The Locus Control Region Activates Serpin Gene Expression through Recruitment of Liver-Specific Transcription Factors and RNA Polymerase II†

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The human serine protease inhibitor (serpin) gene cluster at 14q32.1 comprises 11 serpin genes, many of which are expressed specifically in hepatic cells. Previous studies identified a locus control region (LCR) upstream of the human α1-antitrypsin (α1AT) gene that is required for gene activation, chromatin remodeling, and histone acetylation throughout the proximal serpin subcluster. Here we show that the LCR interacts with multiple liver-specific transcription factors, including hepatocyte nuclear factor 3β (HNF-3β), HNF-6α, CCAAT/enhancer binding protein alpha (C/EBPα), and C/EBPβ. RNA polymerase II is also recruited to the locus through the LCR. Nongenic transcription at both the LCR and an upstream regulatory region was detected, but the deletion of the LCR abolished transcription at both sites. The deletion of HNF-3 and HNF-6 binding sites within the LCR reduced histone acetylation at both the LCR and the upstream regulatory region and decreased the transcription of the α1AT, corticosteroid binding globulin, and protein Z-dependent protease inhibitor genes. These results suggest that the LCR activates genes in the proximal serpin subcluster by recruiting liver-specific transcription factors and components of the general transcription machinery to regulatory regions upstream of the α1AT gene.

Many mammalian genes are organized into structurally related gene clusters that are expressed in a tissue- or stage-specific manner. These clusters are often controlled by complex transcriptional regulators called locus control regions (LCRs) (4). The serpin gene cluster at 14q32.1 is one such cluster (27). This ∼370-kb region contains 11 serpin genes that are organized into discrete proximal, central, and distal subclusters (27). The proximal subcluster contains four serpin genes, an α1-antitrypsin (α1AT, or SERPINA1) gene, an antitrypsin-related (SERPINA2) pseudogene, a corticosteroid binding globulin (CBG, or SERPINA6) gene, and a protein Z-dependent protease inhibitor (ZPI, or SERPINA10) gene (27). These genes are highly expressed in the liver, but they are repressed in most other cell types (23). The microcell-mediated transfer of human chromosome 14 from nonexpressing cells to expressing cells results in systematic serpin gene activation and chromatin remodeling of the entire locus into an expressing cell-typical chromatin state (34). This process provides an experimental system for studying the regulation of gene expression and chromatin structure within the serpin domain.

The activation of many hepatic genes is mediated by families of liver-specific transactivator genes, including those encoding hepatocyte nuclear factor 1 (HNF-1), HNF-3, HNF-4, HNF-6, and the CCAAT/enhancer binding protein (C/EBP). Each of these families contains several members. Furthermore, these transcription factors also regulate the expression of one another, thus forming a complex network that maintains the liver-specific transcription of albumin, α1AT, transthyretin, and α-fetoprotein genes and other hepatic genes (10). α1AT gene activation has been studied in detail previously, and HNF-1, HNF-4, HNF-3β, and HNF-6 have been shown to be involved (3, 35, 37). The data demonstrate that multiple liver-specific transcription factors are required for the activation of the α1AT gene.

More recently, homologous modifications of the chromosomal α1AT locus have identified an LCR that is required for cell-specific gene activation and chromatin remodeling throughout the proximal serpin subcluster. The deletion of five expression-associated DNase I-hypersensitive sites (DHSs) within an ∼8.0-kb genomic DNA segment upstream of the α1AT gene results in a mutant serpin allele that is completely refractory to cell-specific gene activation, DHS formation, and domain-wide histone acetylation (1, 22). Experiments with subdeletions within the 8.0-kb chromosomal segment indicated that the serpin LCR consists of multiple regulatory elements (22).

