Carboxy-terminal elements of c-Myb negatively regulate transcriptional activation in cis and in trans

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The c-Myb protein plays a key role in normal hematopoiesis, and truncation results in its activation to a transforming protein. Truncation of the c-Myb carboxyl terminus also greatly increases its transcriptional activating activity. The role of specific carboxy-terminal domains in negative regulation was investigated using Myb and Myb fusions with GAL4, LexA, or VP16. Negative regulatory activity of the carboxyl terminus in cis resides in at least two regions. A sequence in one of these regions can also inhibit transcriptional activation by Myb, Myb–VP16, or LexA–Myb proteins in trans. Regulation in trans, or suppression, is independent of c-Myb DNA binding and, therefore, likely involves protein–protein interaction. Suppression does not require the presence of a predicted heptad leucine repeat structure on either molecule. The target of suppression is a sequence that contains part of the minimal Myb transcriptional activation domain. This sequence can confer suppressibility on fusion proteins containing heterologous DNA-binding or transcriptional activation domains.

[Key Words: Negative regulation; transcription; suppression; c-Myb; oncogene]

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Additional work suggests that association of Myb with other proteins may be critical to the transformation process. Several mutations in the DNA-binding domain of Myb-related proteins have been described that can bind DNA in vitro but that are defective for transcriptional activation and/or transformation in vivo (Lane et al. 1990; Goff et al. 1991; Grasser et al. 1992). Furthermore, when two of the amino acid changes present in the v-Myb DNA-binding domain are individually back-mutated, the lineage of the transformed cells is altered (Introna et al. 1990). A heptad leucine repeat sequence located carboxy-terminal to the transcriptional activation domain in both v-Myb and c-Myb has recently been suggested to play a role in transactivation and transformation (Kanei-Ishii et al. 1992). These investigators hypothesize that binding of Myb to another protein through this putative leucine zipper is necessary for negative regulation of its activity.

Truncation of the carboxy-terminal region from c-Myb, in addition to activating its transforming potential, greatly increases the transcriptional activation activity of the protein (Sakura et al. 1989; Kalkbrenner et al. 1990). Although there seems to be some correlation between transactivating and transforming functions of v-Myb (Lane et al. 1990), not all transcriptionally active Myb proteins are competent for transformation. For example, a fusion of v-Myb to the transcriptional activation domain of the herpes simplex virus protein VP16 is a very strong activator (Ibanez and Lipsick 1990) but does not transform (U. Engelke and J. Lipsick, unpublished). Furthermore, internal deletion of a region in the carboxyl terminus of c-Myb that is highly conserved between vertebrates and Drosophila also fails to activate transformation (Grasser et al. 1991). We therefore performed a comprehensive deletion analysis to map the functional domains in the carboxyl terminus involved in transcriptional regulation by c-Myb and to elucidate their mechanism of action. The negative regulatory region of c-Myb is comprised of two domains, either one of which is sufficient in cis. One of these domains also inhibits transcriptional activation in trans [suppresses] through a specific Myb target sequence that can confer suppressibility on heterologous DNA-binding or activation domains. Our results show that the c-Myb carboxyl terminus contains functionally complex sequences capable of negatively regulating Myb transcriptional activation activity both in cis and in trans. We present possible models for these functions of the c-Myb carboxyl terminus.

Results
Identification of two negative regulatory regions in the carboxy terminus of chicken c-Myb

Transcriptional activation studies using murine and human c-Myb have shown that deletions in the carboxy-terminal region can greatly increase its transcriptional stimulation capacity (Sakura et al. 1989; Kalkbrenner et al. 1990). These studies placed the "negative regulatory domain" in two essentially nonoverlapping regions of the carboxy terminus corresponding to amino acids 416–499 and 497–641 of chicken c-Myb. To test for the presence of a negative regulatory domain in the chicken homolog, we first tested deletions of c-Myb corresponding approximately to the truncations present in the v-Myb protein, both individually and in combination. These deletions [d] remove either amino-terminal sequences [dCC], carboxy-terminal sequences [CCd], or both [dCd] from the full-length c-Myb protein, CCC (Fig. 1) (Grasser et al. 1991). Expression of the chloramphenicol acetyltransferase (CAT) gene driven by a tandem array of Myb-binding sites upstream of the E1b TATA sequence was assayed using transient transfections of viral vectors in QT6 quail fibroblasts (Fig. 1). Full-length c-Myb was essentially inactive. Amino-terminal truncation had little effect on transcriptional levels, whereas carboxy-terminal truncation consistently increased transcription. A control reporter plasmid lacking Myb-binding sites failed to give CAT activity above background. Essentially the same results were observed in HD-11 cells, a v-Myc-transformed macrophage cell line that does not express c-Myb (data not shown). Thus, the carboxyl terminus of chicken c-Myb contains sequences capable of negatively regulating its transcriptional activation capability.

