ALY, a context-dependent coactivator of LEF-1 and AML-1, is required for TCRα enhancer function

Laurakay Bruhn, Audrey Munnerlyn, and Rudolf Grosschedl

Howard Hughes Medical Institute and Departments of Microbiology and Biochemistry, University of California, San Francisco, California 94143-0414 USA

LEF-1 is a transcription factor that participates in the regulation of the T-cell receptor α (TCRα) enhancer by facilitating the assembly of multiple proteins into a higher order nucleoprotein complex. The function of LEF-1 is dependent, in part, on the HMG domain that induces a sharp bend in the DNA helix, and on an activation domain that stimulates transcription only in a specific context of other enhancer-binding proteins. With the aim of gaining insight into the function of context-dependent activation domains, we cloned ALY, a novel LEF-1-interacting protein. ALY is a ubiquitously expressed, nuclear protein that specifically associates with the activation domains of LEF-1 and AML-1 (CBFα2, PEBP2αB), which is another protein component of the TCRα enhancer complex. In addition, ALY can increase DNA binding by both LEF-1 and AML proteins. Overexpression of ALY stimulates the activity of the TCRα enhancer complex reconstituted in transfected nonlymphoid HeLa cells, whereas down-regulation of ALY by anti-sense oligonucleotides virtually eliminates TCRα enhancer activity in T cells. Similar to LEF-1, ALY can stimulate transcription in the context of the TCRα enhancer but apparently not when tethered to DNA through a heterologous DNA-binding domain. We propose that ALY mediates context-dependent transcriptional activation by facilitating the functional collaboration of multiple proteins in the TCRα enhancer complex.

[Key Words: LEF-1; AML-1(CBFα2,PEBP2αB); TCRα enhancer; ALY; transcription activation]

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was replaced with the DNA-binding domain of LexA (Giese and Grosschedl 1993). We screened an embryonic day 10 mouse cDNA library in which the cDNA inserts were isolated that conferred reporter gene activation only in combination with the LEF-1–LexA fusion construct but not with three control LexA fusion proteins, lamin, Ste3, and Fus1. Four of these contained identical inserts of 230 bp and one contained a longer 310-bp insert from the same gene.

The 310-bp insert was used as a probe to isolate full-length 1.1-kb cDNA clones that encode 255 amino acids followed by stop codons in all three reading frames (Fig. 1). On the basis of its ability to interact with LEF-1 and AML-1 (see below), we refer to the protein product of this cDNA as ALY (Ally of AML-1 and LEF-1). The size of the open reading frame (ORF) of the ALY cDNA corresponds well with the 32-kD molecular mass of recombinant ALY polypeptide transcribed and translated in vitro (see Fig. 3B, below), and with the size of the endogenous protein that can be detected on immunoblots by polyclonal antibodies directed against ALY (data not shown).

Amino acid sequence comparison of ALY with proteins in the GenBank database revealed no identities but did reveal potential homology with RNA-binding proteins. In particular, the protein contains a putative RNP (ribonucleoprotein) RNA recognition motif and Arg/

**Results**

**Cloning of cDNAs encoding LEF-1-interacting proteins**

To identify proteins that interact and functionally collaborate with LEF-1, we used the yeast two-hybrid assay (Fields and Song 1989). For the screen, we used a LEF–LexA fusion protein in which the HMG domain of LEF-1 was replaced with the DNA-binding domain of LexA (Giese and Grosschedl 1993). We screened an embryonic day 10 mouse cDNA library in which the cDNA inserts are fused to sequences encoding a VP16 transcription activation domain (Hollenberg et al. 1995). From ~2.7 × 10^6 yeast transformants, eight library plasmids were isolated that conferred reporter gene activation in vivo (Giese et al. 1995). Studies of the TCRα enhancer have revealed two distinct context-dependent functions of LEF-1, an architectural function and a transcription activation function. The architectural role of LEF-1 is mediated by the HMG domain that, by inducing a directed DNA bend in the enhancer, is thought to facilitate interactions between factors bound at the ATF/CREB-binding sites and AML: Ets-binding sites, which are necessary for the formation of a stable enhancer complex (Giese et al. 1995).

The context-dependent transcription activation functions of LEF-1 are mediated by regions outside of the carboxy-terminally located HMG domain. Analysis of chimeric proteins in which the HMG domain of LEF-1 was replaced by heterologous DNA-binding domains delineated a context-dependent activation domain (CAD) between residues 99 and 217 of LEF-1 (Carlsson et al. 1993; Giese and Grosschedl 1993). This region differs from typical activation domains in that it is unable to stimulate transcription from multimerized binding sites but rather, activates transcription when tethered to DNA in the context of the natural arrangement of factor-binding sites in the TCRα enhancer or the HIV enhancer (Carlsson et al. 1993; Giese and Grosschedl 1993; Sheridan et al. 1995).

A molecular explanation for the function of the context-dependent activation domain of LEF-1 remains unclear. One simple model is that these regions interact, directly or indirectly, with other enhancer-bound factors. Previous attempts to identify direct LEF-1 interactions with other known TCRα enhancer-bound factors have been unsuccessful (Giese et al. 1995). Here we used a yeast two-hybrid screen to isolate LEF-1-interacting proteins that may contribute to its transcription activation functions. We have identified a novel, nuclear protein that interacts with two TCRα enhancer-bound factors, LEF-1 and AML-1. We show that this factor is required for TCRα enhancer function in T cells and plays a role in regulating the activity of this multiprotein enhancer complex.

**Figure 1.** Nucleotide sequence of the murine ALY cDNA and predicted amino acid sequence of the encoded protein. The polypeptide encoded by the 310-bp two-hybrid isolate, termed ALYANC, is underlined. This underlined region encompasses the putative RNP (ribonucleoprotein) RNA-binding motif. The putative RNP1 (RNP octamer) and RNP2 (RNP hexamer) sequences are underlined in bold.

