Identification of a novel zinc finger protein binding a conserved element critical for Pit-1-dependent growth hormone gene expression

Steven M. Lipkin,1,2 Anders M. Näär,1,3 Kristin A. Kalla,4 Ram A. Sack,1,4 and Michael G. Rosenfeld1,4

1Howard Hughes Medical Institute, 1Eukaryotic Regulatory Biology Program, 2Graduate Program in Neurosciences, 3Graduate Program in Molecular Pathology, University of California, San Diego, School and Department of Medicine, La Jolla, California 92093-0648 USA

The growth hormone (GH) and prolactin genes require the pituitary-specific POU domain transcription factor Pit-1 for their activation. However, additional factors are necessary for the effective expression of these genes. Analysis of evolutionarily conserved sequences in the proximal GH promoter suggests the critical importance of one highly conserved element located between the two Pit-1 response elements. Mutation of this site decreases expression of a transgene in mice >100-fold. We have identified a major activity binding to this site as a novel member of the Cys/His zinc finger superfamily, referred to as Zn-15. The Zn-15 DNA-binding domain comprises three zinc fingers separated by unusually long linker sequences that would be expected to interrupt specific DNA site recognition. Zn-15 synergizes with Pit-1 to activate the GH promoter in heterologous cell lines in which this promoter is only minimally responsive to Pit-1 alone. Our data suggest that functional interactions between the tissue-specific POU domain factor Pit-1 and this novel zinc finger factor binding to an evolutionarily conserved region in the GH promoter may constitute an important component of the combinatorial code that underlies the effective expression of the GH gene.

[Key Words: Pit-1; growth hormone; zinc finger; pituitary; transgenic]

Received May 3, 1993; revised version accepted June 24, 1993.

Defining the molecular mechanisms by which select combinations of transcriptional regulators cooperate to regulate tissue-specific gene expression in discrete cell types is a central issue in mammalian development. The growth hormone (GH) gene, expressed solely in the somatotroph cell type in the anterior pituitary, has been studied extensively as a model of tissue-specific gene expression (Voss and Rosenfeld 1992). The proximal GH promoter is capable of targeting somatotroph-specific cis-active elements (Nelson et al. 1986; Bodner and Karin 1987; Nelson et al. 1988; Sharp and Cao 1990). The characterization of a protein that binds to and activates these cell-specific cis-active elements permitted the cloning of Pit-1 (Ingraham et al. 1988; Bodner et al. 1988), a POU domain transcription factor (He et al. 1989) expressed solely in pituitary somatotrophs, lactotrophs, and thyrotrophs. During rat embryogenesis, Pit-1 mRNA is initially detected on e15, 24–36 hr prior to GH and PRL gene activation (Simmons et al. 1990, Dolle et al. 1990). The critical importance of Pit-1 for the survival of somatotrophs, lactotrophs, and thyrotrophs, and for GH gene expression, is illustrated by Snell and Jackson dwarf mice, which possess Pit-1 genomic mutations and lack these cell types in the mature animal (Li et al. 1990). Furthermore, the physiological importance of Pit-1 for GH and PRL transcription has been demonstrated by the identification of human dwarfs who are deficient in both PRL and GH and possess either single-amino-acid substitutions in the Pit-1 POU domain or the introduction of a premature stop codon (Ohta et al. 1992; Pfaflle et al. 1992; Radovick et al. 1992; Tatsumi et al. 1992).

However, the activation of GH, PRL, and TSHβ target genes by Pit-1, and the expression of each in distinct cell types, requires the coordinate actions of factors additional to Pit-1. For example, in transient transfection, the prolactin distal enhancer is effectively stimulated by Pit-1 only in combination with the ligand-activated estrogen receptor (Day et al. 1990; Simmons et al. 1990). For the rat GH gene, T₃ has been suggested to be an important positive regulator. In pituitary tumor cells, T₃ causes a dramatic increase in newly initiated rat GH mRNA transcripts (Evans et al. 1982). The identification of a T₃ response element (T₃RE) in the rat GH promoter suggests that T₃ mediates its effects through the binding of T₃ receptors to this promoter (Glass et al. 1987; Brent
et al. 1989]. However, the sequence of the T₃RE is not conserved in the human GH promoter; in transfection, T₃ negatively regulates human GH [Zhang et al. 1992], suggesting that it may not be a required component of GH activation.

Mapping of the critical sequences for expression in transgenic mice revealed that the proximal 180 bp of the GH promoter contained all of the information necessary to target expression to the anterior pituitary gland [Lira et al. 1988, 1993]. Therefore, we systematically examined evolutionarily well-conserved sequences exclusive of the Pit-1 and T₃ response elements in the proximal GH promoter and tested the effects of their disruption on transcription in a somatotroph cell line (GC); we found that the functionally most important sequence (−110 to −95) lies between the proximal and distal Pit-1-binding sites and is unusually well conserved across species. When this sequence was mutated in the context of the proximal 320 bp of the rat GH promoter as a transgene, promoter activity was impaired >100-fold. On the basis of the functional importance of this element in vivo, we identified a major pituitary cell-binding activity to this site, referred to as Zn-15 (pronounced Zen-15), a novel transcription factor possessing a structurally unusual DNA-binding domain consisting of three CysX₂₋₄CysX₁₁₋₁₆HisX₆ zinc fingers of similar sequence. These three zinc fingers are separated by surprisingly extensive linker sequences that would be predicted to interrupt their ability to form a unitary DNA-binding domain. Zn-15 synergizes with Pit-1 to activate the GH promoter in heterologous cell lines in which this promoter is only minimally responsive to Pit-1 alone. These data suggest that synergistic interactions between Pit-1 and this novel zinc finger factor binding an evolutionarily conserved region in the GH promoter may be required for the effective activation of the GH gene.

