the specific replacement of Gly-119 with Arg promoted the GH antagonist effect [29]. This single Gly-119 amino acid substitution is sufficient to promote a dwarf phenotype in mice transgenic for the GH antagonist (Fig. 2) [31].

Interestingly, the Gly at this position is conserved in all members of the GH family. The GH antagonist is able to bind with high affinity to the preformed GHR dimer while blocking subsequent signal transduction (Fig. 3) [30, 31, 33]. Gly's side chain is made up of a single hydrogen atom, which, in the context of other amino acids in the vicinity, creates a cleft in a region of the 3<sup>rd</sup> helix (Fig. 4). The substitution of this Gly with an amino acid containing a bulky side chain fills this gap, which ultimately generates the GH antagonist [29]. It is important to note that these types of GH antagonist bind to the GHR with affinities similar to wild-type GH and do not inhibit GHR dimerisation but perturb proper or functional GHR dimerisation.







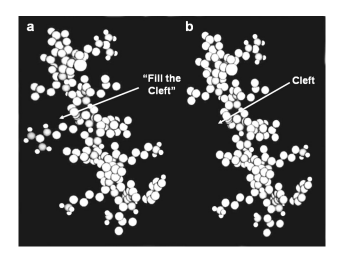


**Figure 3.** Model representing the interaction of the GH antagonist with the receptor. **A.** A preformed dimer of GHR is shown embedded in the lipid bilayer with the GH antagonist in the extracellular space. **B.** Site 1 within the GH antagonist binds with high affinity to one monomer of the preformed GHR dimer. **C.** Improper binding at Site 2 within the GH antagonist blocks subsequent intracellular signal transduction. (Reprinted with permission from Cold Spring Harbor Laboratory Press [91])

**Rycina 3.** Na schemacie przedstawiono oddziaływanie między anta-gonistą GH a receptorem. **A.** Wbudowany w dwuwarstwę lipidową preformowany dimer GHR; antagonista GH znajduje się w przestrzeni zewnątrzkomórkowej. **B.** Miejsce 1. (w obrębie GH) wiąże się z dużym powinowactwem do jednego z monomerów proformowanego dimeru GHR. **C.** Nieprawidłowe wiązanie miejsca 2. (w obrębie cząsteczki antagonisty GH) do drugiego monomeru GHR blokuje późniejsze wewnątrzkomórkowe przewodzenie sygnału

# Development of a long-acting, effective GH antagonist

Owing to GH's relatively short half life (30 minutes), it has proved challenging to create a GH antagonist molecule that was an effective therapeutic agent. In order to counteract kidney excretion of low molecular weight GH, the addition of polyethylene glycol (PEG) was used to significantly increase the molecular mass of the



**Figure 4.** Partial space filling model of the third alpha helix of bGH and bGH-G119R. **A.** Structural representation of the third alpha helix when the glycine is substituted with an arginine. **B.** Structural representation of the wild-type helix. The position of the cleft is indicated. (Reprinted, with permission, (38) [Ó The Endocrine Society])

**Rycina 4.** Częściowy model przestrzenny trzeciej alfa helisy bGH i bGH-G119R. A. Struktura trzeciej alfa helisy po podstawieniu glicyny argininą. B. Struktura helisy typu dzikiego. Zaznaczono pozycję szczeliny

protein [34]. This technology was adapted for the GH antagonist. The PEG addition decreased the affinity of the GH antagonist for its receptor but still proved an effective antagonist because the serum half-life was improved [21]. Furthermore, in an attempt to improve the affinity of the pegylated GH antagonist for its receptor, 8 amino acid substitutions were generated at Site 1, each of which had previously been shown to improve the affinity for GH binding protein [35]. This 8 amino acid substituted and pegylated antagonist (containing lysine at Gly 120) had improved binding affinity for membrane receptors as compared to the pegylated Gly 120K antagonist, resulting in a more effective molecule [21].