To investigate whether negative regulation by the carboxyl terminus of chicken c-Myb required the presence of the highly conserved Myb DNA-binding domain, we constructed a fusion protein in which the amino-terminal DNA-binding region of chicken c-Myb was replaced...
by the unrelated DNA-binding domain of the yeast transcriptional activator GAL4 (Kakidani and Ptashne 1988). Amino acids 1-147 of GAL4, which also contain a nuclear localization signal, were fused to amino acids 233-641 of c-Myb at a site between the c-Myb DNA-binding and activation domains. This construct is similar to that used by Kalkbrenner et al. (1990), except that it contains additional GAL4 sequences necessary for efficient dimerization by GAL4, and will be referred to hereafter as GAL4-c-Myb (GAL4-CC). An additional construct was made that is truncated at c-Myb amino acid 428 near the position of v-Myb carboxy-terminal truncation [deletion (d) 428–641 in Fig. 2; also called GAL4-Cd]. Transfection of these plasmids, along with a reporter plasmid having five GAL4-binding sites upstream of the Elb TATA sequence and CAT gene, demonstrated that negative regulation by carboxy-terminal sequences of chicken c-Myb also functions in the context of a heterologous DNA-binding domain. Transcriptional activation by the full-length GAL4-c-Myb fusion (GAL4-CC) was negligible; however, the truncation of the Myb sequence in d428–641 resulted in greatly increased levels of activation (Fig. 2). Transcriptional activation from a control reporter plasmid lacking GAL4 sites was negligible (data not shown). The GAL4 DNA-binding domain alone, designated GAL4(DBD) in Figure 2, did not activate. Similar results were obtained with BK3A cells, an avian leukemia virus-transformed B-cell line that expresses high levels of endogenous c-Myb (Lüscher et al. 1990a; data not shown).

Progressive truncation from the carboxyl terminus (Table 1) identified a region between amino acids 428 and 462 [MscI–SacI] that is necessary for strong negative regulatory function. Transcriptional activation by the fusion protein truncated to BglII (amino acid 499) resulted in low levels of CAT activity (Fig. 2). Further truncation to the upstream SacI site (amino acid 463) resulted in only a slight increase in CAT activity. However, additional truncation to the MscI site (amino acid 428) significantly increased CAT expression [about sixfold]. This protein exhibited the greatest activity even though it was considerably less abundant than the others as shown by immunoblotting of extracts from the same cells (Fig. 2). Each of the other constructs produced similar steady-state levels of fusion protein of the expected molecular weight. Therefore, differences in transcriptional activation were not the result of different protein stabilities.