In this study, we investigated the mechanism of the serpin LCR functions. This investigation was accomplished by detailed mapping of histone acetylation and transcription factor loading at a variety of DNA sites within the proximal serpin subcluster. These sites included serpin gene promoters, intergenic regions, the serpin LCR, and an upstream regulatory region (URR) 20 to 25 kb upstream of the α1AT gene. Our results demonstrate that histones in these regions are hyperacetylated in expressing cells. However, the domain of histone acetylation is discontinuous, as hyperacetylation in the region between the α1AT and CBG genes was not observed. Significantly, the highest concentrations of liver-specific transcription factors, including HNF-3β, HNF-6α, C/EBPα, and C/EBPβ,
were found not at serpin gene promoters but at the DHS of kb −7.5 (relative to the α1AT start site) of the serpin LCR. The kb −20.8 DHS of the URR was also highly enriched with HNF-6α, C/EBPα, and C/EBPB, and the deletion of a 461-bp core element within the serpin LCR eliminated factor binding at the kb −20.8 URR site. These data suggest that the recruitment of transcription factors to the LCR and the URR is important for gene activation and chromatin remodeling in the proximal serpin domain. Interestingly, RNA polymerase II (Pol II) loading and nongenic transcription at these sites were also observed.

MATERIALS AND METHODS

Cell lines and culture conditions. F(14n)14 and F(461wt n)9 are independent ret hepatoma microcell hybrids that contain a single, wild-type copy of human chromosome 14, and R(h14n)6 is a rat fibroblast microcell hybrid that also contains a wild-type copy of human chromosome 14 (34). F(Δ50.0), F(Δ8.0)6, and F(Δ0.0.015) are independent microcell hybrid clones that contain a mutant human chromosome 14 [F(Δ50.0), F(Δ8.0)6, F(Δ0.0.015)] that lacks an 8.0-kb segment of genomic DNA just upstream of the hepatic α1AT gene promoter (22). The cells were grown in 1:1 Ham’s F12-Dulbecco’s modified Eagle’s medium with 5% fetal bovine serum and 500 μg of Geneticin/ml.

Generation of F(Δ461/−7.8) mutant cells. (i) Generation of mutant allele. The following primer pairs were used to generate the DNA homology segments for the p(Δ461/−7.8) targeting vector: a proximal arm forward primer, AGCACTAACATCCCCACATAAGACTTTTG. Amplified arms for homology segments were initially cloned into pCR2.1-TOPOII (Invitrogen). The primers contained either a SalI or an XhoI tail (underlined) that was used for cloning into the SalI or XhoI site of the pLAHL-PGK-DipA targeting vector (22). A unique PvuI site in the vector backbone was used to linearize the recombination sub-

(ii) DNA transfections. Stable and transient transfections were performed as described previously (6). Stable transfectants were selected in DT40-conditioned medium supplemented with fetal bovine serum, chicken serum, tryptone phosphate, and 1.5 mg of l-histidinol (Sigma/ml).

(iii) Excision of the histidinol selection cassette. Cells were transfected with a bicistronic plasmid encoding Cre recombinase and enhanced green fluorescent protein. After 24 h, >104 enhanced green fluorescent protein-positive cells were harvested using a Vantage SE turbo fluorescence-activated cell sorter. The pooled cells were grown in DT40-conditioned medium.

AQPCR analysis of serpin gene expression. Total RNA (totRNA) was harvested using TRIzol per the protocol of the manufacturer (Invitrogen). The cDNA was analyzed and quantitated using the Agilent 2100 bioanalyzer. Three micrograms of total RNA was treated with amplification-grade DNAse I (Invitrogen) according to the manufacturer’s instructions. cDNA was obtained from the treated totRNA by using the high-capacity cDNA archive kit per the protocol of the manufacturer (Applied Biosystems). Absolute quantitative PCR (AQPCR) was performed with 250 ng of cDNA by using intron-spanning primers and TaqMan MGB probes (Applied Biosystems) specific for human α1AT, CBG, and ZPI genes (see Table S1 in the supplemental material). Expression levels for each totRNA sample were normalized using GeNorm (40) and three internal control genes, the Dicer1 gene and the legumain and isoforms 1,3,4-triphosphate 5/6 kinase genes (catalog no. Hs00715991_m1 and Hs00365461_m1, respectively; Applied Biosystems).