This pegylated GH antagonist has been termed pegvisomant and the approved marketed name is Somavert ® (pegvisomant for injection). Many papers have documented the clinical efficacy of pegvisomant and these will not be further reviewed here. However, readers should visit the following papers and reviews for specific details concerning the many clinical trials [36–45].

## **Pegvisomant and diabetes**

Although it has been known for many decades that GH inhibits insulin's action[46–49], the mechanism responsible for this effect has remained elusive. Recent data have started to illuminate possible mechanisms. For example, a recent link between a specific GHR poly-

morphism and resistance to Type 2 diabetes (T2DM) has been presented [50]. Furthermore, mice transgenic for bGH are insulin resistant, while mice that lack GH signalling are insulin sensitive despite their obesity [51–53]. In terms of intracellular signalling events that account for GH-induced insulin resistance, disruption of p85alpha, a subunit of PI 3-kinase, will increase insulin sensitivity, while elevated p85alpha levels are associated with insulin resistance [54-57]. A recent study by del Rincon et al. reports that GH up-regulates expression of p85alpha in white adipose tissue and suggests this may be responsible for alterations in insulin sensitivity seen in mouse models of altered GH action [58]. A similar situation also occurs in muscle [56]. Thus the diabetogenic effect of GH may be due to "cross-talk" between the GH and insulin signalling pathways.

The established impact of GH on insulin sensitivity led researchers to monitor parameters of insulin action in human subjects given pegvisomant. Healthy subjects given pegvisomant for 7 days did not show altered glucose tolerance or stimulated insulin secretion [59]. As pegvisomant began to be used to treat acromegaly, a disease often accompanied by insulin resistance and diabetes, clinicians were able to examine the effect of this drug on insulin sensitivity and other measures of diabetes. In 2002 Rose and Clemmons reported that treatment with pegvisomant lowered fasting insulin, glucose and haemoglobin A(1)C levels in patients with acromegaly [60]. Later studies have further confirmed an improvement in insulin sensitivity following pegvisomant treatment of patients with acromegaly [61–63].

Clearly, pegvisomant can improve insulin and glucose levels in patients with acromegaly, but what about patients with other insulin-related conditions? Williams et al. treated young, Type 1 diabetic adults with 5 or 10 mg/day of pegvisomant for 3 weeks [64]. No changes in insulin sensitivity under hyperinsulinaemic euglycaemic clamp conditions were observed; however, both doses of pegvisomant decreased the amount of insulin required overnight to maintain euglycaemia. Thus although there has been limited research to date, pegvisomant shows promise for treating not only acromegalics with insulin resistance but also young adult patients with Type 1 diabetes. Further research is required to determine if pegvisomant treatment might benefit patients with type 2 diabetes as well.

#### Pegvisomant and nephropathy

Long and short term renal changes can be caused by GH and IGF-1. Transgenic mice expressing GH antagonist are dwarf and have reduced circulating IGF-1 levels [30, 32]. When GH antagonist mice are made diabetic, they are protected from renal damage [65]. In

addition, treatment of control and diabetic mice with GH antagonist protects them from renal damage [66, 67] and prevents compensatory renal growth in uni-nephrectomised mice [68]. The mechanism in which GH antagonist protects the kidney has not been determined, but studies point to several possibilities. When exogenous GH antagonist is administered in increasing doses to adult female Balb/C mice, there is a dose-dependent decrease in hepatic and serum IGF-1 levels, no effect on hepatic or renal IGFBP-1 and 3 levels, and an increase in hepatic and circulatory IGFBP-4 levels [69]. In effect, this would create a significant decrease in IGF-1 bioavailability. Additionally, variable concentrations of pegvisomant have a significant impact on the GHR/GHRBP gene transcription in stable cell lines of T-SV40 immortalised glomerular mesangial cells [70, 71]. Interestingly, GH antagonist has been reported to inhibit GHR/GHRBP gene transcription directly at the cellular level in human mesangial cells at all concentrations of pegvisomant tested [72]. Collectively, this data indicates that pegvisomant administration may influence kidney function.