Results

The GH Z box is required for effective expression of the GH promoter

Because the proximal 180 bp of the rat GH promoter contains sufficient information to target pituitary-specific expression in transgenic mice, we hypothesized that evolutionarily conserved sequences in this promoter region additional to the Pit-1-binding sites were likely to contribute to effective expression of the GH gene. We therefore mutated three well-conserved sites in this promoter region and functionally evaluated the effects of these mutations in transiently transfected rat pituitary tumor cells (GC) [Fig. 1A]. Mutation of two conserved sequences [a putative AP-2-binding site (−159/−148) [Lefevre et al. 1987] and a putative Spl site (−142/−130) [Lemaigre et al. 1990; Schaufele et al. 1990] had little effect [Fig. 1B]. In contrast, mutation of a third conserved sequence (−110/−95) located between the proximal and distal Pit-1-binding sites markedly decreased activity [referred to as the GH Z box; Fig. 1A,C]. Previously, footprinting experiments using GC cell extracts had revealed the presence of DNase I hypersensitive sites over this sequence, suggesting that a binding activity to this site might exist in somatotrophs [Mangalam et al. 1989; H. Mangalam, pers. comm.].

To evaluate the importance of the GH Z box in vivo, pedigrees of transgenic mice carrying either a wild-type or mutated GH Z box in the context of the proximal 320 bp rat GH promoter fused to a human GH (hGH) reporter gene (Fig. 2A) were established, and the pituitary hGH content from these mice quantitated by radioimmunoassay. The GH Z box mutation in vivo resulted in a decrease of >100-fold in reporter gene expression [Fig. 2B]. No ectopic expression of the hGH transgene was observed in any pedigree further characterized [Fig. 2C; data not shown]. To confirm the dramatic effect of the GH Z box mutation on transgene expression, in situ hybridization was performed. Sections of adult anterior pituitary glands from mice carrying the wild-type GH promoter transgene showed intense hybridization signal compared with that observed in pituitaries from mice carrying the GH Z box mutation when probed with a 35S-labeled oligonucleotide probe specific for hGH [Fig. 3]. As expected, pituitaries from mice carrying either transgene construct hybridized strongly to a mouse GH (mGH)-specific probe [Fig. 3].

Characterization and Cloning of the GH Z box-binding factor

When a 32P-labeled oligonucleotide encompassing the GH Z box site was used to screen rat pituitary and somatotroph cell (GC) Àgt11 expression libraries, and a clone that bound specifically to this site was further characterized. This clone contained partial sequences of a novel member of the superfamily of transcriptional regulators containing several CysX₂₋₄CysX₁₁₋₁₆HisX₆ zinc fingers [Rhodes and Klug 1993]. Multiple overlapping clones were isolated to obtain the full-length cDNA [Fig. 5]. Stop codons in all three frames precede an ATG that initiates a 6.5-kb continuous open reading frame (ORF) containing 15 putative CysX₂₋₄CysX₁₁₋₁₆HisX₆ zinc fingers [Fig. 5]. Because of this distinguishing structural feature, we call the protein Zn-15.

Zn-15 has no described gene homolog in any species. The full Zn-15 ORF predicts a protein with a molecular
mass of 240 kD. However, in vitro translation of the full coding sequence produces a protein that migrates on denaturing SDS-polyacrylamide gels with a mobility of ~180 kD, in addition to a minor 75-kD product [Fig. 6C]. To resolve this discrepancy between predicted and apparent molecular mass on SDS-polyacrylamide gels, we raised a polyclonal antisera reactive against the Zn-15 carboxyl terminus. Western blot analysis of nuclear extracts made from rat somatotroph (GC) cells [Fig. 6C] revealed that antisera specific for Zn-15 reacted specifically against a 180-kD protein and also a 75-kD protein that appears to represent a proteolytic product [Fig. 6C; data not shown]. Because of the concordance between the size of in vitro-translated Zn-15 and the higher molecular mass species present in cell extracts, we consider it likely that the full-length Zn-15 ORF migrates at a size anomalous to its predicted molecular mass. Aberrant electrophoretic behavior has also been observed for several other large members of the CysX2â’4CysX11â’16HisX3â’6His class of transcription factors [Ruppert et al. 1990; Read and Manley 1992; Settleman et al. 1992].

Probing GC cell mRNA with a Zn-15 carboxy-terminal coding fragment revealed that the rat Zn-15 mRNA is expressed as a single band of ~7.4 kb in size [Fig. 6D].RNase protection analysis of a panel of rat tissues with an 863bp fragment from the Zn-15 carboxy-terminal zinc finger cluster demonstrates that Zn-15 mRNA is expressed in pituitary and nonpituitary tissues, including spleen, pituitary, and heart [Fig. 6E]. Thus, unlike Pit-1, Zn-15 expression is not restricted to the anterior pituitary gland.

Zn-15 possesses a DNA-binding domain consisting of three zinc fingers with extremely elongated linker sequences

A fragment encompassing six of the seven zinc fingers in
A novel factor critical for growth hormone expression