RT assay for nongenic transcription. Reverse transcription (RT) reactions were performed as described previously (17), with modifications. Briefly, 5 μg of RNA was treated with RNase-free DNase for 15 min at 25°C. The RNA was then reverse transcribed using Superscript II TM and random hexamer primers according to the protocol of the manufacturer (Invitrogen). The cDNA was amplified by real-time QPCR. The difference (n-fold) in the amount of a given target sequence was determined by dividing the amount of the target sequence in the cDNA sample by the amount of the target sequence in genomic DNA. The relative level of transcription of a given target sequence was then obtained by normalizing the difference (n-fold) in the amount of the sequence with the difference (n-fold) obtained for primers within Dicer1 exon 21.

ChIP assays and data analysis. Histone acetylation and binding of Pol II, HNF-3β, and HNF-6α at specific genomic sites in vivo were studied by chromatin immunoprecipitation (ChIP) as described previously (14). Briefly, 2.5 × 107 cells were cross-linked with 1% formaldehyde for 10 min at room temperature, and chromatin was sheared to 500- to 1,000-bp fragments by sonication. The chromatin was preclayed by incubation overnight with protein asogare beads at 4°C. An aliquot of preclayed chromatin served as the input sample. The chromatin was subjected to reactions with specific antibodies and was precipitated using protein agarose beads. The eluted chromatin and the reserved input sample were reverse cross-linked by incubation at 65°C overnight. Purified DNA samples were analyzed by real-time QPCR using specific primers and TaqMan probes.

RESULTS

The α1AT and CBG genes reside in distinct chromatin sub-domains. As demonstrated previously, the deletion of an ~8.0-kb chromosomal segment containing the serpin LCR results in a mutant serpin allele that is refractory to cell-specific gene activation, fails to form expression-associated DHSs, and reduces histone acetylation throughout the proximal subcluster (1, 22). To determine whether this region comprises a continuous chromatin domain, we assayed levels of histone acetylation across the entire domain. ChIP experiments were performed using antibodies against histone H3 acetylated at lysine residues 9 and 14 and against all forms of acetyl-histone H4. The immunoprecipitated samples were tested by real-time QPCR using multiple ampiclons as depicted in Fig. 1. The level of enrichment of a sample with a particular DNA sequence was normalized according to the level of enrichment with sequences 5′ of the Goosecoid gene, a nonexpressed gene located ~380 kb from the α1AT gene toward the distal end of human chromosome 14 (27).

Averages of results from three independent experiments are shown in Fig. 2. In expressing F(14n)14 cells, peaks of histone H3 and H4 hyperacetylation at the kb −6.2 DHS of the serpin LCR, at the macrophage and hepatic promoters of the α1AT gene, and at the CBG and ZPI gene promoters were seen (Fig. 2), consistent with our previous data (1). Lower levels of his-
tone hyperacetylation at the kb −7.5, −5.3, and −4.3 DHSs within the LCR were detected, but the entire LCR was hyperacetylated to some degree. Notably, DNA sequences at kb −3.2, which is between the LCR core (kb −8.1 to −5.8) (22) and the α1AT gene promoters, were also hyperacetylated. This finding suggests that histone hyperacetylation in the region between the serpin LCR and the α1AT gene, which is the first gene of the proximal serpin subcluster (Fig. 1), is continuous. Histones at DHSs in the region from kb −20 to −24 were moderately (5- to 12-fold) acetylated. In contrast, histones in the kb −15.9 region were hypoacetylated in both expressing [F(14n)14] and nonexpressing [R(h14n)6] cells. This finding suggests that the URR at kb −24 to −20 and the serpin LCR at kb −8.4 to −0.3 are separated by a hypoacetylated chromatin zone.

The level of histone acetylation throughout the region in nonexpressing R(h14n)6 cells was generally lower than that in expressing F(14n)14 cells, which demonstrates the tissue specificity of the acetylated chromatin domain. One exception to this rule was the macrophage promoter of the α1AT gene, which showed constitutive hyperacetylation in both cell types. It is noteworthy that the macrophage promoter in both F(14n)14 and R(h14n)6 cells is also constitutively sensitive to DNase I (34). Levels of H3 and H4 acetylation in the KIAA1622 gene (Fig. 2), which is expressed in both cell types, were also similar.