#### Pegvisomant and retinopathy

The role of GH in the development of retinopathy was first described after ablation of the pituitary gland resulted in reduction of the disease [73, 74]. This result, coupled with the fact that diabetic dwarfs do not develop retinopathy [75], suggests that the use of GH antagonists for the treatment of diabetic retinopathy may be beneficial. Furthermore, results using mice expressing a GH antagonist to study non-diabetic ischemiainduced retinal neovascularisation showed an inhibition of neovascularisation despite elevated levels of vascular endothelial growth factor receptor [76]. Pegvisomant treatment of diabetic patients with severe retinopathy ensued. In this 12-week study, where type 1 and type 2 diabetic patients were treated daily with pegvisomant, no regression of retinopathy was seen [77]. However, considering the short length of the study as well as the advanced retinopathy of the subjects, further studies are warranted.

#### Pegvisomant and cardiovascular disease

Acromegaly has been shown to be associated with an increased cardiovascular risk. Thus it is not surprising that CRP (C-reactive protein) levels, a common marker for cardiovascular risk, were found to be lower with the administration of pegvisomant in humans [78]. Since pegvisomant blocks GHR activation and decreases IGF-1 production, the effects observed on CRP could be mediated both by the decrease in IGF-1 and the direct effect of GHR blockade. GH antagonist treatment in patients with acromegaly is also known to induce a reduction in diastolic blood pressure in hypertension and improve glucose metabolism [62]. A recent study by Pivonello et al. also showed that pegvisomant can reverse left ventricle hypertrophy and progressively improve left ventricular diastolic and systolic performance in acromegalics [79]. Thus long term treatment with pegvisomant has positive effects on cardiovascular function and may prevent the development or progression of cardiac insufficiency, at least for acromegalics.

# Pegvisomant and cancer

The IGF-1/GH axis has been implicated in contributing to the growth and formation of many different cancers [80, 81]. IGF-1 has been shown to be a growth factor for numerous types of cancer and neoplastic growth [82]. Additional studies have also shown that some neoplasms are also capable of producing autocrine and/or paracrine IGF-1 [82]. Transfection of MCF-7 cells with the hGH gene showed that these cells synthesised and secreted hGH into the media, and these cells were found to have higher levels of STAT5-mediated transcriptional activation than controls [83]. The disruption of excess GH stimulation and therefore reduction in IGF-1 levels may therefore be useful in the treatment of numerous cancers.

Multiple studies using both animals and humans have attempted to show the beneficial use of GH antagonist to prevent or slow the growth of various tumours. GH antagonist mice were found to have lower IGF-1 levels and a decreased mammary tumour incidence in relation to litter-mate controls when exposed to a chemical carcinogen [84]. Additional studies using GHR/-and C3(1)/Tag mouse models showed an inhibition of oestrogen-independent mammary carcinogenesis [85]. Recently a study using the spontaneous dwarf rat (an animal known to have lower levels of GH and IGF-1) injected with GH showed that these animals were more vulnerable to mammary carcinogenesis with increasing levels of circulating IGF-1 and GH [86]. Pegvisomant administration to virgin female mice caused a 70-80% reduction in serum IGF-1 levels and a 30% reduction in the volume of MCF-7 xenografts [87]. In mice the growth of human meningioma xenografts significantly decreased following pegvisomant treatment, and in some cases tumour regression was observed [88]. Additional studies xenografting human colorectal cancer lines into female nude mice with subsequent pegvisomant treatment reported a 39% reduction in tumour volume with a reduction in both IGF-1 and IGFBP-3 levels [89]. Studies involving GHR disrupted and Tag mice suggest that the disruption of GH signalling may also reduce prostate carcinogenesis [90]. These results indicate a potential therapeutic use of pegvisomant in the prevention and treatment of certain cancers.