Six small deletions of between 35 and 61 amino acids

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**Figure 2.** Negative regulation of GAL4-c-Myb by carboxy-terminal sequences in cis. (A) The transcriptional activities of GAL4-c-Myb and its deletion mutants are shown relative to the best activator, d428–641 [GAL4-Cd]. The schematics showing GAL4 (cross-hatched) and c-Myb sequences, and promoter elements that direct CAT expression are as in Fig. 1, except that five GAL4-binding sites replace the nine sites specific for Myb. The asterisk (*) indicates the deletion that removes the potential Myb leucine zipper structure. The double deletion d425–464 + 499–558 was tested in a separate experiment along with d428–641 to which its activity was also normalized. Transfections of reporter (1 μg) and activator (5 μg) plasmids were as in the legend to Fig. 1. Activity of 100% represents 6,600 and 23,700 cpm, respectively. (B) Immunodetection of GAL4-c-Myb proteins in transfected cell extracts. Samples were analyzed by SDS-PAGE as described in Materials and methods and detected using a mixture of anti-Myb-2.2 and Myb-2.7 antibodies. Lines at left indicate molecular mass markers of 180, 116, 84, 58, 48, 36, and 26 kD, in descending order.
were constructed throughout the carboxy-terminal region (amino acids 367–641 of chicken c-Myb) to locate the negative regulatory region more precisely (Table 1). These deletions span the entire carboxy-terminal region deleted in the viral protein (amino acids 443–641 of chicken c-Myb). Transcriptional analysis by transient transfection of the appropriate activator and reporter plasmids into quail fibroblasts revealed that only one deletion [d499–558; BglII–NsiI] of the six tested resulted in any detectable activation of transcription by the GAL4–c-Myb fusion proteins (Fig. 2). None of the other five small deletions resulted in transcriptional activation, including the one that removes the predicted Myb leucine zipper [d367–427]. Internal deletion of an amino acid sequence (amino acids 425–464, PflMI–SacI), almost identical to the region necessary for strong negative regulation (amino acids 428–462) that was identified by progressive truncation, had no effect on transcriptional activation. These experiments therefore suggested that the carboxy terminus of c-Myb contains a distal regulatory sequence carboxy-terminal to amino acid 499 and a proximal regulatory sequence between amino acids 428 and 462 [MscI–SacI], either of which is sufficient for strong negative regulation in cis. Simultaneous deletion of both regions resulted in ~60% of the activity of the most active carboxy-terminal truncation [d428–641], demonstrating that each of the regions can function independently as a negative regulatory element (Fig. 2).

The c-Myb carboxyl terminus inhibits trans-activation in vivo, not DNA binding

To assess the effect of the carboxy terminus on nuclear transport and DNA binding in vivo, we also made a fusion of the carboxy-terminal portion of c-Myb with the bacterial repressor LexA. Fusions of this type have been used to demonstrate in vivo DNA binding of chimeric proteins to repressible promoters in animal cells (Godowski et al. 1988; Martin et al. 1990). This LexA–c-Myb fusion protein is analogous to the GAL4–c-Myb fusion except that the 147-amino-acid GAL4 DNA-binding domain was replaced by the 202 amino acids of LexA. A second construct was made by deleting the carboxy terminus of c-Myb from the LexA fusion protein near the position of viral truncation. The in vivo DNA-binding [repression] assay was performed with reporter plasmids containing the SV40 promoter/enhancer driving CAT gene expression. The test plasmid contained one Lex operator positioned near the site of transcription initiation, whereas the control plasmid contained no Lex operator sites (Fig. 3). Thus, LexA proteins should bind to the Lex operator of the test plasmid and repress transcription from the SV40–Lex promoter, but not from the SV40 promoter of the control plasmid.

Because the LexA fusion proteins were used as a model to assess the effect of carboxy-terminal truncation on in vivo DNA binding, it was important to confirm that they demonstrated the same relative transcriptional activation behavior as the GAL4–c-Myb fusions. The same LexA–c-Myb fusions were therefore tested, along with the GAL4 and LexA DNA-binding domains alone, for transcriptional activation activity using the E1b TATA-CAT reporter plasmid with or without six upstream Lex sites (Fig. 3). Lex site-dependent transcriptional activation was not observed with the GAL4 DNA-binding domain [GAL4(DBD)], LexA, or the full-length LexA fusion [LexA–CC], but strong activation was seen with the truncated fusion [LexA–Cd]. These results were similar to those obtained with the corresponding GAL4 fusion proteins [GAL4–CC and GAL4–Cd [d428–641]; see Fig. 2] and the GAL4 reporter plasmid. Immunoblotting of transfected cell extracts [Fig. 3] showed the presence of similar levels of LexA–c-Myb fusion proteins of the expected molecular weights.