Figure 2. Functional analysis of the GH Z box mutation in the expression of a transgene in mice. (A) Depiction of the GH wild-type and GH Z box mutant transgene constructs. The wild-type proximal GH promoter (GH WT) spanning from −320 to +08 relative to the start of transcription, and the the mutated GH Z box (−110/−95) (Z BOX MUT) in the identical promoter context were fused to a human genomic GH reporter gene [hGH]. (B) Quantitative analysis of GH WT and GH Z BOX MUT promoter-driven hGH transgene expression in the pituitary gland. The two promoter/hGH fusion constructs were injected into mouse oocytes; three separate lines of founder mice containing stably integrated copies of the GH WT construct (W7, W19, and W26), and four separate lines containing the Z BOX MUT construct (mZ18, mZ25, mZ33, and mZ48) were established and used for further analysis. Pituitary glands were excised from adult mice, and the hGH content was quantitated by radioimmunoassay (Nicholls Institute) after normalization for protein (Bio-Rad). The expression levels of each transgenic line (nanograms of hGH/10 μg of protein ± S.E.M.) are, respectively, W7, 60.9 ± 19.1 (n = 4); W19, 68.0 ± 13.0 (n = 6); W26, 66.6 ± 12.3 (n = 3); mZ18, 0.1 ± 0.1 (n = 8); mZ25, 0.1 ± 0.1 (n = 1); mZ33, 0.37 ± 0.2 (n = 3); mZ48, 1.58 ± 1.2 (n = 8). (C) Tissue specificity of GH WT transgene expression. A panel of nonpituitary tissues from all transgenic lines further characterized were excised from adult mice and quantitated for hGH content. Shown are representative results from a mouse carrying the GH WT transgene (W19 pedigree).

the carboxy-terminal cluster encoded in the original λgt11 phage insert was expressed in bacteria (Zn-15 C6), and the specificity of Zn-15 binding to the GH Z box was examined. Zn-15 C6 bound with high affinity to a 32P-labeled GH Z box oligonucleotide but not to the mutated site that had been substituted for the Z box in the initial scanning mutagenesis of the GH promoter, nor to several other sites tested (Fig. 7A). Unexpectedly, we found that Zn-15 can also bind with high affinity to the T3RE from the rat α-myosin heavy chain gene [MHC T3RE] but not to other T3REs tested (Fig. 7A). Alignment of the GH Z box and the α-MHC T3RE sequences shows a high degree of sequence identity (see Fig. 8C).

To determine which of these six zinc fingers are critical for binding to these two sites, a series of six mutant Zn-15 C6 proteins were expressed in bacteria in which the first conserved histidine residue of each of the six fingers was mutated to a serine. Mutation of either fingers IX, X, or XI abolished all binding to the GH Z box while mutation of fingers XII, XIII, or XIV was without significant effect (Fig. 7B). The same mutations also abolished αMHC T3RE binding (data not shown). Therefore, zinc fingers IX, X, and XI are each necessary for binding to both recognition elements. The cooperation of these three zinc fingers to form a unitary DNA-binding domain is somewhat unexpected in that the linker sequence between zinc fingers IX and X is 21 amino acids and the linker sequence between zinc fingers X and XI extends a full 143 amino acids. Previously, in all of the other zinc finger proteins whose DNA-binding abilities have been directly tested, only zinc fingers separated by short (usually 7 amino acids) linker sequences in tandem have been shown to cooperate to form a single DNA-binding domain. Because zinc loop fingers that are spaced widely are a common feature among members of this superfamily, our data suggest that such widely spaced fingers may contribute DNA-binding domains with DNA site specificity comparable to those of factors that contain tandem arrays of closely spaced zinc fingers.

To examine the ability of the amino-terminal six zinc fingers to bind DNA, this region of Zn-15 was also expressed in bacteria (Zn-15 N6). Interestingly, several of the zinc fingers possess residues that are predicted to disrupt the ability of this structure to bind DNA (Pav-
Zn-15 represents the major GH Z box-binding activity in cells expressing the endogenous GH gene.

**Zn-15 is a trans-activator that synergizes with Pit-1 to activate the GH promoter**

As determined by EMSA analysis, GH Z box-binding activity is low in African green monkey kidney cells (CV-1) (Fig. 4A). Thus, this cell line was chosen initially to examine the transcriptional effects of Zn-15. As shown in Figure 9A, transfection of Zn-15 was able to stimulate transcription from constructs containing either one copy of the GH Z box fused to a minimal prolactin promoter or one copy of the αMHC T3RE fused to the herpes sim-
A novel factor critical for growth hormone expression

Figure 5. Zn-15 cDNA amino acid sequence. Each of the 15 putative CysX_2-CysX_2-HisX_3-His zinc fingers are numbered and reverse highlighted.
Figure 6. Zn-15 protein and mRNA expression. (A) Schematic diagram of the Zn-15 cDNA-coding sequence. Vertical lines depict individual putative zinc fingers. (B) Alignment of Zn-15 putative zinc fingers. The conserved cysteine and histidine residues in each zinc finger are reverse highlighted. Amino acids that are not consistent with zinc finger consensus residues at their respective positions are circled. The three zinc fingers that are critical for binding to the GH Z box (see Fig. 7) are lightly shaded. (C) Comparison of endogenous somatotroph Zn-15 protein and in vitro-translated Zn-15 SDS-polyacrylamide gel migration. Ten micrograms of GC cell nuclear extract and 4 μl of [35S]-labeled in vitro-translated Zn-15 cDNA (TnT System, Promega) were run on a 5% SDS-polyacrylamide gel. The gel was transferred to nitrocellulose, and the lane containing the GC cell nuclear extract was probed with an antibody directed against the carboxyl terminus of Zn-15 (αZn-15) and visualized with a chemiluminescent secondary antibody (ECL, Amersham). The lane containing the [35S]-labeled in vitro-translated Zn-15 (Zn-15 I.V.T.) was exposed directly to film. Protein molecular weight markers are indicated. (D) RNA blot of GC somatotroph cell mRNA. Twelve micrograms of denatured size-fractionated poly(A)-selected mRNA was transferred to nitrocellulose and probed with a random-primed 1-kb PvuII fragment from the Zn-15 carboxyl terminus. mRNA size standards (in kb) are indicated at left. (E) RNase protection panel of Zn-15 expression. A 32P-labeled antisense cRNA probe from the Zn-15 DNA-binding domain was incubated with 20 μg of total RNA from rat pituitary, testes, heart, spleen, liver, muscle, and kidney. A rat actin antisense cRNA probe was used to normalize the quality and quantity of RNA added to each reaction. The arrow indicates the size of the protected probe (863 bp). Size standards are indicated at right. [tRNA] Yeast transfer RNA control; [actin] rat β-actin probe.