# Conclusion

Since the initial discovery of a growth hormone antagonist [30] both basic and clinical studies have advanced. In terms of human use, the growth hormone antagonist Somavert® (pegvisomant for injection) has been approved for lowering IGF-1 levels in acromegalic individuals. Further studies are likely to provide insight into its therapeutic potential for the treatment of diabetes, diabetic complications and cancer indications. Finally, the growth hormone antagonist is now a commonly used reagent that specifically antagonises the effects of growth hormone in many basic research scenarios. In the future, the growth hormone antagonist will also assist researchers in uncoupling the biological effects of growth hormone from those of IGF-1.

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#### References

- 1. Hill DJ, Riley SC, Bassett NS et al. Localization of the growth hormone receptor, identified by immunocytochemistry, in second trimester human fetal tissues and in placenta throughout gestation. J Clin Endocrinol Metab 1992; 75 (2): 646–650.
- Mercado M, Da Vila N, McLeod JF et al. Distribution of growth hormone receptor messenger ribonucleic acid containing and lacking exon 3 in human tissues. J Clin Endocrinol Metab 1994; 78 (3): 731–735.
- 3. Mertani HC, Morel G. In situ gene expression of growth hormone (GH) receptor and GH binding protein in adult male rat tissues. Mol Cell Endocrinol 1995; 109 (1): 47–61.
- Nyberg F, Burman P. Growth hormone and its receptors in the central nervous system — location and functional significance. Horm Res 1996; 45 (1–2): 18–22.
- Rapaport R, Sills IN, Green L et al. Detection of human growth hormone receptors on IM-9 cells and peripheral blood mononuclear cell subsets by flow cytometry: correlation with growth hormone-binding protein levels. J Clin Endocrinol Metab 1995; 80 (9): 2612–2619.
- Kopchick JJ, Andry JM. Growth hormone (GH), GH receptor, and signal transduction. Mol Genet Metab 2000; 71 (1–2): 293–314.
- Goodman HM. Multiple effects of growth hormone on lipolysis. Endocrinology 1968; 83 (2): 300–308.
- Green H, Morikawa M, Nixon T. A dual effector theory of growth-hormone action. Differentiation 1985; 29 (3): 195–198.
- 9. Gharib H, Cook DM, Saenger PH et al. American Association of Clinical Endocrinologists medical guidelines for clinical prac-

tice for growth hormone use in adults and children — 2003 update. Endocr Pract 2003; 9 (1): 64–76.