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**Table 1.** Nucleotide sequence of the murine ALY cDNA and predicted amino acid sequence of the encoded protein. The polypeptide encoded by the 310-bp two-hybrid isolate, termed ALYANC, is underlined. This underlined region encompasses the putative RNP (ribonucleoprotein) RNA-binding motif. The putative RNP1 (RNP octamer) and RNP2 (RNP hexamer) sequences are underlined in bold.
Gly-rich regions in the amino and carboxyl termini that are loosely homologous to RGG RNA-binding motifs (for review, see Burd and Dreyfuss 1994). Biochemical and structural studies of RNP domains have shown that they are sufficient to mediate RNA binding and that three highly conserved aromatic amino acids found in the RNP1 (RNP octamer) and RNP2 (RNP hexamer) sequences are involved in critical stacking interactions with RNA bases (for review, see Nagai et al. 1995). However, the putative RNP1 and RNP2 sequences of ALY (Fig. 1, bold underlined) contain only a single aromatic amino acid. Moreover, preliminary studies demonstrated that ALYANC, which includes the putative RNP domain, is not capable of binding RNA at any detectable level, although very weak nonspecific binding to RNA and DNA was observed with full-length ALY protein (data not shown). Thus, ALY may represent a protein that is related to RNA-binding proteins and has some propensity to interact with nucleic acids, but in which the RNP domain may perform other functions.

**ALY encodes a ubiquitously expressed, nuclear protein**

To examine the expression pattern of ALY, we analyzed poly(A)+ RNA from various mouse tissues and cell lines for the presence of transcripts that hybridize with an ALY cDNA probe. A single 1.1-kb ALY transcript was detected at varying levels in all tissues and cell lines examined suggesting that ALY may be expressed ubiquitously (Fig. 2). The subcellular localization of ALY was examined by indirect immunofluorescence microscopy of COS7 cells that had been transfected transiently with a plasmid expressing a T7 epitope-tagged version of ALY. This approach was necessary because our anti-ALY antiserum failed to recognize native protein (data not shown). By comparing the immunofluorescence staining of T7-ALY with DAPI staining of DNA, we detected ALY primarily in the nucleus of transfected cells (Fig. 2C,D). We conclude from these results that ALY is a ubiquitously expressed nuclear protein.

**ALY interacts with LEF-1 and AML proteins**

To establish that full-length ALY can interact directly with LEF-1, we performed in vitro association assays with purified recombinant glutathionine S-transferase (GST)-ALY fusion protein. 35S-Labeled LEF-1 protein was bound by immobilized GST–ALY but not GST alone (Fig. 3A, lane 2). Because ALY contains regions with some homology to RNA-binding domains, it was possible that the ALY:LEF-1 interaction was mediated indirectly by nucleic acids. However, inclusion of ethidium bromide in the GST association experiments, which has been shown to disrupt potential nonspecific interactions caused by contaminating nucleic acids (Lai and Herr 1992), and preincubation of GST–ALY and the 35S-labeled proteins with RNase did not affect the association of GST–ALY with LEF-1 (data not shown).

Previous studies suggested that the function of LEF-1, in particular the CAD, could involve interactions with other transcriptional regulators of the TCRα enhancer (Travis et al. 1991; Carlsson et al. 1993; Giese and Groschedel 1993). We tested the ability of GST–ALY to interact with various 35S-labeled transcription factors including those that are known to bind the TCRα enhancer (Fig. 3A). Strikingly, in addition to LEF-1 (lane 2), GST–ALY interacted specifically with AML-3 (lane 3) and AML-1 (see below). In contrast, GST–ALY did not detectably associate with CBFβ (lane 4), the heterodimeric binding partner of AML proteins (Ogawa et al. 1993b; Wang et al. 1993) and failed to interact with Ets-1 or an amino-terminally truncated derivative of Ets-1 (lanes 7,8), which was shown to interact strongly with AML-3 (PEBP2αA, CBFα1; Giese et al. 1995). Members of the ATF/CREB family including ATF-2, CREB-B, and c-Jun, also failed to interact strongly with GST–ALY, although a very low level of binding of CREB-B and c-Jun was observed (lane 6; data not shown). This weak interaction of ALY with CREB-B was unaffected by phosphorylation of CREB-B by protein kinase A (data not shown), which was found previously to regulate its association with the
coactivator CREB-binding protein (CBP) [Kwok et al. 1994]. Two other transcription factors—Myb, which collaborates with AML proteins to activate the TCRβ enhancer [Hernandez-Munain and Krangel 1995], and Pit-1, which contains a functional domain with homology to the CAD of LEF-1 [Holloway et al. 1995]—also failed to associate detectably with GST-ALY (Fig. 3A, lanes 1,9). These results were confirmed by reciprocal associations that demonstrated that 35S-labeled ALY could bind to immobilized GST–LEF-1 and GST–AML-3, but not significantly to GST–CBFβ, GST–Ets-1, or GST–ATF2 [data not shown].

To examine whether ALY has the potential to multimerize, we incubated 35S-labeled ALY with immobilized GST–ALY and GST–ALYΔNC (Fig. 3B). Association was detected only with GST–ALY indicating that ALY can multimerize primarily through regions at the amino or carboxyl terminus of the protein. As for the interaction of ALY with LEF-1 and AML proteins, RNase treatment of the protein preparations did not affect the associations (Fig. 3B; data not shown). The finding that ALY may function as a multimer could allow, in principle, for simultaneous interactions between ALY and multiple factors in the enhancer complex even if the interactions with different factors were mediated by the same region of ALY.