Plasmids encoding the LexA DNA-binding domain [LexA], the full-length fusion protein [LexA–CC], the truncated fusion protein [LexA–Cd], and GAL4(DBD) were each cotransfected with either the test plasmid [SV40–Lex–CAT] or the control plasmid (SV40–CAT). The GAL4(DBD) protein should not affect transcription from either reporter and would therefore result in constitutive CAT expression levels against which to compare the LexA and LexA–c-Myb fusion proteins. Transcription from the control plasmid, SV40–CAT, was reduced somewhat in the presence of the LexA proteins relative to the GAL4(DBD) (Fig. 3). Perhaps there is a cryptic Lex operator in the SV40 promoter/enhancer region. However, all three LexA-containing proteins greatly reduced the level of transcription from the SV40–Lex–CAT plasmid, relative to GAL4(DBD). These results showed that the repression was dependent on the presence of both the LexA DNA-binding domain and Lex operator sequences. Expression of the CAT gene from the test plasmid containing the Lex operator relative to

### Table 1. Deletions within the carboxyl terminus of c-Myb

| Deletion | DNA sequence | Amino acid sequence | Region deleted |
|----------|--------------|---------------------|----------------|
| d428–641 | TGT GGC TAG  | Cys-Gly-Stp          | Msc-end        |
| d463–641 | GAG,CTA GAT  | Glu-Leu-Asp-Lys     | Sac-end        |
| d499–641 | GAT CCT AGA  | Asp-Pro-Arg         | Bgl-end        |
| d367–427 | CCT GAC CAC  | Pro-Asp-His         | Bsu–Msc        |
| d425–464 | CCA GCT CCA  | Pro-Ala-Pro         | PflMI–Sac      |
| d463–497 | GAG,GAT      | Glu-Asp             | Sac–Bgl        |
| d499–558 | GAT CAT GCA  | Asp-His-Ala         | Bgl–Nsi        |
| d561–598 | GCA GCT AGC  | Ala-Ala-Ser         | Nsi–Nhe        |
| d601–641 | GCT AGC TAG  | Ala-Ser-Stp         | Nhe-end        |

*a The chicken c-Myb amino acids deleted [d] in each mutant are given.

*b The nucleotide sequence at the deletion junction is shown. Carets indicate the point of DNA fusion, except where an adaptor was used. The underlined nucleotide in the adaptor was added to bring the coding sequence in-frame. Codons through the deletion junction are separated by spaces.

*c Amino acid sequence of the junction. Underlined amino acids are those not found in c-Myb. [Stp] Stop codon.
The carboxyl terminus of c-Myb inhibits transcriptional activation by v-Myb or a v-Myb–VP16 fusion in trans

We also tested the ability of c-Myb carboxy-terminal sequences in trans to affect transcription levels stimulated by v-Myb. A proviral vector that expresses v-Myb was cotransfected with the Myb-responsive E1b TATA/CAT gene reporter and plasmids that express different parts of the c-Myb protein (Fig. 4). Cotransfection with DNA encoding c-Myb proteins that contained the carboxyl terminal significantly inhibited v-Myb transcriptional activation relative to vector DNA lacking any c-Myb-coding sequence. The level of inhibition by the carboxyl terminus was similar whether (CCC) or not (dCC) the amino terminus was present. The doubly truncated c-Myb protein (dCd) resulted in no inhibition, whereas the singly truncated proteins dCC and CCd resulted in strong and moderate inhibition of v-Myb activation, re-
spectively. The results of this experiment were complicated by the fact that the c-Myb proteins used as inhibitors can also bind to Myb-binding sites. This binding might result in a complex composite reaction by the c-Myb protein being tested, reflecting both its own transcriptional activating activity and its inhibition of v-Myb transcriptional activation. Therefore a c-Myb protein from which the DNA-binding domain had been deleted was also tested. Transcriptional activation was inhibited as strongly by this protein, CCC(dDBD), as by the other c-Myb proteins containing the carboxyl terminus, providing strong evidence that c-Myb was inhibiting v-Myb activation in trans. As seen with the simple activation experiments using different c-Myb proteins to examine negative regulation in cis (Fig. 1), the ability to inhibit v-Myb transcriptional activation activity in trans localized to the carboxyl terminus of c-Myb.