- Molitch ME, Clemmons DR, Malozowski S et al. Evaluation and treatment of adult growth hormone deficiency: an Endocrine Society Clinical Practice Guideline. J Clin Endocrinol Metab 2006; 91 (5): 1621–1634.
- 11. Clayton PE, Cianfarani S, Czernichow P et al. Management of the child born small for gestational age through to adulthood: a consensus statement of the International Societies of Pediatric Endocrinology and the Growth Hormone Research Society. J Clin Endocrinol Metab 2007; 92 (3): 804–810.
- 12. INCRELEX<sup>™</sup> Package Insert. 2005.
- Abdel-Meguid SS, Shieh HS, Smith WW et al. Three-dimensional structure of a genetically engineered variant of porcine growth hormone. Proc Natl Acad Sci USA 1987; 84 (18): 6434–6437.
- de Vos AM, Ultsch M, Kossiakoff AA. Human growth hormone and extracellular domain of its receptor: crystal structure of the complex. Science 1992; 255 (5042): 306–312.
- Chen XZ, Shafer AW, Yun JS et al. Conversion of bovine growth hormone cysteine residues to serine affects secretion by cultured cells and growth rates in transgenic mice. Mol Endocrinol 1992; 6 (4): 598–606.
- Hara K, Hsu Chen CJ, Sonenberg M. Recombination of the biologically active peptides from a tryptic digest of bovine growth hormone. Biochemistry 1978; 17 (3): 550–556.
- Bazan JF. Haemopoietic receptors and helical cytokines. Immunol Today 1990; 11 (10): 350–354.
- Leung DW, Spencer SA, Cachianes G et al. Growth hormone receptor and serum binding protein: purification, cloning and expression. Nature 1987; 330 (6148): 537–543.
- Cunningham BC, Ultsch M, De Vos AM et al. Dimerization of the extracellular domain of the human growth hormone receptor by a single hormone molecule. Science 1991; 254 (5033): 821–825.
- 20. Harding PA, Wang X, Okada S et al. Growth hormone (GH) and a GH antagonist promote GH receptor dimerization and internalization. J Biol Chem 1996; 271 (12): 6708–6712.
- Ross RJ, Leung KC, Maamra M et al. Binding and functional studies with the growth hormone receptor antagonist, B2036-PEG (pegvisomant), reveal effects of pegylation and evidence that it binds to a receptor dimer. J Clin Endocrinol Metab 2001; 86 (4): 1716–1723.
- Gent J, van Kerkhof P, Roza M et al. Ligand-independent growth hormone receptor dimerization occurs in the endoplasmic reticulum and is required for ubiquitin system-dependent endocytosis. Proc Natl Acad Sci USA 2002; 99 (15): 9858–9863.
- Brown RJ, Adams JJ, Pelekanos RA et al. Model for growth hormone receptor activation based on subunit rotation within a receptor dimer. Nat Struct Mol Biol 2005; 12 (9): 814–821.
- 24. Cunningham BC, Jhurani P, Ng P et al. Receptor and antibody epitopes in human growth hormone identified by homologscanning mutagenesis. Science 1989; 243 (4896): 1330–1336.
- Cunningham BC, Wells JA. High-resolution epitope mapping of hGH-receptor interactions by alanine-scanning mutagenesis. Science 1989; 244 (4908): 1081–1085.
- Pearce KH Jr, Ultsch MH, Kelley RF et al. Structural and mutational analysis of affinity-inert contact residues at the growth hormone-receptor interface. Biochemistry 1996; 35 (32): 10 300– -10 307.
- Fuh G, Cunningham BC, Fukunaga R et al. Rational design of potent antagonists to the human growth hormone receptor. Science 1992; 256 (5064): 1677–1680.
- Bass SH, Mulkerrin MG, Wells JA. A systematic mutational analysis of hormone-binding determinants in the human growth hormone receptor. Proc Natl Acad Sci USA 1991; 88 (10): 4498– -4502.
- 29. Chen WY, Wight DC, Chen NY et al. Mutations in the third alpha-helix of bovine growth hormone dramatically affect its

intracellular distribution in vitro and growth enhancement in transgenic mice. J Biol Chem 1991; 266 (4): 2252–2258.