**Mapping the protein domains required for interaction of ALY with LEF-1 and AML-1**

To begin to map the protein domains required for the ALY:LEF-1 interaction, we tested the ability of GST–ALY and GST–ALYΔNC to interact with various amino- and carboxy-terminally truncated versions of LEF-1 (Fig. 4). Amino-terminal truncation of residues between 99 and 166, which coincide with the CAD of LEF-1 [Carlsson et al. 1993; Giese and Grosschedl 1993], completely abrogated binding to GST–ALYΔNC and reduced binding to GST–ALY 5- to 10-fold. A form of LEF-1, lacking residues downstream of residue 243, showed increased
AML proteins, GST-ALY bound with similar efficiency residues carboxy-terminal to 287 of AML-1 decreased the function as a transcription activator both at the TCRβ nal half of AML proteins is required for its ability to ALY. Thus, the amino- or carboxy-terminal regions of ALY may mediate interactions with the HMG domain.

Figure 4. ALY interacts with the context-dependent activation and HMG domains of LEF-1. Schematic diagrams of LEF-1 and deletion derivatives (top). The β-catenin-binding domain (BD), context-dependent activation domain (CAD), and high mobility group (HMG) domain are boxed. Association assays between in vitro-translated 35S-labeled LEF-1 polypeptides [lanes 1–5, 10% of input] and GST-ALYANC [lanes 6–10] or GST–ALY [lanes 11–15] immobilized on glutathione–agarose beads. Similar molar amounts, normalized for the number of 35S-labeled methionines, of labeled full-length and truncated versions of LEF-1 were used in the association reactions. The input and the bound proteins were analyzed by SDS-PAGE and visualized by autoradiography. The amounts of 35S-labeled proteins bound to GST, GST-ALYANC, and GST–ALY were quantified from the association experiment shown at bottom using a PhosphorImager (Molecular Dynamics) and are presented as percentages relative to full-length LEF-1, which was designated as 100% (top). Similar results were obtained in three independent association experiments.

Figure 5. ALY interacts with the transcription activation domain of AML-1. Schematic diagrams of AML-1 and deletion derivatives [top]. The Runt domain that mediates DNA binding and interaction with CBFβ, Ets-1, and C/EBP is represented by a solid rectangle. Association assays between in vitro-translated 35S-labeled AML-1 polypeptides [lanes 1–4, 10% of input] and GST–ALYANC [lanes 5–8] or GST–ALY [lanes 9–12] immobilized on glutathione agarose beads. Similar molar amounts, normalized for the number of 35S-labeled methionines, of labeled full-length and truncated versions of AML-1 were used in the association reactions. The input and bound proteins were analyzed by SDS-PAGE and visualized by autoradiography. The amounts of 35S-labeled proteins bound to GST, GST–ALYANC, and GST–ALY were quantified from the association experiment shown at bottom using a PhosphorImager (Molecular Dynamics) and are presented as percentages relative to full-length AML-1, which was designated as 100% (top). Similar results were obtained in three independent association experiments.

ALY augments DNA binding by full-length LEF-1 and AML proteins

To examine whether ALY can interact with DNA-bound LEF-1 or AML proteins, we used an electrophoretic mobility-shift assay with the 98-bp TCRα enhancer probe.
Incubation of recombinant purified LEF-1 with high concentrations of purified GST–ALY yielded a complex that migrated with a slower mobility than that formed with LEF-1 alone, consistent with the formation of a ternary ALY:LEF-I:DNA complex (Fig. 6A, lane 3). No slower migrating complex was detected when GST–ALY was coincubated with the HMG domain (Fig. 6A, lane 7). In addition, GST–ALY had no detectable DNA-binding activity by itself in this assay (Fig. 6A, lanes 10–12). Notably, ALY increased DNA binding approximately fivefold by full-length LEF-1 and the HMG domain, both of which interact with ALY. In contrast, GST–ALYΔNC, at a 10-fold higher concentration, had no effect on DNA binding by LEF-1 or the HMG domain (data not shown). A recombinant purified hexa-histidine-tagged version of ALY had similar effects as GST–ALY on DNA-binding by LEF-1 (data not shown).

The ternary ALY:LEF-1:DNA complex formed with the 98-bp TCRα enhancer probe but not with a 50-bp TCRα enhancer probe lacking sequences upstream of the LEF-1 site (Fig. 6A; data not shown). This observation raised the possibility that the formation of a ternary complex may require a particular length or sequence of DNA. In quantitative DNase I protection assays, in which we confirmed the effect of ALY on increasing the efficiency of DNA binding by LEF-1, we did not detect differences in the footprint pattern of the LEF-1:TCRα(98 bp) complex in the presence or absence of ALY (Fig. 6B). Thus, this putative ALY:DNA interaction may not involve a specific sequence or may be unstable.

![Figure 6](https://genesdev.cshlp.org)
To examine whether ALY has an effect on DNA binding by AML proteins, we incubated the 98-bp TCRα enhancer probe with purified recombinant AML-3, which could be obtained from bacteria in soluble form. A complex that migrated with a slower mobility than the AML-3:DNA complex was detected (Fig. 6C). In the ALY:AML-3:DNA ternary complex, DNA binding was increased ~20-fold. In contrast, coincubation of ALY with the Runt domain of AML-3 increased DNA binding less than twofold and did not result in formation of a detectable ternary complex (Fig. 6C). The Runt domain binds DNA more efficiently than full-length AML-3 protein but does not interact with ALY in the GST association assay. Therefore, the association of AML proteins with ALY may antagonize the function of an inhibitory domain, consistent with the previous observations that carboxy-terminally truncated versions of AML-1 bind DNA with higher affinity than full-length protein (Bae et al. 1994; Tanaka et al. 1995).