bp of the rat GH promoter [Fig. 9B]. In the same experiments, transfection of a Pit-1 cDNA strongly stimulated expression (74-fold) of the rat prolactin distal enhancer and proximal promoter [data not shown]. On the GH promoter, transfection of Zn-15 also activated expression weakly. Interestingly, when both Pit-1 and Zn-15 were cotransfected, the GH promoter was stimulated by an amount greater than the multiplicative product of their separate folds of activation. No synergism was observed when Pit-1 and Zn-15 were cotransfected with the GH promoter containing a mutated GH Z box. Similar results were obtained in another cell line of macrophage origin [U937] [data not shown].

The mechanisms underlying transcriptional synergism are poorly understood. To determine whether synergism requires only transcription factor binding or whether both factors must also be transcriptionally competent, we tested several deletion mutants of Zn-15 and Pit-1 for their ability to synergize with each other on the GH promoter. Truncation of the amino-terminal zinc finger cluster [Zn-15 Δ1-855] did not appreciably affect the ability of Zn-15 to trans-activate or synergize with Pit-1 [Fig. 10A]. A further amino-terminal deletion [Zn-15 Δ1-1214] decreased, but did not abolish, both trans-activation and synergism with Pit-1 [Fig. 10A]. However, a Zn-15 fragment containing only the carboxy-terminal zinc finger cluster [Fig. 10A, Zn-15 DBD] was incompetent for both trans-activation from the GH Z box sequence and synergism with Pit-1. While these results are in concordance with a model in which the DNA-binding domain and carboxy terminus of Zn-15 contain sufficient information for both trans-activation and synergism functions of the various Zn-15 deletion mutants may reflect different levels of expression in the nucleus. To determine which regions of Pit-1 are required for synergism with Zn-15, a series of Pit-1 amino
A novel factor critical for growth hormone expression

Figure 7. Analyses of Zn-15 DNA-binding activity. (A) EMSA assay of the Zn-15 carboxy-terminal zinc finger cluster binding specificity. $^{32}$P-Labeled sites encompassing the sequences of the GH Z box (GH Z BOX), mutated GH Z box (mut Z BOX), α-myosin heavy chain T$_2$ response element (MHC TRE), PRL Pit-1-binding site (Prl 1P), rat GH T$_3$ response element (rGH TRE), palindromic T$_3$ response element (TRE PAL), and herpes simplex virus octamer-binding site (HSV OCT) were used as probes for bacterially expressed Zn-15 carboxy-terminal zinc finger cluster [Zn-15 C6]. (B) Mutational analysis of Zn-15 zinc finger binding to the GH Z box. The first conserved histidine residue in each of the six Zn-15 carboxy-terminal cluster zinc fingers (fingers IX–XIV) was altered separately to a serine residue by conventional mutagenesis. Each of these six point mutants was expressed in bacteria in the context of the whole Zn-15 carboxy-terminal zinc finger cluster and analyzed for GH Z box-binding ability. (C) EMSA of the Zn-15 amino-terminal zinc finger cluster. The bacterially expressed amino-terminal zinc finger cluster (Zn-15 N6) and carboxy-terminal cluster were tested for binding to a $^{32}$P-labeled probe consisting of completely degenerate 16-mers representing all possible binding sites [(H.B.S.) heterogeneous binding sites] and the GH Z box. Approximately equal amounts of Zn-15 N6 and Zn-15 C6 protein were added to the binding reaction (data not shown). Three times the amount of cpm for the heterogeneous binding site oligonucleotide compared with the homogenous Z box gel shift probe was incubated with the bacterial protein extract, because only a fraction of the probe sequences would be expected to function as potential binding sites.

acid substitution mutants were constructed and functionally evaluated in transfection. A transcriptionally incompetent Pit-1 molecule consisting of the POU homeo domain and the carboxyl terminus [AN Pit-1] is unable to synergize with Zn-15, as is a second transcriptionally incompetent Pit-1 mutant protein containing a proline for alanine substitution in the POU domain that causes hereditary dwarfism in humans [Pit-1 A158P; Pfaeffle et al. 1992] (Fig. 10B).

Discussion

On the basis of the analysis of mouse and human dwarfs who carry Pit-1 genomic mutations, Pit-1 is clearly required for GH gene activation. Yet, the actions of other transcription factors are also necessary for activation of Pit-1 target genes. For example, Pit-1 does not effectively activate the prolactin distal enhancer, in the absence of ligand-bound estrogen receptor [Day et al. 1990; Simmons et al. 1990]. Similarly, Pit-1 synergizes with ligand-bound retinoic acid receptor on the Pit-1 gene enhancer [Rhodes et al. 1993]. Here, we describe a new sequence conserved among hominid and quadruped GH promoters that constitutes an important element in regulating GH expression. By transgenic mouse analyses we show that the GH Z box, the binding site for Zn-15, constitutes a crucial regulatory element because its mutation decreases transgene expression >100-fold. These data, coupled with the observation that Zn-15 represents a major binding activity to this site, suggest that this factor may be an important component of GH gene expression. In concert with this model, Zn-15 is capable of synergizing with Pit-1 in GH gene activation in heterologous cells in which Pit-1 alone produces only ineffective stimulation.