- Chen WY, Wight DC, Wagner TE et al. Expression of a mutated bovine growth hormone gene suppresses growth of transgenic mice. Proc Natl Acad Sci USA 1990; 87 (13): 5061–5065.
- Chen WY, Wight DC, Mehta BV et al. Glycine 119 of bovine growth hormone is critical for growth-promoting activity. Mol Endocrinol 1991; 5 (12): 1845–1852.
- 32. Chen WY, White ME, Wagner TE et al. Functional antagonism between endogenous mouse growth hormone (GH) and a GH analog results in dwarf transgenic mice. Endocrinology 1991; 129 (3): 1402–1408.
- Harding PA, Wang X, Okada S et al. Growth hormone (GH) and a GH antagonist promote GH receptor dimerization and internalization. J Biol Chem 1996; 271 (12): 6708–6712.
- Clark R, Olson K, Fuh G et al. Long-acting growth hormones produced by conjugation with polyethylene glycol. J Biol Chem 1996; 271 (36): 21969–21977.
- Lowman HB, Wells JA. Affinity maturation of human growth hormone by monovalent phage display. J Mol Biol 1993; 234 (3): 564–578.
- Trainer PJ, Drake WM, Katznelson L et al. Treatment of acromegaly with the growth hormone-receptor antagonist pegvisomant. N Engl J Med 2000; 342 (16): 1171–1177.
- van der Lely AJ, Hutson RK, Trainer PJ et al. Long-term treatment of acromegaly with pegvisomant, a growth hormone receptor antagonist. Lancet 2001; 358 (9295): 1754–1759.
- Kopchick JJ, Parkinson C, Stevens EC et al. Growth hormone receptor antagonists: discovery, development, and use in patients with acromegaly. Endocr Rev 2002; 23 (5): 623–646.
- 39. Kopchick JJ. Discovery and mechanism of action of pegvisomant. Eur J Endocrinol 2003; 148 (supl 2): S21–S25.
- Muller AF, van der Lely AJ. Pharmacological therapy for acromegaly: a critical review. Drugs 2004; 64 (16): 1817–1838.
- 41. Paisley AN, Trainer PJ, Drake WM. The place of pegvisomant in the acromegaly treatment algorithm. Growth Horm IGF Res 2004; 14 (supl A): S101–S106.
- 42. van der Lely AJ, Kopchick JJ. Growth hormone receptor antagonists. Neuroendocrinology 2006; 83 (3–4): 264–268.
- 43. Paisley AN, Drake WM. Treatment of pituitary tumors: pegvisomant. Endocrine 2005; 28 (1): 111–114.
- Paisley AN, Trainer PJ. Recent developments in the therapy of acromegaly. Expert Opin Investig Drugs 2006; 15 (3): 251–256.
- Burt MG, Ho KK. Newer options in the management of acromegaly. Intern Med J 2006; 36 (7): 437–444.
- Houssay B, Biasotti A. The hypothesis, carbohydrate metabolism and diabetes. Endocrinology 1931; 15: 511.
- 47. Hansen I, Tsalikian E, Beaufrere B et al. Insulin resistance in acromegaly: defects in both hepatic and extrahepatic insulin action. Am J Physiol 1986; 250 (3 Pt 1): E269–E273.
- Davidson MB. Effect of growth hormone on carbohydrate and lipid metabolism. Endocr Rev 1987; 8 (2): 115–131.
- Foss MC, Saad MJ, Paccola GM et al. Peripheral glucose metabolism in acromegaly. J Clin Endocrinol Metab 1991; 72 (5): 1048–1053.
- 50. Strawbridge RJ, Karvestedt L, Li C et al. GHR exon 3 polymorphism: association with type 2 diabetes mellitus and metabolic disorder. Growth Horm IGF Res 2007.
- Coschigano KT, Clemmons D, Bellush LL et al. Assessment of growth parameters and life span of GHR/BP gene-disrupted mice. Endocrinology 2000; 141 (7): 2608–2613.
- Berryman DE, List EO, Coschigano KT et al. Comparing adiposity profiles in three mouse models with altered GH signaling. Growth Horm IGF Res 2004; 14 (4): 309–318.
- 53. Olsson B, Bohlooly YM, Fitzgerald SM et al. Bovine growth hormone transgenic mice are resistant to diet-induced obesity but develop hyperphagia, dyslipidemia, and diabetes on a highfat diet. Endocrinology 2005; 146 (2): 920–930.
- 54. Mauvais-Jarvis F, Ueki K, Fruman DA et al. Reduced expression of the murine p85alpha subunit of phosphoinositide

3-kinase improves insulin signaling and ameliorates diabetes. J Clin Invest 2002; 109 (1): 141–149.