We tested a v-Myb–VP16 fusion to determine whether trans-inhibition would operate on a strong heterologous transcriptional activation domain [Triezenberg et al. 1988]. The fusion was made at a site in v-Myb within its transcriptional activation domain and is referred to as the v-Myb–VP16 fusion protein [VVP] [Ibanez and Lipsick 1990]. Truncation of native v-Myb at this site abolishes transformation and transcriptional activation [LX2 [Ibanez and Lipsick 1988, 1990]]. When expressed from a proviral vector, this VVP fusion protein stimulated high levels of transcription from a reporter plasmid containing Myb-binding sites in front of the Elb TATA/CAT gene combination but not from a reporter without Myb-binding sites [Grasser et al. 1992]. The high activity of the VVP protein required less DNA for each transfection and resulted in a greater range of activation over which to examine inhibition. c-Myb inhibited VVP activity by at least 80% at plasmid ratios between 0.3 : 1 and 30 : 1 (data not shown). All of those c-Myb proteins [CCC, dCC, GAL4-CC] that contain the carboxyl terminus strongly inhibited activation by VVP, whereas both c-Myb proteins that lack carboxy-terminal sequences [CCd and dCd] failed to inhibit VVP activity significantly (data not shown). Although it is unable to bind Myb-binding sites, the GAL4–c-Myb fusion protein was able to strongly inhibit VVP transcriptional activation [Fig. 5]. Furthermore, the DNA-binding domain deletion of c-Myb [CCC(dDBD)] also inhibited VVP (data not shown). These results demonstrate that inhibition of VVP activity by the carboxy-terminal sequence of c-Myb was not due simply to competition between the c- and v-Myb molecules for Myb-binding sites on the reporter DNA. We refer to trans-inhibition as “suppression” to distinguish it from negative regulation in cis. The region of c-Myb that performs this suppression is defined as having suppressor activity, and the region of v-Myb necessary for inhibition is called the target of suppression.

Suppression by c-Myb maps to one of the carboxy-terminal negative regulatory elements

The six carboxy-terminal deletions of the GAL4–c-Myb fusion protein previously tested for transcriptional activation were used to identify the region in the c-Myb carboxyl terminus required to suppress VVP activity in trans. Each of these deletion constructs was cotransfected along with VVP and the reporter plasmid containing upstream Myb-binding sites. Figure 5 shows that all of the deletions except d499–558 [BglII–NsiI] were capable of suppressing VVP activity. Specifically, deletion of the heptad leucine repeat sequence located between amino acids 376 and 404 of the chicken c-Myb sequence had no effect on suppressor activity (see d367–427 in Fig. 5). Immunoblot analysis of the same cells assayed for CAT activity demonstrated the presence of similar levels of the expected GAL4–c-Myb fusion proteins [Fig. 5]. Deletion of amino acids 499–558 from GAL4–CC increased transcriptional activation by VVP to between 55 and 70% of uninhibited levels in different experiments. It is interesting that the amino acid 499–558 region might function both in the trans-inhibition (suppression) of v-Myb–VP16 transcriptional activation (Fig. 5) and in

Figure 5. Mapping of c-Myb carboxy-terminal elements responsible for trans-inhibition. (A) Activities are shown relative to VVP in the presence of the GAL4(DBD). Schematics show protein sequences as in Fig. 2, except experiments included VVP and reporter that contained Myb-binding sites (top). The asterisk [*] indicates the deletion that removes the potential Myb leucine zipper structure. Transfections included reporter [1 μg], and plasmids expressing VVP [0.1 μg] and different GAL4-CC deletion mutants [3 μg] as described. Activity of 100% equals 70,500 cpm. (B) Immunodetection of GAL4–c-Myb proteins in transfected-cell extracts. Immunoblotting was performed, and markers are the same as in the legend to Fig. 2.
the negative regulation of its own transcriptional activity in cis (Fig. 2).