**ALY specifically stimulates the activity of the TCRα enhancer**

Previously we have found that expression of LEF-1 and the lymphoid-specific factors AML (CBFα, PEBP2α) and Ets-1 confers high levels of TCRα enhancer activity in nonlymphoid HeLa cells (Giese et al. 1995). Given the lower ALY mRNA levels in HeLa cells compared to most T-cell lines (see Fig. 2B), we used the reconstituted HeLa cell system to determine the effect of ALY overexpression on TCRα enhancer activity. For these and other transfection experiments, we used AML-1, which is expressed in T cells much more abundantly than AML-3 (Satake et al. 1995). Overexpression of ALY resulted in a dose-dependent increase in the levels of TCRα-CAT expression with a maximum of fivefold (Fig. 7A). ALY alone or the truncated ALYΔNC in combination with the lymphoid-specific factors had no effect on TCRα enhancer activity.

To determine whether the ALY-mediated increase in reporter gene expression was specific for the TCRα enhancer, we tested three other reporter gene constructs controlled by the cytomegalovirus (CMV), Rous sarcoma virus (RSV), or SV40 viral enhancer/promoter regions, which are activated by endogenous proteins in HeLa cells. In addition, we examined the effect of ALY overexpression on the activity of two reporter genes stimulated by exogenously expressed AP1 or Gal-4–VP16. In these experiments, we transfected minimal amounts of reporter and effector plasmids to ensure that potential ALY-mediated effects on reporter gene activity could be observed. Except for a minor, twofold effect on the SV40-controlled reporter gene, the expression levels of all other reporters were unaffected by overexpression of ALY (Fig. 7B).

**ALY is required for TCRα enhancer function in T cells**

Although overexpression of ALY resulted in a significant increase in TCRα enhancer activity in HeLa cells, overexpression in T cell lines conferred negligible increases in TCRα expression (L. Bruhn, unpubl.). This could reflect the higher levels of endogenous ALY transcript present in most T cell lines (see Fig. 2B). To determine the contribution of ALY to TCRα enhancer activity in T cells in which the enhancer is naturally functional, we attempted to decrease the levels of endogenous ALY protein using C-5 propyne pyrimidine-substituted phosphorothioate anti-sense oligonucleotides (Wagner et al. 1993). Cotransfection of an ALY anti-sense oligonucleo-

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**Figure 7.** ALY stimulates specifically the activity of the TCRα enhancer. (A) Full-length ALY stimulates TCRα enhancer activity reconstituted in nonlymphoid cells. HeLa cells were transfected with expression plasmids for AML-1 (200 ng), Ets-1 (200 ng), and LEF-1 (100 ng) alone or with different amounts of expression plasmids for ALY or ALYΔNC together with 250 ng of a TCRα reporter plasmid. The reporter plasmid (top) contains the minimal 98-bp TCRα enhancer linked to the minimal fos promoter (−56 to +109), and the chloramphenicol acetyltransferase (CAT) gene. For these and subsequent transfection assays, luciferase or CAT activity was normalized to the activity of an RSV-β-gal expression plasmid as an internal transfection standard. Fold activation was quantitated relative to CAT activity from cells transfected with the reporter alone, which was designated as 1. (B) Effects of ALY on other reporter constructs. HeLa cells were transfected with various reporter and effector plasmids in the absence or presence of 500 ng of ALY expression plasmid. Data from TCRα-CAT with LEF-1, Ets-1, and AML-1 is duplicated from A. The amounts of other reporter and effector plasmids were as follows: 500 ng of CMV-CAT, RSV-β-gal, or SV40-CAT reporter constructs, 2 μg of AP1–luciferase plus 250 ng each RSV-c-Jun and RSV-c-Fos, 2 μg of Gal4–luciferase plus 250 ng RSV-Gal4–VP16.
ALY into Jurkat T cells and examined the levels of T7-epitope-tagged ALY in whole cell extracts by immunoblot analysis. Strikingly, the expression of T7-ALY was decreased in a dose-dependent manner with the ALY anti-sense oligonucleotide but was unaffected by similar concentrations of the shuffled control oligonucleotide (Fig. 8C). Thus, ALY plays an important role in the regulation of the TCRα enhancer.

Figure 8. Down-regulation of ALY expression in T cells abrogates TCRα enhancer function. Jurkat T cells were transfected transiently with 1 μg of TCRα-CAT (A), RSV-βgal (B), or CMV-T7-ALY (C), together with increasing concentrations of ALY anti-sense or shuffled oligonucleotides. The activities of the reporters in the presence of different concentrations of oligonucleotides were calculated relative to that with no oligonucleotide, which was designated as 100%. Whole cell extracts of cells transfected transiently with T7-ALY were analyzed by immunoblot using T7 monoclonal antibody. Data in A and B are the average of at least two transfections carried out in duplicate; error bars represent standard error of the mean.

Transcriptional activation by ALY is dependent on multiple TCRα enhancer-binding proteins

One simple model of ALY function is that this protein increases the activity of the TCRα enhancer by providing the enhancer complex with a transcription activation domain through binding to LEF-1. To explore this possibility, we tested whether ALY is capable of activating transcription when tethered to DNA by the Gal4 DNA-binding domain. ALY–Gal4 did not confer activation from multimerized Gal4 sites in transfected T cells and likewise, a LexA–ALY fusion protein did not activate expression of a reporter containing multimerized LexA sites (Fig. 9A; data not shown). In addition, neither ALY–Gal4 nor LexA–ALY fusion proteins significantly activated expression of TCRα enhancer constructs in which the LEF-1 binding site was replaced with a Gal4 or LexA site (Fig. 9B, data not shown). Efficient expression and DNA binding of the fusion proteins was confirmed by electrophoretic mobility-shift assays with nuclear extracts from COS7 cells transfected with these expression plasmids (Fig. 9C). These data suggest that ALY lacks a typical activation domain and probably does not stimulate TCRα enhancer function by recruitment of a transcription activation domain to the enhancer complex.