The molecular mechanisms by which transcription factors synergize are not well defined. In some cases, transcriptional synergism appears to reflect cooperative DNA-binding interactions [Jiang and Levine 1993]. How-
ever, this does not appear to be the case for Zn-15 and Pit-1, as in EMSA studies we can detect no cooperativity in DNA binding between Pit-1 and the bacterially expressed Zn-15 carboxyl terminus (data not shown). Synergy requires that both proteins be trans-activationally competent, and for Pit-1 the amino acid sequences required are distinct from those necessary for synergism with the estrogen receptor in the activation of the prolactin distal enhancer (J. Holloway, in prep.), an observation that has intriguing implications with respect to cell type-specific expression of the GH and PRL genes.

Zn-15 is a novel member of the Cys/His zinc finger gene family

The structural aspects of Zn-15 may be informative in the study of other members of the zinc finger superfamily of transcription factors. We have found that fingers IX, X, and XI cooperate to form a unitary DNA-binding domain. Intriguingly, the linker sequences between these three fingers [21 and 143 amino acids, respectively] are much longer than the canonical 7 amino acid linkers observed to space zinc fingers in tandem arrays in all other members of this gene family in which DNA binding has been directly tested. Therefore, under the appropriate circumstances, widely spaced zinc fingers can cooperate in DNA binding. Previously, binding site selection (SAAB) studies with the Drosophila CF2 gene had suggested that zinc fingers separated by long linker sequences were unable to contribute to binding site recognition (Hsu et al. 1992). In our studies of Zn-15 DNA-binding specificity, we have found that Zn-15 can bind to the αMHC T3RE. Alignment of these two sequences shows that they have a high degree of base pair identity over the area with which Zn-15 interacts with guanine residues. Intriguingly, in the αMHC T3RE, the Zn-15 binding site overlaps with bases contacted by T3 receptor/RXR heterodimers, and in EMSA analysis Zn-15 can compete for binding with T3 receptor/RXR heterodimers (A. Näär, unpubl.). These data suggest that Zn-15 is ca-
A novel factor critical for growth hormone expression

Figure 9. Transient transfection analyses of Zn-15 transcriptional activity in CV-1 cells. (A) Stimulation of transcription by Zn-15 and Zn-15 deletion mutants from the GH Z box and α-MHC T3REs. Expression plasmids encoding Zn-15 or Zn-15 deletion mutants under the control of the RSV promoter [Simmons et al. 1990] were cotransfected with a luciferase cDNA under the control of the PRL minimal promoter (P36), one copy of the GH Z box in front of the PRL minimal promoter [ZRE P36], the HSV TK promoter (TK), or one copy of the α-MHC T3RE in front of the TK promoter (MHC TK). Background was determined by using an RSV expression vector driving the expression of a gene encoding neomycin resistance (RSV Neo) [see Materials and Methods]. Results are the average of triplicate determinations, and error bars represent S.E.M. Similar results were obtained in four experiments of similar design. (Zn-15 A1-855) Zn-15 carboxy-terminal coding fragment deleting the first 855 amino acids; (Zn-15 A1-1214) Zn-15 carboxy-terminal coding fragment deleting the first 1214 amino acids; (Zn-15 A1215-2177) Zn-15 amino-terminal coding fragment deleting the carboxy-terminal 963 amino acids; (Zn-15 DBD) Zn-15-coding fragment spanning amino acids 1214-1753. (B) Cooperative interactions of Zn-15 and Pit-1 on the GH promoter and lack of cooperativity on the GH promoter containing a mutated Z box. Transfections were performed as in (A) [also see Materials and methods]. Transfections were balanced with RSV Neo so that the same amount of plasmid DNA was added to each plate of cells. [T3R] human β thyroid hormone receptor.

The role of the other 12 putative zinc finger sequences in the Zn-15-coding sequence that do not participate in GH Z box or αMHC T3RE binding is not entirely clear. Given the widespread distribution of Zn-15 mRNA and protein, these fingers might be involved in binding to other gene promoters. We have been unable to demonstrate any capacity for DNA binding of the first six amino-terminal zinc fingers. However, it is still possible that sequences exist to which these zinc fingers bind that we could not detect in our assay. In the CF2 zinc finger protein, a similar inability of amino-terminal zinc fingers to bind DNA specifically has also been observed [Gogos et al. 1992]. Recently, basic amino acid sequences adjacent to zinc fingers have been shown to influence finger/DNA interactions [Keller and Maniatis 1992]. It is thus possible that adjacent sequences necessary for specific DNA binding are lacking in both Zn-15 and CF2 amino-terminal fingers. Alternatively, zinc fingers have been demonstrated to bind to specific secondary structures of of RNA [Theunissen et al. 1992], and it is possible that the Zn-15 amino-terminal group of zinc fingers might subserve such a function. The observation that Zn-15, a novel member of the zinc finger transcription factor superfamily [Rhodes and Klug 1993], binds to an evolutionarily conserved GH promoter sequence required for effective activation of this promoter in vivo raises intriguing questions about the role of this protein in development. Previously, several DNA–protein-binding activities on the GH promoter were characterized, and Zn-15 is likely to correspond to a binding activity referred to as GHF-5 [Schaufele et al. 1990].

In conclusion, Zn-15 is a novel zinc finger protein that binds to an evolutionarily conserved sequence in the GH gene required for effective Pit-1-dependent activation of this promoter in vivo and is capable of synergizing with Pit-1 in experiments in cultured cells. These observations may help to define the combinatorial code that underlies the molecular basis of somatotroph and lactotroph-specific gene transcription during mammalian organogenesis.