*Specific sequence in the Myb activation domain functions as a target of suppression*

To determine the region in v-Myb that was the target of suppression, we constructed deletions of v-Myb and VVP and tested them for their ability to be inhibited in trans by GAL4-c-Myb (Fig. 6). Most deletions of native v-Myb abolish transcriptional activation activity (Ibanez and Lipsick 1990; R.-H. Chen and J. Lipsick, unpubl.), even if they retain the minimal transcriptional activation domain defined by previous GAL4-v-Myb fusion studies (Weston and Bishop 1989). Because these v-Myb deletion proteins have no intrinsic activity, it is not possible to test them as targets for suppression. However, we constructed a deletion of a less well-conserved region, whose amino acid sequence suggests random coil character, located between the DNA-binding and transcriptional activation domains. A v-Myb mutant that was deleted of this possible "hinge" region (v-Myb-dH) retained ~50% of wild-type transcriptional activation activity (Fig. 6). Stimulation of transcription from the Myb-binding site reporter plasmid by this v-Myb-dH mutant was strongly suppressed by the full length GAL4-c-Myb fusion protein, as was wild-type v-Myb (Fig. 6). Both the trans-activating protein [v-Myb or v-Myb-dH] and the suppressing protein [GAL4-CC] were detected by immunoblotting of the same cells tested for CAT activity (Fig. 6).

These results indicated that the target of suppression must be in another part of the v-Myb protein, in the DNA-binding region, transcriptional activation region, or both. Because other v-Myb deletions fail to trans-activate, the v-Myb–VP16 fusion protein was used to further map the target sequence. The VP16 protein was suppressed by c-Myb even though it is missing the putative heptad leucine repeat. Therefore, the leucine repeat on the target molecule is not involved in suppression. A deletion of the region corresponding to amino acids 169–239 of v-Myb between Smal [Xmal] and HindII [SalI] sites in VVP was constructed [VVP–dTA] that removes the remainder of a negatively charged region implicated in transcriptional activation by v-Myb. This yields a protein consisting essentially of the v-Myb DNA-binding domain fused to the VP16 trans-activating sequence (Fig. 6). When tested for suppression by the c-Myb carboxy-terminal sequence, the VVP–dTA protein exhibited the same high activation levels in the presence of the GAL4–CC fusion protein or GAL4[DBD] alone (Fig. 6). Western blots using extracts from the same cells as those assayed for CAT activity showed that lack of suppression was not the result of the absence of the GAL4–CC protein (Fig. 6). The v-Myb DNA-binding domain alone is therefore not sufficient to function as a target of suppression, nor is the VP16 transcriptional activation domain alone a sufficient target. Rather, amino acids in the transcriptional activation domain of the v-Myb sequence between amino acids 169 and 239 [SmaI–HindII] are necessary for suppression by c-Myb.

**Figure 6.** Mapping of Myb target sequences that mediate trans-inhibition. [A] Activities are shown relative to each wild-type protein in the presence of GAL4[DBD] (left column). Schematics show protein sequences and reporter plasmid as in previous figures. Transfections included reporter [1 µg], and plasmids expressing v-Myb [5 µg], v-Myb-dH [5 µg], VVP [10.2 µg], VVP–dTA [0.5 µg], or LexA–Cd [3 µg], and GAL4[DBD] or GAL4–c-Myb [3 µg] as described. Results from the transfection of 0.25 µg of VVP–dTA (data not shown) were similar to those using 0.5 µg. Activity of 100% represents v-Myb, 10,000; VVP, 26,500; and LexA–Cd, 39,400 cpm. (B) Immunodetection of Myb and Myb fusions in transfected cell extracts. Immunoblotting was performed as in Fig. 2. VP16 fusion proteins are not detected owing to the low amount of plasmid transfected. The positions of v-Myb (double arrowhead), v-Myb-dH (single arrowhead), and GAL4–c-Myb (arrow) are indicated. Fig. 3B shows the stable production of immunoreactive LexA–Cd protein of the expected molecular mass.
To determine whether the Myb DNA-binding domain itself was necessary for suppression, we tested the highly active LexA–Cd protein for suppression by GAL4–CC. Neither of these proteins contains a Myb DNA-binding domain, furthermore, the use of LexA–Cd allowed us to assess the suppressibility of a c-Myb transcriptional activation domain. The activation domain of c-Myb differs from that of v-Myb by three amino acid substitutions. Figure 6 shows that transcriptional activation by LexA–Cd from the Lex site reporter was inhibited by GAL4–CC relative to GAL4[DBD], the GAL4 DNA-binding domain alone. This result demonstrated that the process of suppression does not require a Myb DNA-binding domain on either protein. In addition, the c-Myb activation domain of LexA–Cd is also capable of being suppressed, despite the three different amino acids in the target sequence.