To more extensively map the histone hyperacetylation domain, we designed and analyzed amplicons throughout the ~150-kb region (Fig. 1). This hypoacetylation suggests that the histone hyperacetylation domain within the proximal serpin subcluster is discontinuous. However, histone hyperacetylation throughout the domain requires the serpin LCR, as hyperacetylation in the region from kb −24 to +100 in F(Δ8.0)1 (Fig. 2) and F(Δ8.0)6 and F(Δ8.0)15 (data not shown) cells, which lack the serpin LCR, was not observed. This finding confirms the dominant function of the LCR in establishing an active chromatin domain in this ~125-kb region.

**Pol II is recruited to the LCR.** ChIP experiments were performed to determine the distribution of Pol II across the proximal serpin subcluster. As expected, no significant Pol II enrichment in nonexpressing R(h14n)6 cells was detected (Fig. 3A). In contrast, the α1AT gene promoter in F(14n)14 cells was highly enriched (~20-fold) with Pol II, consistent with the high level of expression of α1AT in this cell type (Fig. 3A). Moderate levels (~5-fold) of Pol II enrichment at the promoters of the CBG and ZPI genes, which are expressed at levels 50- to 100-fold lower than the α1AT gene, were also observed. Significantly, both the kb −7.5 and the kb −6.2 DHSs of the serpin LCR were enriched ~5- to 10-fold with Pol II (Fig. 3A). However, Pol II enrichment in the region upstream of the α1AT gene was discontinuous, as the α1AT gene promoter and the serpin LCR were enriched with Pol II but the regions between them were not (Fig. 3A, kb −5.3 and −4.3 amplicons). Similarly, the region just upstream of the LCR (Fig. 3A, kb −9.5 amplicon) was not enriched with Pol II. These results suggest that Pol II is specifically recruited to the kb −7.5 and −6.2 DHSs of the serpin LCR, which is consistent with the observation that core LCR activity lies within the 2.3-kb DNA segment between kb −8.1 and −5.8 (22). It is also possible that these sites may be enriched with Pol II by reduced rates of transcriptional elongation of nongenic transcripts (see below) in these regions. Pol II is also found at the macrophage pro-
moter of the α1AT gene, which is active at low levels in this cell type (Fig. 3A).

We demonstrated previously that α1AT and CBG expression decreases dramatically when the LCR is deleted (22). To test whether the deletion of the LCR affected the recruitment of Pol II to serpin gene promoters, ChIP assays with F(14n)14 cells, which lack the LCR, were performed. In these cells, Pol II recruitment to the α1AT, CBG, and ZPI gene promoters was abolished (Fig. 3A). Thus, the serpin LCR is important not only for recruitment to the neighboring α1AT gene promoter, but also for Pol II loading at the CBG and ZPI gene promoters, which are 67 and 100 kb away, respectively.

Nongenic transcription occurs at the LCR and upstream regions. To determine whether RNA transcripts were generated at the LCR and other regions of Pol II enrichment, RT-PCR experiments were performed. The enrichment of cDNA samples with specific target sequences was tested by real-time QPCR, and the levels of enrichment were normalized according to the levels of enrichment with sequences at exon 21 of the human Dicer gene, a constitutively expressed locus which is 700 kb toward the distal end of human chromosome 14. Normalization with levels of other gene sequences yielded similar results. In F(14n)14 cells, transcription at the LCR core was detected, with levels ~6 to 10% of those at Dicer exon 21 (Fig. 3B, ~7.5 and ~6.2). Notably, nongenic transcripts at the kb ~4.3 DHS and kb ~3.2 were also detected (Fig. 3B), although these regions were not significantly enriched with Pol II (Fig. 3A). This finding suggests that there may be progressive transcription between the LCR core and the α1AT gene promoter. Nongenic transcripts at the kb ~24.2, ~22.8, ~15.9, and ~9.5 sites were also detected (Fig. 3B). This finding suggests that nongenic transcripts in F(14n)14 cells are generated over a broad region. Nongenic transcription in R(h14n)6 cells was also detected, but the levels were ~3- to 5-fold lower than those in F(14n)14 cells. Nongenic transcripts in F(Δ8.0/−8.1)1 cells were not detected (Fig. 3B), indicating that the nongenic transcription we observed is LCR dependent.