The findings that ALY can form multimers and can interact with both LEF-1 and AML proteins raised the possibility that it may serve to functionally bridge these factors within the TCRα enhancer complex. According to this view, the function of ALY in the reconstituted HeLa system would be dependent on both LEF-1 and AML-1. Coexpression of ALY with any two of the three lymphoid-specific factors stimulated the activity of TCRα four- to fivefold, although the levels of reporter expression were ~10-fold lower than those observed with all three factors (cf. Figs. 7A and 9D). In contrast, no significant ALY stimulation was observed with individual lymphoid-specific factors, with the exception of a twofold ALY stimulation with Ets-1. On basis of the presence of AML-2 [CBFo3, PEBP2αC] in all cell types, this modest increase of TCRα-CAT expression could be attributable to a stimulation by ALY in concert with this endogenous isoform. Consistent with this interpretation, the increase was not observed with a TCRα enhancer in which both AML-binding sites were mutated [Fig. 9D, TCRαmA]. Taken together, these data indicate that the stimulation of TCRα enhancer activity by ALY is dependent on the presence of any two of the three lymphoid-specific factors LEF-1, AML-1, and Ets-1, presumably in combination with a member of the ubiquitously expressed CREB/ATF family. However, this
stimulation appears to be independent of a simultaneous interaction of ALY with LEF-1 and AML-1.

**Discussion**

In this report, we present the identification and characterization of a ubiquitously expressed nuclear protein, termed ALY, that interacts specifically with two TCRα enhancer-binding factors, LEF-1 and AML-1. We find that overexpression of ALY in a reconstituted HeLa cell system stimulates the activity of the TCRα enhancer and that down-regulation of ALY using anti-sense oligonucleotides virtually abrogates TCRα enhancer activity in T cells. Thus, ALY seems to play a critical role in the function of the TCRα enhancer. Intriguingly, ALY interacts with domains of LEF-1 and AML-1 that mediate transcriptional activation within a specific context of other DNA-binding factors. However, ALY does not seem to contain an activation domain capable of stimulating transcription when tethered to multimerized DNA sites through fusion to a Gal4 or LexA DNA-binding domain. This finding distinguishes ALY from typical coactivators that contain transcriptional activation domains. For example, CBP is a coactivator that interacts with both various transcription factors and TFIIB, a component of the basal transcription machinery (for review, see Janknecht and Hunter 1996). Another characteristic of ALY is its ability to stimulate DNA binding of transcription factors including LEF-1 and AML. We propose that ALY is a novel type of transcriptional regulator that may function to coordinate the activity of CADs within multiprotein enhancer complexes or to regulate the DNA-binding properties of proteins within the TCRα enhancer complex.

**ALY acts in the context of multiple TCRα enhancer-binding proteins**

The finding that ALY associates with both LEF-1 and AML proteins may account for the marked dependence of the TCRα enhancer on ALY. Down-regulation of ALY in T cells by anti-sense oligonucleotides decreases the activity of the enhancer by at least 100-fold. In contrast, mutations of individual factor-binding sites in the enhancer reduces its activity 5- to 10-fold (Giese et al. 1995). Thus, ALY may be important for the function of both LEF-1 and AML-1, and possibly other factors that participate in regulating the activity of the multiprotein TCRα enhancer complex. The context dependence of ALY function is reminiscent of the requirement of LEF-1 and AML-1 to act in a specific context of proteins. Part of the context dependence of LEF-1 can be attributed to the binding of DNA by the HMG domain of LEF-1 (Giese et al. 1992, 1995), but the activation domain of LEF-1 tethered to a heterologous DNA-binding domain also shows a context dependence (Carlsson et al. 1993; Giese and Grosschedl 1993). Part of the context-dependent function of AML-1 protein may reflect its direct interaction with Ets-1, which binds an adjacent site in the TCRα and TCRβ enhancers (Wotton et al. 1994; Giese et al. 1995; Sun et al. 1995). The ternary AML:Ets-1:DNA complex
is, however, rather unstable and is, by itself, not competent to stimulate transcription from a minimal promoter (Giese et al. 1995). Therefore, additional interactions may be involved in mediating the context dependence of LEF-1 and AML proteins.

Although the interactions of ALY with both LEF-1 and AML-1 could, in principle, contribute to the context dependence by bridging these DNA-binding proteins, ALY also stimulates the activity of the TCRα enhancer in the absence of either LEF-1 or AML-1. Moreover, we failed to detect a further stabilization of the ALY:LEF-1:DNA ternary complex in vitro in the presence of AML or vice versa [data not shown]. Thus, ALY may mediate transcriptional stimulation by facilitating the interaction of the CADs of LEF-1 and AML-1 with other components of the TCRα enhancer complex. In addition to the association with LEF-1 and AML-1, ALY shows some ability to interact weakly with members of the ATF/CREB family in vitro, raising the possibility that ALY may function to couple LEF-1 or AML-1, or both, to factors bound at the ATF/CREB site. Interestingly, the formation of the ternary complex of ALY with LEF-1 or AML proteins and DNA is dependent on the 98-bp TCRα enhancer fragment and is not observed with a 50-bp fragment that lacks sequences upstream of the LEF-1 binding site. These observations raise the possibility that ALY may also interact weakly with upstream sequences, although we did not detect a clear footprint in DNase I protection assays. Thus, ALY may participate in the multiple protein–protein and protein–DNA interactions involved in the TCRα enhancer complex, which, together, account for the context dependence of individual protein components.