Discussion

The differences between the transforming oncprotein v-Myb and its normal cellular counterpart, c-Myb, are responsible for the oncogenicity of the virus AMV. Deletion of either one or both ends of the c-Myb protein, which are absent in v-Myb, has been shown to be sufficient for activation of its transforming potential [Grasser et al. 1991]. Deletion of the carboxyl terminus of c-Myb also markedly increases its transcriptional activation capacity [Sakura et al. 1989; Kalkbrenner et al. 1990] [Fig. 1]. Previous work from a number of laboratories has attempted to define the elements necessary for negative regulation of c-Myb transcriptional activity (Sakura et al. 1989; Kalkbrenner et al. 1990; Hu et al. 1991; Kanie-Ishii et al. 1992). We undertook the present study to investigate the role of specific carboxyl-terminal domains in this regulation.

Our results show that no single small deletion within the carboxyl terminus results in the large increase in transcriptional activation activity observed with carboxyl-terminal truncations extending upstream of amino acid 463 [Fig. 2]. In contrast to the results of Kanie-Ishii et al. (1992), deletion of the region containing the heptad leucine repeat structure did not increase transcriptional activation above background, suggesting that it is not involved in the negative regulation examined here. Internal deletion of amino acids 499–558 was the only one that increased transcription by itself, albeit modestly. These results suggest that cis negative regulatory function exists in at least two regions of the c-Myb carboxyl terminus, either of which is sufficient. Although internal deletion of the more carboxy-terminal element (amino acids 499–558) increased transcription only slightly, removal of this region by truncation from the carboxyl terminus does activate a low level of transformation [Grasser et al. 1991; Hu et al. 1991], providing additional evidence that amino acids 499–558 may indeed be involved in negative regulation. The large increase in transcriptional activation observed upon simultaneous deletion of both regions implicated in negative regulation (d425–464 + 499–558) confirmed that either region can function as a strong negative regulatory element in the absence of the other. These experiments also resolve the discrepancy between previous reports that had assigned the negative regulatory domain to different nonoverlapping regions [Sakura et al. 1989; Kalkbrenner et al. 1990; Hu et al. 1991]. In addition, a sequence in the carboxyl terminus of c-Myb can negatively regulate transcriptional activation by v-Myb, or by a v-Myb–VP16 or LexA–c-Myb fusion protein in trans, a process we refer to as suppression. Suppressor activity correlates with one of the cis-acting negative regulatory regions and is independent of DNA binding by c-Myb. The target of suppression is a Myb sequence containing part of the transcriptional activation domain. Suppression does not require the presence of the Myb DNA-binding domain on either molecule.

These results provide evidence of a possibly complex domain organization of the carboxy-terminal region of c-Myb. Interestingly, an amino acid sequence within each negative regulatory region appears to be conserved in a number of Myb-related proteins, including the yeast BAS1 protein, which was previously believed to have homology only to the Myb DNA-binding domain [Tice-Baldwin et al. 1989] [Table 2]. In BAS1 the two regions of homology occur in the inverse order relative to c-Myb, suggesting that they may be modular in nature. Because the amino acid 499–558 deletion was one of the largest used in this study [60 amino acids] and deletions may result in gross overall structural changes, smaller deletions through this domain might provide more detailed information. Additionally, less disruptive linker insertion mutations throughout the entire carboxyl terminus may reveal even more complexity in the domain structure of the carboxy-terminal region.

The presence of the carboxyl terminus in cis seems to have little, if any, effect on in vivo DNA binding by a LexA–c-Myb fusion protein, although it strongly inhibits transcriptional activation. Ramsay et al. (1991, 1992) suggested that carboxy-terminal truncation may activate c-Myb in part by increasing its affinity for Myb-binding sites. We have also reported a progressive increase in v-Myb DNA binding that parallels progressive truncation from the carboxyl terminus [Garcia et al. 1991]. Perhaps these distal amino acid sequences sterically hinder or perturb the structure of the Myb DNA-binding domain. However, although it remains possible that changes in DNA-binding affinity play a part in the activation of c-Myb by carboxy-terminal truncation, the action of the cis negative regulatory domains described in this paper does not involve inhibition of DNA binding.