GENES & DEVELOPMENT 1683
Figure 10. Deletional analysis of regions of Zn-15 and Pit-1 necessary for synergistic interactions on the GH promoter. (A) Mapping of Zn-15 domains necessary for synergism with Pit-1. Transfections were performed as in Fig. 9. Transfections were balanced with RSV Neo so that the same amount of plasmid DNA was added to each plate of cells. (B) Mapping of Pit-1 domains necessary for synergism with Zn-15. [AN PIT-1] Deletion mutant of Pit-1 lacking the amino-terminal trans-activation domains (Ingraham et al. 1988). [PIT-1 A158P] Pit-1 mutant containing a proline for alanine substitution at amino acid 158 in the POU-specific domain (Pfäffle et al. 1992).

Materials and Methods

Cloning of Zn-15

A multimerized Z box sequence [5'-AATTCAGCACAAGCTGTCAGTGGAATT-3'] was used to screen rat pituitary GC cell Xgt11 expression libraries as described previously (Ingraham et al. 1988). Multiple overlapping clones were isolated from rat pituitary and lactotroph cell line (MMQ) cDNA libraries. A large number of oligonucleotides were synthesized to sequence the full Zn-15 ORF (Sequenase, U.S. Biochemical).

Bacterial and reticulate lysate expression of Zn-15, antibody generation, and Western analysis

For in vitro translation, the full Zn-15 cDNA was subcloned into the vector pCITE 2a (Novagen) and translated in the presence of [35S]methionine using the TnT System-coupled transcription and translation system (Promega), with the addition of 100 mM KCl to enhance full-length translation.

A 1.6-kb coding fragment containing zinc fingers IX-XIV [Zn-15 C6; XbaI (nucleotide 3642)/Msci (nucleotide 5259)] and a 1.7-kb coding fragment containing zinc fingers I-VI [Zn-15 N6; BamHI (nucleotide -6)/PstI (nucleotide 1768)] were expressed as glutathione S-transferase (GST) fusion protein according to standard protocols (Pharmacia). For the generation of antiseras reactive against Zn-15, the carboxyl terminus of Zn-15 [133 amino acids] was also expressed as a GST fusion protein and purified over glutathione–agarose (Sigma). GST Zn-15 carboxyl terminus fusion protein was excised from protein gels, emulsified with RIBI adjuvant, and injected into white, female New Zealand rabbits. Antisera used in this manuscript are all from the third bleed. For Western analysis, GC and 235-1 cell nuclear extracts were made as described previously (Dignam et al. 1983), with the addition of extra protease inhibitors (4 mM PMSF, 2 U/ml of antipain, 2 U/ml of leupeptin, 2 U/ml of aprotinin, 4 mM benzamidine) to staunch Zn-15 protein degradation. Blots were incubated with 1 : 1000 dilution of antiseras under the manufacturer's suggested protocol for the ECL Western Blot Kit (Amersham).

EMSA, antibody perturbation, and methylation interference

Double-stranded oligonucleotides encoding the GH Z box
DNA was used because the Zn-15-coding sequence (6.55 kb) is luciferase reporter plasmid, and, as indicated, 1 lag of CMV Pit-1 receptor, RSV Zn-15, RSV Zn-15A1215-2177. More Zn-15-containing plasmid (5'-AATTCAGACAAAGCTGTACGGTGGCAGAGAT-3'), or palindromic T3RE (5'-AAGGGGATCCGGTAAGATTCATGAAGGTG-3'), or rat GH Z box sequences beyond nucleotide 3560, and religating.

Generation of transgenic mice
Plasmid pGH320-hGH was constructed by fusing the proximal 320 bp (-310 to +8) rat GH of the rat GH promoter to the genomic hGH gene [Nicholls Institute, San Juan Capistrano, CA]. Plasmid pGH320 GH Z box Mut-hGH was similarly constructed, except that the GH Z box sequence had been transverted by conventional mutagenesis, as described in EMSA methods. The rat GH promoter/hGH structural gene fusion DNA was purified and injected (1-5 μg/ml) into mouse oocytes as described previously [Lira et al. 1988]. All transgenic lines further characterized were shown to carry intact copies of the transgene as determined by Southern analysis.

hGH Reporter gene quantification in transgenic mice
Immunooquantification of transgenic mouse pituitary hGH was performed as described previously [Lira et al. 1988].

Transfection analyses
Reporter plasmid vectors containing rat GH promoter information (-310 to +8), rat GH promoter with mutated -159/-148 site (5'-TCTACTGGTTTCTC-3'), -142/-130 (5'-GGTGTGGTTCTTCTT-3') site, or GH Z box sequences (see legend to Fig. 1 and EMSA methods above), rat PRL 1P Pit-1-binding site (Prl 1P: 5'-GAATGCTAATGATATTCT'~3'), or GH Z box sequences (see legend to Fig. 1) were transfected by calcium phosphate precipitation with 1 lag of plasmid DNA. DNA was used because the Zn-15-coding sequence (6.55 kb) is larger than the entire RSV expression vector (3.9 kb). Twenty-four hours post-transfection, cells were washed with serum-free DMEM, incubated in 3 ml of DMEM containing 10% charcoal-stripped serum [Scantibodies Laboratories], and cells were harvested and luciferase activity was assayed on the following day as described previously [Drolet et al. 1989].

Plasmid construction
For generation of the zinc finger IX-XIV DNA-binding mutants, the first conserved histidine residue and the preceding amino acid were replaced by a glycine and a serine residue (BamHI site) by conventional mutagenesis (T7 Polymerase Mutagen, Bio-Rad). Zn-15 Δ1-855 was constructed by excising a 5' flanking (nucleotide 2565)/XbaI (3' polylinker) fragment from pCITE2a Zn-15, which was inserted into an RSV expression vector. Similarly, RSV Zn-15Δ1-1214 was constructed by subcloning a Zn-15 Xbal (nucleotide 3642)/XbaI (3' polylinker)-coding fragment into the same vector. RSV Zn-15Δ1-2177 was constructed by cutting RSV Zn-15 with Xbal, which excised all coding sequences beyond nucleotide 3560, and religating.