**LEF-1-interacting proteins contribute to distinct transcriptional regulatory functions**

Consistent with the context-dependent function of ALY, interaction of this protein with LEF-1 does not mediate activation of transcription from multimerized LEF-1 binding sites [S.-C. Hsu and R. Grosschedl, unpubl.]. However, another LEF-1-interacting protein, β-catenin, which has been shown recently to associate with the amino-terminal 76 residues of LEF-1 [Behrens et al. 1996; Huber et al. 1996], confers on LEF-1 and the related HMG-domain protein XTCF-3 the ability to stimulate transcription from multimerized sites, although β-catenin does not augment LEF-1 function at the TCRα enhancer [Behrens et al. 1996; Molenaar et al. 1996; S.-C. Hsu and R. Grosschedl, unpubl.]. Thus, interactions of LEF-1 with distinct proteins mediate either context-dependent or context-independent regulatory functions.

β-Catenin is involved in both cell adhesion and the wnt signal transduction pathway, suggesting that LEF-1 may function as a nuclear target of this developmentally important signaling pathway. Consistent with this putative function of LEF-1, targeted inactivation of the *Lef1* gene in the mouse resulted in severe developmental defects in the formation of tissues that express this gene during embryogenesis, including teeth, mammary glands, and hair follicles [van Genderen et al. 1994]. Although targeted mutation of the *Lef1* gene did not abrogate the expression of the TCRα gene [van Genderen et al. 1994], recent analysis of mice carrying a mutation in both the *Lef1* gene and the closely related *TCf1* gene revealed a complete block in T-cell differentiation and a lack of TCRα gene expression [R. Okamura, H. Clevers, and R. Grosschedl, unpubl.]. Thus, LEF-1 and TCF-1 may play redundant roles in regulating TCRα gene expression in vivo. TCF-1 is significantly more abundant than LEF-1 in thymocytes [Travis et al. 1991; van de Wetering et al. 1991]. However, TCF-1 is 10-fold less efficient than LEF-1 in stimulating TCRα enhancer activity in transfected tissue culture cells [van de Wetering et al. 1996; L. Bruhn, unpubl.]. Interestingly, TCF-1, which contains an HMG domain virtually identical to that of LEF-1, is divergent in the ALY-interacting region and ALY associates only weakly with TCF-1 [L. Bruhn, unpubl.]. Thus, the ability of DNA-bound LEF-1 to interact stably with ALY may, at least in part, explain the higher efficiency of LEF-1 in stimulating TCRα enhancer function relative to TCF-1.

**The ALY-interaction domain of AML-1 is translocated in many leukemias**

Studies of AML function have demonstrated that this protein plays an important role in transcriptional regulation of many targets in addition to TCRα. The block in fetal liver hematopoiesis displayed by mice carrying a targeted mutation in the *AML-1* gene demonstrates that AML-1 plays a role in regulating genes required for definitive hematopoiesis of all lineages [Okuda et al. 1996; Wang et al. 1996]. The abundant expression of ALY observed in the fetal liver [L. Bruhn, unpubl.] is consistent with a possible role for ALY in mediating AML function at these putative targets. At least 25% of all childhood leukemias involve translocations in the genes encoding AML-1 or its heterodimeric DNA-binding partner CBFβ [for review, see Ito 1996]. Interestingly, these translocations often replace the carboxy-terminal transcription activation domain of AML-1, which interacts with ALY, with segments of other proteins producing dominant-negative fusion proteins that can bind DNA strongly but do not activate transcription normally. To date, ALY is the only reported protein that interacts with the carboxyl terminus of AML-1, whereas all other known AML-interacting proteins, including CBFβ, Ets-1, and C/EBP, associate with the Runt domain. In addition to its role at the TCRα enhancer, ALY may facilitate the function of AML-1 in combination with other proteins such as Myb and C/EBP, which have been shown to stimulate transcription in a synergistic manner with AML proteins at the TCRβ enhancer and the macrophage colony-stimulating factor receptor promoter, respectively [Hernandez-Munain and Krangel 1995; Zhang et al. 1996].

**Potential functions of ALY in regulating the TCRα enhancer**

The ability of ALY to bind weakly to RNA raises the
possibility that, by analogy with HIV Tat protein (for review, see Jones and Peterlin 1994), ALY may stimulate gene expression by interacting with the nascent transcript. However, stimulation of the activity of the TCRα enhancer by ALY is observed with either the tk or fos promoters linked to the bacterial CAT reporter gene (Fig. 7A; data not shown). Thus, ALY-mediated stimulation of the TCRα enhancer does not appear to require specific promoter or coding sequences.

The association of ALY with the CADs of LEF-1 and AML-1 can also occur with DNA-bound proteins, although the protein–protein interactions appear to be weak because high concentrations of ALY are required. One notable feature of the interaction of ALY with LEF-1 and AML is the 5- to 20-fold increase in DNA binding by both proteins. This effect of ALY on DNA binding involves the amino- and/or carboxy-terminal regions of ALY because ALY-ANC does not increase DNA binding by LEF-1 or AML proteins. The stimulation of DNA binding by full-length ALY may be related to its potential to interact nonspecifically with nucleic acids. Consistent with this view, DNA binding by other proteins, such as c-Jun and glucocorticoid receptor that do not significantly interact with ALY, can also be augmented by addition of high concentrations of ALY in vitro (data not shown). However, the stimulation of transcription by ALY appears to require specific interaction domains because ALY does not stimulate the function of c-Jun or glucocorticoid receptor in transfection assays (Fig. 7B; data not shown).