The serpin LCR functions as a recruitment center for transcription factors. Hepatic genes are regulated by a set of liver-specific transcription factors. For example, HNF-3 and HNF-6 bind to the transthyretin, α1AT, and α1AT gene promoters in vitro, and this binding activates linked reporter gene expression in transfection assays (3, 35). Transcriptional activation of the α1AT gene also requires the binding of HNF-1α and HNF-4α (13, 33) to the α1AT gene promoter/enhancer (37). ChIP studies have shown that HNF-6 and C/EBPα and
C/EBPβ are bound at the mouse HNF-3β gene promoter in vivo (44).

LCRs and enhancers generally activate target genes by recruiting transcriptional activators. To determine whether the serpin LCR functions in this way, we assayed the binding of different liver-specific transactivators to the locus by using ChIP. In these experiments, amplicons 5’ of the Goosecoid gene or within Dicer1 exon 21 served as negative controls, and an amplicon within the rat HNF-3β gene promoter was the positive control. Results from three independent experiments are summarized in Fig. 4. As expected, the rat HNF-3β gene promoter in F(14n)14 cells was highly enriched with HNF-3β, HNF-6α, C/EBPα, and C/EBPβ (Fig. 4, insets). Within the human serpin locus, the kb 7.5 DHS of the LCR and the CBG gene promoter were highly enriched with HNF-3β (Fig. 4A). Low-level HNF-3β enrichment at the kb 20.8 DHS, the kb 6.2 DHS, and the hepatic a1AT gene promoter was also found (Fig. 4A). In contrast, HNF-6α was found only at the kb 7.5 and 20.8 DHSs (Fig. 4A). The kb 7.5 and 6.2 DHSs of the serpin LCR were highly enriched with C/EBPα and C/EBPβ, and the kb 20.8 DHS was less so (Fig. 4B). These results suggest that the LCR is a center for the recruitment of liver-specific transcription factors. Consistent with this view, the deletion of the LCR in F(Δ8.0/−8.1)1 cells abolished the binding of all four transcription factors at other sites in the proximal subcluster (Fig. 4).

Establishment of the histone acetylation domain requires HNF-3β and HNF-6α binding to the LCR. To test the function of HNF-3β and HNF-6α bound at the kb 7.5 DHS of the LCR, we generated a mutant serpin allele (Δ461/−7.8) in which the core region of the kb 7.5 DHS was deleted specifically (Fig. 5A). Four independent F(Δ461/−7.8) clones carrying this mutant allele showed reduced levels of expression of the α1AT (~4-fold), CBG (~4-fold), and ZPI (~3-fold) genes as assessed by AOPCR (Fig. 5B).

ChIP experiments were carried out to define the chromatin landscape of the Δ461/−7.8 mutant allele. As the amplicon at kb 7.5 was deleted in this mutant allele, a new primer-probe set at kb 7.3 was used to monitor this region. The binding of HNF-3β and HNF-6α at the kb 7.5 DHS in the URR was dramatically reduced by the 461-bp deletion.

FIG. 3. Pol II distribution and nongenic transcription across the proximal serpin locus. (A) ChIP experiments were performed as described in the legend to Fig. 2, except that antibody to Pol II was used. The relative enrichment is expressed as n-fold. (B) RT-PCR was performed to detect the level of nongenic transcription. The level of nongenic transcription at each of the amplicons is depicted as the percentage of the transcription from the constitutively expressed Dicer1 exon 21 amplicon. Enrichment data for the LCR are absent in F(Δ8.0/−8.1)1 cells as this is the region deleted by the Δ8.0/−8.1 modification. α1ATpM, macrophage α1AT gene promoter; α1ATpH, hepatic α1AT gene promoter; CBGp, CBG gene promoter.
The binding of C/EBPα and C/EBPβ to the LCR and the upstream region was modestly affected (Fig. 6B).

Pol II loading at the hepatic α1AT gene promoter and the kb -6.2 DHS of the LCR was decreased two- to threefold in the F(Δ461/Δ7.8) mutant relative to that in the wild type, but Pol II loading at the macrophage α1AT gene promoter or the CBG or ZPI gene promoter was not affected (Fig. 6C). This finding suggests that the recruitment of Pol II to the LCR core requires HNF-3β and HNF-6α binding at this site. Nongenic transcription in the F(Δ461/Δ7.8) mutant was largely unaffected (Fig. 6D).