In situ hybridization analysis
In situ hybridization with 10-μm sagittal sections of adult transgenic mouse anterior pituitary glands was performed essentially as described previously [Aubry et al. 1993] with oligonucleotides from the 3'-untranslated sequences of either mouse (5'-GGGGCAGGGGAGGCAGGGAGTGCAGGAGAGAC-3') or human (5'-TCAGGGTGAGTGGCTA-3') GH mRNAs.

Acknowledgments
We gratefully acknowledge Dr. J. Voss and B. McEvilly for construction of the original GH promoter mutants, C. Nelson for invaluable assistance with tissue culture, G. DiMattia for assistance with analyses of transgenic mice, L. Erkman and P. Sawchenko for assistance with in situ analysis, H. Mangalam for computer analysis, and Drs. B. Andersen and C. Glass for critical reading of this manuscript. M.G.R. is an investigator with the Howard Hughes Medical Institute. These studies were supported in part by a grant from the National Institutes of Health (DK18477).

The publication costs of this article were defrayed in part by payment of page charges. This article must therefore be hereby marked "advertisement" in accordance with 18 USC section 1734 solely to indicate this fact.

Note added in proof
The Zn-15 nucleotide sequence has been submitted to the GenBank data library.

References
Aubry, J.-M., M.-F. Schulz, S. Pagliai, P. Schulz, and J.Z. Kiss. 1993. Coexpression of dopamine D2 and substance P (Neurokinin-1) receptor messenger RNAs by a subpopulation of cholinergic neurons in the rat striatum. Neuroscience 53: 417-424.
Bodner, M. and M. Karin. 1987. A pituitary-specific trans-acting factor can stimulate transcription from the growth hormone promoter in extracts of nonexpressing cells. Cell 50: 267-275.
Bodner, M., J.L. Castrillo, L.E. Theill, T. Deerinck, M. Ellisman, and M. Karin. 1988. The pituitary-specific transcription factor GHF-1 is a homeobox-containing protein. Cell 55: 505–518.

Brent, G.A., J.W. Harney, Y. Chen, R.G. Warne, D.D. Moore, and P.R. Larsen. 1989. Mutations of the rat growth hormone promoter which increase and decrease response to thyroid hormone define a consensus thyroid hormone response element. Mol. Endocrin. 3: 1996–2007.

Day, R.N., S. Koike, M. Sakai, M. Muramatsu, and R.A. Maurer. 1990. Both Pit-1 and the estrogen receptor are required for estrogen responsiveness of the rat prolactin gene. Mol. Endocrin. 4: 1964–1971.

Dignam, J.D., R.M. Lebovitz, and R. Roeder. 1983. Accurate transcriptional initiation by RNA polymerase II in a soluble extract from isolated mammalian nuclei. Nucleic Acids Res. 11: 1475–1489.

Dolle, P., J.L. Castrillo, L. Theill, T. Deerinck, and M. Ellisman. 1990. Expression of c-GHF-1 protein in mouse pituitaries correlates both temporally and spatially with the onset of growth hormone gene activation. Cell 60: 808–820.

Drolet, D.W., K.M. Scully, D.M. Simmons, M. Wegner, K.T. Chu, L.W. Swanson, and M.G. Rosenfeld. 1991. TEF, a transcription factor expressed specifically in the anterior pituitary during embryogenesis, defines a new class of leucine zipper proteins. Genes & Dev. 5: 1739–1753.

Evans, R.M., N.C. Birnberg, and M.G. Rosenfeld. 1982. Glucocorticoids and thyroid hormone transcriptionally regulate growth hormone gene expression. Proc. Natl. Acad. Sci. USA 79: 7659–7663.

Glass, C.K., R. Franco, C. Weinberger, V.R. Albert, R.M. Evans, and M.G. Rosenfeld. 1987. A c-erbA binding site in rat growth hormone gene mediates trans-activation by thyroid hormone. Nature 329: 738–741.

Gogos, J.A., T. Hsu, J. Bolton, and F.C. Kafatos. 1992. Sequence discrimination by alternatively spliced isoforms of a DNA binding zinc finger domain. Science 257: 1951–1954.

He, X., M.N. Tracey, D.M. Simmons, H.A. Ingraham, L.W. Swanson, and M.G. Rosenfeld. 1989. Expression of a large family of POU-domain regulatory genes in mammalian brain development [published erratum appears in Nature [1989] Aug. 24, 340(6235):662]. Nature 340: 35–41.

Hsu, T., J.A. Gogos, S.A. Kirsh, and F.C. Kafatos. 1992. Multiple zinc finger forms resulting from developmentally regulated alternative splicing of a transcription factor gene. Science 257: 1946–1950.

Ingraham, H.I, R. Chen, H. Mangalam, H. Elsholtz, S. Flynn, C. Lin, D. Simmons, L. Swanson, and M.G. Rosenfeld. 1988. A tissue-specific transcription factor containing a homeodomain specifies a pituitary phenotype. Cell 55: 519–529.

Jiang, J. and M. Levine. 1993. Cooperative interactions with bHLH activators delimit threshold responses to the dorsal gradient morphogen. Cell 72: 741–752.

Keller, A.D. and T. Maniatis. 1992. Only two of the five zinc fingers of the Drosophila etonal gene encode zinc finger proteins with distinct DNA binding specificities. EMBO J. 11: 1035–1044.

Li, S., E.B. Crenshaw III, E.J. Rawson, D.M. Simmons, L.W. Swanson, and M.G. Rosenfeld. 1990. Dwarf locus mutants lacking three pituitary cell types result from mutations in the POU-domain gene pit-1. Nature 347: 528–533.