Several proteins have been found to increase DNA binding by other factors. For example, HMGI(Y), a small basic DNA-binding protein, stably interacts with both ATF-2 and nuclear factor-κB (NF-κB) on the γ-interferon enhancer and increases DNA binding of these proteins and their assembly into a stereospecific multiprotein complex (Thanos and Maniatis 1992, 1995). The unrelated proteins HMG-1 and HMG-2 have also been shown to augment DNA binding of various proteins without participating in formation of a stable ternary complex (Onate et al. 1994; Zwillling et al. 1995). In addition, the protein hnRNP-K, which contains a KH motif RNA-binding domain and interacts with single-stranded nucleic acids, has been shown to interact with TATA-binding protein [TBP] in vitro and to function as a transcription factor in vivo (Michelotti et al. 1996). Finally, dimerization cofactor of hepatocyte nuclear factor 1 [DcoH], a transcriptional coactivator containing a potential RNP RNA-binding motif augments the dimerization and DNA binding by hepatocyte nuclear factor-1 (Mendel et al. 1991; Endrizzi et al. 1995). Taken together, these proteins appear to increase DNA binding by facilitating structural changes in DNA or proteins that accompany binding of individual proteins or the assembly of multiprotein complexes.

By analogy with these proteins, a simple model for ALY function at the TCRα enhancer could involve the increase of DNA binding by both LEF-1 and AML proteins. According to this view, the RNP motif of ALY, which mediates association with the CADs of LEF-1 and AML-1, would recruit ALY into the enhancer complex to allow the basic RGG motifs of ALY to facilitate the structural distortion of DNA that accompanies binding by both LEF-1 and AML-1 (Giese et al. 1992; Love et al. 1995; Golling et al. 1996). However, this model of ALY function cannot account for all observations. First, ALY stimulates the activity of the TCRα enhancer in vivo by fivefold in the presence of both LEF-1 and AML-1 and a similar increase is observed when only one of these factors is present (Fig. 9D). Second, compared to the ALY-mediated stimulation of DNA binding by either LEF-1 or AML proteins individually, ALY does not appear to further augment binding of both factors together in electrophoretic mobility-shift assays (data not shown). In addition, preliminary experiments suggest that addition of purified ALY protein to a mixture of LEF-1, AML-3, Ets-1, and ATF-2 or CREB is not sufficient to promote stable complex formation in DNase I footprint assays (A. Munnerlyn, unpubl.). Previously, we have shown that stable formation of the ternary AML:Ets-1:DNA complex is dependent on LEF-1 and Hela cell nuclear extract, which may have provided the correct ATF/CREB family member or other factors required for stable complex formation (Giese et al. 1995).

Another possible, not mutually exclusive, model is that the association of ALY with the CADs of LEF-1 and AML-1 facilitates the coordination of multiple activation domains in the multiprotein TCRα enhancer complex. This stimulation could be mediated by direct interactions of a composite activation surface with basal factors or by recruitment of as yet unidentified coactivators. Thus, ALY interaction with multiple TCRα enhancer-bound factors may be involved in the functional synergy observed between these factors. For both models ALY may be incorporated stably into the TCRα enhancer complex, possibly by operating as a molecular glue. Alternatively, ALY may act in a transient manner to facilitate interactions between multiple proteins or the DNA and may not be a structural part of the transcriptionally active multiprotein complex. Whatever molecular mechanism accounts for the function of ALY in regulating the TCRα enhancer, the association of ALY with the CADs of LEF-1 and AML-1, along with its observed functional properties, suggest that ALY may function to integrate and coordinate multiple protein–protein and/or protein–DNA interactions within specific higher order nucleoprotein complexes.

Materials and methods

Plasmids and oligonucleotides

Oligonucleotide-directed mutagenesis was used to replace the LEF-1 site in TCRα-fos–CAT (Travis et al. 1991) with a Gal4-binding site and to mutate the AML-binding sites (underlined) in TCRαmA-fos–CAT 5′-CTCAGaAGAActaACATCC; mutated residues are in lowercase letters. Plasmid CMV–AML-1 was generated by inserting a BsaBI–EcoRI fragment from pKS+PEBP2aB1 (Ba et al. 1994) into pEVRF0 (Matthias et al. 1989), pCMV–ALY and pCMV–T7–ALY were generated by inserting a Klenow-filled NcoI–XbaI fragment from pl38-20z containing ALY cDNA, into the Smal site of pEVRF0 and pEVRF0-
The latter vector contains coding sequences for the T7 epitope [MASMTGGQQMG]. pCMV-ALYANC contains an Eagl-BbsI fragment (codons 72-191) in pEVRF0. pCMV-ALY-Gal4 was constructed by inserting a fragment containing Gal4 codons 1-147 at a BglII site introduced at the stop codon of ALY by site-directed mutagenesis. Gal4[1-147] was also inserted into pEVRF0 to create pCMV-Gal4. 35S-Labeled ALY protein was expressed in vitro from p138-20z and full-length LEF-1 was expressed from pML-PEBP2~B1 (Bae et al. 1994). Amino- and carboxy-terminal deletion derivatives of LEF-1 and AML-1 were constructed using naturally occurring restriction sites and PCR-generated fragments. Mouse AML-3(94-513) (Fig. 3) was expressed from pET-PEBP2aAl(94-513) (K. Giese and R. Grosschedl, unpubl.). To allow 1-147 at a C-5 propyne pyrimidine-substituted phosphorothioate oligonucleotides used for the anti-sense experiments are as follows: The anti-sense ALY oligonucleotide [5'-GACATGTCCTTTTGTCCGC] is complementary to codons 2-8 of the ALY transcript. The shuffled control oligonucleotide [5'-TATCGCGGT-TGGCATACGTC] has the same base composition in a random order generated by the GCG sequence analysis program Shuffle.