- Cornier MA, Bessesen DH, Gurevich I et al. Nutritional upregulation of p85alpha expression is an early molecular manifestation of insulin resistance. Diabetologia 2006; 49 (4): 748–754.
- 56. Barbour LA, Mizanoor Rahman S, Gurevich I et al. Increased P85alpha is a potent negative regulator of skeletal muscle insulin signaling and induces in vivo insulin resistance associated with growth hormone excess. J Biol Chem 2005; 280 (45): 37 489–37 494.
- Taniguchi CM, Tran TT, Kondo T et al. Phosphoinositide 3-kinase regulatory subunit p85alpha suppresses insulin action via positive regulation of PTEN. Proc Natl Acad Sci USA 2006; 103 (32): 12 093–12 097.
- del Rincon JP, Iida K, Gaylinn BD et al. Growth hormone regulation of p85alpha expression and phosphoinositide 3-kinase activity in adipose tissue: mechanism for growth hormonemediated insulin resistance. Diabetes 2007; 56 (6): 1638–1646.
- Parkinson C, Drake WM, Roberts ME et al. A comparison of the effects of pegvisomant and octreotide on glucose, insulin, gastrin, cholecystokinin, and pancreatic polypeptide responses to oral glucose and a standard mixed meal. J Clin Endocrinol Metab 2002; 87 (4): 1797–1804.
- Rose DR, Clemmons DR. Growth hormone receptor antagonist improves insulin resistance in acromegaly. Growth Horm IGF Res 2002; 12 (6): 418–424.
- 61. Drake WM, Rowles SV, Roberts ME et al. Insulin sensitivity and glucose tolerance improve in patients with acromegaly converted from depot octreotide to pegvisomant. Eur J Endocrinol 2003; 149 (6): 521–547.
- 62. Colao A, Pivonello R, Auriemma RS et al. Efficacy of 12-month treatment with the GH receptor antagonist pegvisomant in patients with acromegaly resistant to long-term, high-dose somatostatin analog treatment: effect on IGF-I levels, tumor mass, hypertension and glucose tolerance. Eur J Endocrinol 2006; 154 (3): 467–477.
- 63. Lindberg-Larsen R, Moller N, Schmitz O et al. The impact of pegvisomant treatment on substrate metabolism and insulin sensitivity in patients with acromegaly. J Clin Endocrinol Metab 2007; 92 (5): 1724–1728.
- 64. Williams RM, Amin R, Shojaee-Moradie F et al. The effects of a specific growth hormone antagonist on overnight insulin requirements and insulin sensitivity in young adults with Type 1 diabetes mellitus. Diabetologia 2003; 46 (9): 1203–1210.
- Chen NY, Chen WY, Kopchick JJ. A growth hormone antagonist protects mice against streptozotocin induced glomerulosclerosis even in the presence of elevated levels of glucose and glycated hemoglobin. Endocrinology 1996; 137 (11): 5163–5165.
- 66. Flyvbjerg A, Bennett WF, Rasch R et al. Inhibitory effect of a growth hormone receptor antagonist (G120K-PEG) on renal enlargement, glomerular hypertrophy and urinary albumin excretion in experimental diabetes in mice. Diabetes 1999; 48: 377–382.
- 67. Segev Y, Landau D, Rasch R et al. Growth hormone receptor antagonism prevents early renal changes in nonobese diabetic mice. J Am Soc Nephrol 1999; 10 (11): 2374–2381.
- Flyvbjerg A, Bennett WF, Rasch R et al. Compensatory renal growth in uninephrectomized adult mice is growth hormone dependent. Kidney Int 1999; 56 (6): 2048–2054.
- 69. van Neck JW, Dits NF, Cingel V et al. Dose-response effects of a new growth hormone receptor antagonist (B2036-PEG) on circulating, hepatic and renal expression of the growth hormone/insulin-like growth factor system in adult mice. J Endocrinol 2000; 167 (2): 295–303.
- Sraer JD, Delarue F, Hagege J et al. Stable cell lines of T-SV40 immortalized human glomerular mesangial cells. Kidney Int 1996; 49 (1): 267–270.
- 71. Mene P, Simonson MS, Dunn MJ. Physiology of the mesangial cell. Physiol Rev 1989; 69 (4): 1347–1424.