Lira, S.A., E.B. Crenshaw III, C.K. Glass, L.W. Swanson, and M.G. Rosenfeld. 1988. Identification of rat growth hormone genomic sequences targeting pituitary expression in transgenic mice. Proc. Natl. Acad. Sci. 85: 4755–4759.

Lira, S.A., K. Kalla, C. Glass, and M.G. Rosenfeld. 1993. Synergistic interactions between Pit-1 and other elements are required for effective somatotroph rat growth hormone gene expression in transgenic mice. Mol. Endocrinol. 7: 694–701.

Mangalam, H.J., V.R. Albert, H.A. Ingraham, M. Kapiloff, L. Wilson, C. Nelson, H. Elsholtz, and M.G. Rosenfeld. 1989. A pituitary POU domain protein, Pit-1, activates both growth hormone and prolactin promoters transcriptionally. Genes & Dev. 3: 946–958.

Nelson, C., E.B. Crenshaw III, R. Franco, S.A. Lira, V.R. Albert, R.M. Evans, and M.G. Rosenfeld. 1986. Discreet cis-active sequences dictate the cell type-specific expression of rat prolactin and growth hormone genes. Nature 322: 557–562.

Nelson, C., V.R. Albert, H.P. Elsholtz, L.I. Lu, and M.G. Rosenfeld. 1988. Activation of cell-specific expression of rat growth hormone and prolactin genes by a common transcription factor. Science 240: 1400–1405.

Pavletich, N.P. and C. Pabo. 1991. Zinc-finger-DNA recognition: Crystal structure of a Zif268-DNA complex at 2.1 angstrom. Science 252: 809–816.

Pfaffl, M.W., G.E. DiMattia, J.S. Parks, M.R. Brown, J.M. Wit, M. Jansen, H. Van der Nat, J.L. Van den Brande, M.G. Rosenfeld, and H.I. Ingraham. 1992. Mutation of the POU-specific domain of pit-1 and hypopituitarism without pituitary hypoplasia. Science 257: 1118–1120.

Radovick, S., M. Nations, Y. Du, L.A. Berg, B.D. Weintraub, and F.E. Wondisford. 1992. A mutation in the POU-Homodomain of Pit-1 responsible for combined pituitary hormone deficiency. Science 257: 1115–1117.

Read, D. and L.J. Manley. 1992. Alternatively spliced transcripts of the Drosophila tramtrack gene encode zinc finger proteins with distinct DNA binding specificities. EMBO J. 11: 1035–1044.

Rhodes, S., R. Chen, G. DiMattia, K. Scully, K. Kalla, S. -C. Lin, V. Yu, and M.G. Rosenfeld. 1993. A tissue-specific enhancer confers Pit-1 dependent morphogen inducibility and auto-regulation upon the Pit-1 gene. Genes & Dev. 7: 913–932.

Rhodes, D. and A. Klug. 1993. Zinc fingers. Sci. Am. 268: 56–65.

Ruppert, J.M., B. Vogelstein, K. Arheden, and K. Kinzler. 1990. GLI3 encodes a 190-kilodalton protein with multiple regions of GLI similarity. Mol. Cell. Biol. 10: 5408–5415.

Schaufele, F., B.L. West, and T. Reudelhuber. 1990. Somaticotroph and lactotroph specific interactions with the homeobox protein binding sites in the rat growth hormone gene promoter. Nucleic Acids Res. 18: 5235–5243.

Schaufele, F., B.L. West, and T. Reudelhuber. 1990. Overlapping Pit-1 and Sp1 binding sites are both essential to full rat growth hormone gene promoter activity despite mutually exclusive Pit-1 and Sp1 binding. J. Biol. Chem. 265: 17189–17196.

Settleman, J., V. Narasimhan, L.C. Foster, and R.A. Weinberg. 1992. Molecular cloning of cDNAs encoding the GAP-associated protein p190: Implications for a signaling pathway from ras to the nucleus. Cell 69: 539–549.

Sharp, D.Z. and Y. Cao. 1990. Regulation of cell-type-specific transcription and differentiation of the pituitary. BioEssays 12(2): 80–85.

Simmons, D.M., J.W. Voss, H.A. Ingraham, J.M. Holloway, R.S.
A novel factor critical for growth hormone expression

Broide, M.G. Rosenfeld, and L.W. Swanson. 1990. Pituitary cell phenotypes involve cell-specific Pit-1 mRNA translation and synergistic interactions with other classes of transcription factors. *Genes & Dev.* 4: 695–711.

Tatsumi, K., K. Miyai, T. Notomi, K. Kaibe, N. Amino, Y. Mizuno, and H. Kohno. 1992. Cretinism with combined hormone deficiency caused by a mutation in the PIT1 gene. *Nat. Genet.* 1: 56–58.

Theunissen, O., F. Rudt, U. Guddat, H. Mentzel, and T. Pieler. 1992. RNA and DNA-binding zinc fingers in Xenopus TFI-IIA. *Cell* 71: 679–690.

Voss, J. and M.G. Rosenfeld. 1992. Anterior pituitary development: Short tales from dwarf mice. *Cell* 70: 527–530.

Zhang, W., R. Brooks, D. Silversides, B.L. West, F. Leidig, J. Baxter, and N. Eberhard. 1992. Negative thyroid hormone control of human growth hormone gene expression is mediated by 3'-untranslated/3'-flanking DNA. *J. Biol. Chem.* 267: 15056–15063.
Identification of a novel zinc finger protein binding a conserved element critical for Pit-1-dependent growth hormone gene expression.

S M Lipkin, A M Näär, K A Kalla, et al.

*Genes Dev.* 1993, 7:
Access the most recent version at doi:10.1101/gad.7.9.1674