**Yeast two-hybrid screen, cDNA isolation, and RNA analysis**

The LEF-1-LexA fusion construct [Giese and Grosschedl 1993] was subcloned into pBTM116 [Hollenberg et al. 1995] and the resulting plasmid was transformed into yeast strain L40, which contains integrated lexA-lacZ and lexA-HIS3 reporter genes [Hollenberg et al. 1995]. The mouse E10 embryonic day 9.5-10.5 cDNA-VP16 fusion library, which contains inserts averaging 300 bp in length, was screened using methods described in Hollenberg et al. [1995]. From 2.7 x 10^9 original transformants, 200 HIS'/LacZ' clones were isolated. Of these, 75 maintained expression of the reporter genes only in the presence of LEF-1-LexA. Library plasmids isolated from these clones were reintroduced into strain L40 containing expression plasmids for LexA, LEF-1-LexA, LexA-lamin [Hollenberg et al. 1995], LexA-STE3, or LexA-FUS1 [kind gifts of S. Givens, B. Ferguson, and G. Sprague, University of Oregon, Eugene], to isolate eight cDNAs that encode peptides that interact specifically with LEF-1. The 310-bp two-hybrid cDNA clone encoding ALYANC was radiolabeled by random priming and used to screen a murine thymocyte cDNA library, a kind gift of M. Davis [Stanford University, CA]. The sequence of a 1.1-kb cDNA clone 138-20z and the predicted amino acid sequence were compared with the GenBank DNA and protein databases using the blast algorithm [Altschul et al. 1990].

**Immunofluorescence and immunoblots**

COS7 cells were transfected transiently by electroporation with pCMV-T7-ALY and were allowed to settle onto poly-l-lysine-coated slides and incubated for 48 hr. Slides were processed as described in Travis et al. [1991]. For immunodetection, T7 monoclonal antibody [Novagen] was used at a 1:1000 dilution, followed by a fluorescein isothiocyanate (FITC)-conjugated donkey anti-mouse antibody. 4,6-Diamidino-2-phenylindole (DAPI, 0.1 μg/ml) was used to stain nuclei. Whole-cell extracts of Jurkat T cells transfected transiently with CMV-T7-ALY and antisense oligonucleotides were prepared by boiling and sonicating the cells in SDS-PAGE sample buffer. Extract from 2 x 10^6 cells was separated on 12.5% SDS polyacrylamide gels, transferred to nitrocellulose membrane, probed with T7 monoclonal antibody [Novagen] followed by peroxidase-conjugated goat anti-mouse secondary antibody [Cappell], and developed with Renaissance chemiluminescence reagent [DuPont NEN].

**Cell culture, transient transfections, and reporter gene assays**

COS7 and HeLa cells were cultured at 37°C in Dulbecco’s modified Eagle medium, supplemented with 10% fetal calf serum (FCS). Lymphoid cells were grown in RPMI medium, supplemented with 10% FCS and 50 μM 2-mercaptoethanol. Transient transfections for the experiments shown in Figures 7 and 9 were performed by the DEAE-dextran/chloroquine procedure with the amounts of plasmids indicated in the figure legends and 250 ng of RSV-β-galactosidase control plasmid [Grosschedl and Baltimore 1985]. Transfections of COS7 cells for the experiment of Figure 9C were performed by electroporation with 20 μg of CMV expression plasmids. Anti-sense oligonucleotides were transfected with Lipofectin according to manufacturer’s instructions [GIBCO-BRL]. CAT, luciferase, and β-galactosidase assays were performed as described by Starr et al. [1996]. After background subtraction, CAT or luciferase activity was normalized to the activity of the β-galactosidase internal transfection control.

**In vitro protein–protein association assays**

35S-labeled proteins were synthesized using a coupled transcription/translation (TnT) kit [Promega]. GST fusion proteins immobilized on glutathione–agarose beads were incubated with radiolabeled proteins in binding buffer [10 mM Tris-HCl (pH 8), 150 mM NaCl, 0.2% NP-40, 1 mM DTT, 0.2% BSA] for 2 hr at 4°C. Beads were collected by centrifugation and washed four times with 1 ml of binding buffer followed by one wash with 1 ml of binding buffer lacking BSA. Bound proteins were eluted from the beads by boiling in sample buffer [62.5 mM Tris-HCl (pH 6.8), 2% SDS, 10% glycerol, 5% β-mercaptoethanol], separated by SDS-PAGE, and visualized by autoradiography. Where indicated, the GST and 35S-labeled proteins were preincubated with 50 μg/ml pancreatic RNase [Boehringer Mannheim] for 15 min at 22°C before the association reactions. This amount of RNase was sufficient to eliminate the ribosomal RNA present in the in vitro translated 35S-labeled proteins as detected by ethidium bromide staining.

**Protein purification, nuclear extracts, and DNA-binding analysis**

GST fusion proteins and the LEF-1-HMG domain polypeptide were overexpressed in bacteria and purified using glutathione-conjugated agarose beads essentially as described in Giese et al. [1995]. The 6-His-tagged proteins LEF-1, AML-3 (94-411), and Runt were expressed in bacteria and purified using Ni-NTA agarose beads [Qiagen] according to manufacturer’s instructions. Nuclear extracts of COS7 cells were prepared according to the method of Schreiber et al. [1989]. Electrophoretic mobility-shift assays and DNase I footprinting were performed as described by Travis et al. [1991]. DNA-binding reactions contained 20 mM HEPES buffer (pH 7.9, 75
mm NaCl, 1 mm DTT, 2 mm MgCl₂, 10% glycerol, 0.1 mg/ml of BSA, 5 μg/ml of salmon sperm DNA, 2.5 μg/ml of poly[d(I-C)], ~10,000 cpm (5 fmole) labeled probe, and the amount of proteins or nuclear extract indicated in the figure legends. DNA-binding reactions with nuclear extract contained 50 μg/ml each of salmon sperm DNA and poly[d(I-C)]. For DNase I footprint analysis, the binding buffer also contained 0.5 mm spermidine.

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Note added in proof

The GenBank accession number for the sequence of ALY is U89876.

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L Bruhn, A Munnerlyn and R Grosschedl

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