Histone acetylation across the entire proximal serpin subcluster was significantly decreased (Fig. 7). These data demonstrate that the establishment of a domain of histone hyperacetylation within the proximal subcluster requires HNF-3β and HNF-6α binding to the kb -7.5 DHS of the serpin LCR.

**DISCUSSION**

LCRs were originally defined based on their ability to induce integration site-independent and copy number-dependent expression of linked genes in transgenic mice (18). These gene activation phenomena are generally correlated with alterations in chromatin structure, for example, increased nuclease accessibility and posttranslational histone modifications (25). It has therefore been assumed that LCRs provide a dominant chromatin-opening activity (4) that is required for gene expression. Consistent with this view, the deletion of one DHS in the human growth hormone (hGH) LCR results in decreased transgene expression and the loss of histone acetylation in an ~32-kb domain (11). Moreover, the deletion of the 5′ HS3 core region of the β-globin LCR affects DNase I accessibility and reduces histone acetylation throughout the locus in transgenic mice (7). However, deletions of the mouse chromosomal
β-globin LCR reduce β-globin gene expression, but the chromatin structure of the locus is largely unaffected (2, 36). These results suggest that the β-globin LCR affects gene expression by a mechanism that functions downstream of activation-specific alterations in chromatin structure, possibly at the level of RNA elongation (36). In marked contrast, chromosomal deletions in the serpin LCR abolish gene expression, DHS formation, and histone acetylation across the entire proximal serpin subcluster, a genomic region of ~130 kb (1, 22). These results suggest that LCRs function in a variety of different ways in vivo.

Although the α1AT, CBG, and ZPI genes are expressed at different levels, the transcription of all three genes is LCR dependent. In liver cells, α1AT mRNA is abundant, constituting ~1% of Pol II transcripts (5), while the CBG and ZPI genes are expressed at much lower levels (unpublished data). Levels of histone acetylation also vary across the locus, with the highest levels at the hepatic α1AT gene promoter, the serpin LCR, and the ZPI gene promoter. However, as shown both in a previous study (1) and in the present study, the zone of histone hyperacetylation across the proximal serpin subcluster is discontinuous, as there are areas of hypoacetylation between genes. This finding suggests that the differential transcription of the α1AT gene versus the CBG and ZPI genes may be due, at least in part, to the residence of these genes in different chromatin subdomains. Further studies of the region using high-density probe arrays may help to resolve this issue.

Whether proximity to the LCR and/or the presence of intervening matrix attachment regions also affects gene activity in the region is presently unknown. In any event, the activation domain(s) that includes the proximal serpin subcluster does not extend to the kb –45 DHS upstream of the locus or to the KIAA 1622 gene downstream.

There is increasing evidence that the activation of target genes through distal enhancer and/or LCR elements is mediated by the recruitment of transcriptional machinery components. For example, an enhancer 25 kb away from the pDβ1 gene in the T-cell receptor β locus binds Pol II (38). Moreover, Pol II is recruited to the upstream enhancer of the androgen-responsive prostate-specific antigen gene in a hormone-dependent but promoter-independent manner (21). Pol II is also found at DHSs within the β-globin LCR in both mouse and human (15, 41). More recently, it was shown that the β-globin genes localize to active Pol II foci in expressing cells and that the β-globin LCR is required for colocalization (28, 30). These findings are consistent with the notion that during transcription, genes are localized to discrete nuclear sites of concentrated RNA polymerase. In the present study, we found that not only serpin gene promoters but also the serpin LCR is enriched with Pol II. Furthermore, deletions of the LCR abolished Pol II binding to the hepatic α1AT gene promoter, suggesting that the promoter alone is not sufficient for the recruitment of Pol II at its native chromosomal site. LCR deletions also abolished Pol II binding near the CBG and ZPI gene promoters, which are 67 and 98 kb away, respectively. These results suggest that the serpin LCR plays a fundamental role in recruiting Pol II to the locus, and genes both nearby and at a distance require Pol II binding at the LCR for their subsequent regulation.