- Meinhardt U, Eble A, Besson A et al. Regulation of growthhormone-receptor gene expression by growth hormone and pegvisomant in human mesangial cells. Kidney Int 2003; 64 (2): 421–430.
- 73. Poulsen J. The Houssay phenomenon in man: recovery from retinopathy in a case of diabetes with Simmond's disease. Diabetes 1953; 2: 7–12.
- 74. Poulsen JE. Diabetes and anterior pituitary insufficiency. Final course and postmortem study of a diabetic patient with Sheehan's syndrome. Diabetes 1966; 15 (2): 73–77.
- Merimee TJ. A follow-up study of vascular disease in growthhormone-deficient dwarfs with diabetes. N Engl J Med 1978; 298 (22): 1217–1222.
- Smith LE, Kopchick JJ, Chen W et al. Essential role of growth hormone in ischemia-induced retinal neovascularization. Science 1997; 276 (5319): 1706–1709.
- 77. The Growth Hormone Antagonist for Proliferate Diabetic Retinopathy Study Group. The effect of a growth hormone receptor antagonist drug on proliferative diabetic retinopathy. Ophthalmology 2001; 108 (12): 2266–2272.
- Sesmilo G, Fairfield WP, Katznelson L et al. Cardiovascular risk factors in acromegaly before and after normalization of serum IGF-I levels with the GH antagonist pegvisomant. J Clin Endocrinol Metab 2002; 87 (4): 1692–1699.
- 79. Pivonello R, Galderisi M, Auriemma RS et al. Treatment with growth hormone receptor antagonist in acromegaly: effect on cardiac structure and performance. J Clin Endocrinol Metab 2007; 92 (2): 476–482.
- 80. Pollak M. Insulin-like growth factor physiology and cancer risk. Eur J Cancer 2000; 36 (10): 1224–1228.
- Pollak MN, Schernhammer ES, Hankinson SE. Insulin-like growth factors and neoplasia. Nat Rev Cancer 2004; 4 (7): 505–518.
- Khandwala HM, McCutcheon IE, Flyvbjerg A et al. The effects of insulin-like growth factors on tumorigenesis and neoplastic growth. Endocr Rev 2000; 21 (3): 215–244.
- Kaulsay KK, Mertani HC, Tornell J et al. Autocrine stimulation of human mammary carcinoma cell proliferation by human growth hormone. Exp Cell Res 1999; 250 (1): 35–50.
- Pollak M, Blouin MJ, Zhang JC et al. Reduced mammary gland carcinogenesis in transgenic mice expressing a growth hormone antagonist. Br J Cancer 2001; 85 (3): 428–430.
- 85. Zhang X, Mehta RG, Lantvit DD et al. Inhibition of estrogenindependent mammary carcinogenesis by disruption of growth hormone signaling. Carcinogenesis 2007; 28 (1): 143–150.
- 86. Shen Q, Lantvit DD, Lin Q et al. Advanced rat mammary cancers are growth hormone dependent. Endocrinology 2007; epub ahead of press.
- Divisova J, Kuiatse I, Lazard Z et al. The growth hormone receptor antagonist pegvisomant blocks both mammary gland development and MCF-7 breast cancer xenograft growth. Breast Cancer Res Treat 2006; 98 (3): 315–327.
- McCutcheon IE, Flyvbjerg A, Hill H et al. Antitumor activity of the growth hormone receptor antagonist pegvisomant against human meningiomas in nude mice. J Neurosurg 2001; 94 (3): 487–492.
- Dagnaes-Hansen F, Duan H, Rasmussen LM et al. Growth hormone receptor antagonist administration inhibits growth of human colorectal carcinoma in nude mice. Anticancer Research 2004; 24 (6): 3735–3742.
- Wang Z, Prins GS, Coschigano KT et al. Disruption of growth hormone signaling retards early stages of prostate carcinogenesis in the C3(1)/T antigen mouse. Endocrinology 2005; 146 (12): 5188–5196.
- Berryman DE, Kopchick JJ. Discovery of a growth hormone antagonist using a structure-function approach. In: Golemis EA, Adams PD (ed). Protein-Protein Interactions: A Molecular Cloning Manual. Cold Spring Harbor: Cold Spring Harbor Laboratory Press. 2005; 873–884.