**FIG. 6.** The region excised by the Δ461/−7.8 modification is crucial for the recruitment of transcription factors. ChIP experiments were performed (as described in the legend to Fig. 2) with F(14n)14 and F(Δ461/−7.8) cells with antibodies to HNF-3β and HNF-6α (A), C/EBPα and C/EBPβ (B), and Pol II (C). The relative enrichment is expressed as n-fold. (D) Levels of nongenic transcription in F(14n)14 and F(Δ461/−7.8) cells were measured (as described in the legend to Fig. 3). α1ATpH, hepatic α1AT gene promoter; α1ATpM, macrophage α1AT gene promoter; CBGp, CBG gene promoter; rHNF3βpro, rat HNF-3β gene promoter; Dicer EX, Dicer1 exon 21.
The core region of the serpin LCR is a 2.3-kb DNA segment that extends from kb −8.1 to −5.8 (22). This region includes two expression-associated DHSs, at kb −7.5 and −6.2. Deletions in this region affect serpin gene expression and DHS formation across the locus (22), and histone hyperacetylation is also affected by the 2.3-kb deletion (unpublished data). ChIP experiments demonstrated that the kb −6.2 DHS within this region was highly enriched with acetylated histones and that C/EBPα and C/EBPβ were also bound at this site. Significantly, the kb −7.5 DHS was highly enriched with HNF-3β, HNF-6α, C/EBPα, and C/EBPβ. The deletion of a 461-bp core element of the kb −7.5 DHS abolished HNF-3β and HNF-6α binding in this region and reduced histone H3 and H4 acetylation across the locus. These results suggest that DNA sequences in the deleted 461-bp segment may serve as a platform for the recruitment of histone acetyltransferases to the locus through HNF-3β and HNF-6α.

The formation of expression-associated DHSs in the region from kb −25 to −20 upstream of the α1AT gene is strongly affected by LCR deletions (22). Here we show that factor recruitment in this region also requires the serpin LCR. For example, the kb −20.8 DHS is enriched with C/EBPα, C/EBPβ, and HNF-6α, but both DHS formation and factor binding at this site are abolished by the 8.0-kb LCR deletion. This finding raises the possibility that the binding of these factors to the kb −20.8 DHS may actually reflect a physical interaction between the LCR and this site by the formation of a chromatin loop (39). Alternatively, it may be due simply to the inaccessibility of sequences in this upstream region to trans-acting factors in the absence of the LCR. An analysis of the 461-bp-deletion mutant allowed us to distinguish between these possibilities. When the kb −7.5 core region, the region most highly enriched with HNF-3β and HNF-6α, is deleted, the kb −20.8 DHS still forms (unpublished data) and this region is enriched with C/EBPα but HNF-6α is no longer bound to this site. Thus, HNF-6α binding at kb −20.8 requires HNF-6 and/or HNF-3 binding at the kb −7.5 DHS of the LCR. Furthermore, there are no HNF-6 binding sites at kb −20.8 that can be recognized in silico, although there are candidate C/EBP binding sites at both kb −20.8 and kb −7.5. As C/EBPα and C/EBPβ can form homo- and heterodimers through leucine zipper sequences at their C termini (16), the serpin URR and LCR may contact each other through dimer formation between C/EBP factors.

The deletion of the URR significantly decreases the expression of the CBG gene but only modestly affects α1AT gene expression (unpublished data), suggesting that the differential activation of α1AT and CBG genes requires distinct combinations of regulatory elements. Interestingly, we found that the
URR and the CBG promoter are enriched with HNF-6α and HNF-3β, respectively. It has been shown previously that HNF-3β and HNF-6 can physically interact (31). This finding suggests that the URR may contact the CBG gene promoter through this interaction, which may be an intermediate step during CBG gene activation by the LCR.

Our findings that Pol II is recruited to the LCR and that nongenic transcripts are generated in the LCR are consistent with predictions of the transcription factory model (43). As proposed, the model suggests that if transcribed regions are being sequestered in Pol II foci, both regulatory elements and target genes will be in proximity to one another. Our observation that transcription factors bound at the LCR may also bind to the region from kb −24 to −20 implies that these two regions may interact physically. Studies of interactions between regulatory elements and target serpin genes, as well as colocalization studies of these DNA sequences, should provide information that will be important for understanding long-range gene control.

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