Negative regulation by the c-Myb carboxyl terminus also functions in trans to suppress transcriptional activation by v-Myb, v-Myb–VP16, or a truncated LexA–c-Myb protein, LexA–Cd. Binding to DNA by the protein bearing the c-Myb carboxy-terminal suppressor sequence is not required for this suppression. DNA-binding independent suppression by the carboxyl terminus eliminates competition for Myb-binding sites on the reporter plasmid as a possible mechanism. Therefore, protein–protein interactions are most likely involved. There are
Table 2. Alignment of carboxy-terminal domains of vertebrate Myb-related proteins and yeast BAS1

|          | Alignment 1 | Alignment 2 |
|----------|-------------|-------------|
| HumC 451–480 | R T P A I K R S I L E S S P R T P T P F K H A L A A Q E I K | Q D V I K Q E S D E S G F V A E F Q E N G P L L K K I K Q E V E S P |
| ChiC 452–481 | R T P A I K R S I L E S S P R T P T P F K H A L A A Q E I K | Q D V I K Q E S D E S G F V A E F Q E N G P L L K K I K Q E V E S P |
| HumA 540–569 | R T P R I R R S I L G T T P R T P T P R K N A L A A Q E R K | E D I R E V L K E E T G T D L F L K E K E D E P A Y K S C K Q E N T A S |
| BAS1 524–553 | R T P N Y N A F S L E A T S H N P A D N A N E L G S Q S N R | Q D I L N V K E N E S S K L P R L K D N D G P I L N D S K P Q A L P P |

Amino acid sequences of two regions in several Myb-related proteins are shown using the standard single-letter code. The residue numbers for each protein segment compared in the two alignments are given. Alignments were performed with the MACAW computer program [Schuler et al. 1991], kindly provided to us by Eric Westin.

Boldface type indicates identity of a residue in all four sequences.

(*) Identity of a BAS1 residue with the same position in at least two other sequences.

(○) Similarity of a BAS1 residue with the same position in at least two other sequences.

many possible models to explain how such interactions might operate [Fig. 7]. First, the interaction between the c-Myb suppressor and Myb acidic target domains may be direct. It could be relevant that several potentially basic amino acid segments have been found in appropriate regions of the c-Myb carboxyl terminus (Gonda et al. 1985). Second, the regulatory state could be modulated by the phosphorylation of either or both domains. There are several known or postulated phosphorylation sites in appropriate regions of c-Myb. Third, the negative regulatory region of c-Myb might contain a binding site for a small effector molecule, such as a hormone or vitamin. The activation of c-Myb by carboxy-terminal truncation does resemble that of the steroid hormone receptor, in that deletion of the hormone-binding domain also creates a constitutively active receptor [Evans 1988]. Finally, regulation could be facilitated by ancillary proteins that specifically interact with the Myb functional domains. The action of such accessory proteins would be similar to the adaptors proposed to mediate interaction between some DNA-binding transcription factors and the general transcription machinery [Pugh and Tjian 1990]. It is also possible that two or more of these mechanisms function in concert.

Mapping of the domains required for suppression indicates that a sequence in the downstream carboxy-terminal negative regulatory element [amino acids 499–558] contains most of the suppressor activity [Fig. 5]. Small amounts of additional suppressor activity residing in other regions may have escaped detection. A predicted Myb leucine zipper structure (Biedenkapp et al. 1988) present in the GAL4–c-Myb fusion protein appears to be unnecessary for suppressor function. Because the suppressible target v-Myb–VP16 lacks the potential Myb leucine zipper, this sequence is not involved in suppression either on the target molecule or on the carboxy-terminal suppressor sequence. The Myb sequence that does function as the target for suppression [amino acids 169–239; Fig. 6] includes the upstream two-thirds of the region defined as the minimal transcriptional activation domain [Weston and Bishop 1989]. Although this sequence contains most of the acidic character of the domain and is necessary as a target for suppression, it is not sufficient to support transcriptional activation either as a GAL4 fusion [Weston and Bishop 1989] or as the carboxy-terminal part of a nontransforming, truncated native v-Myb molecule [LX2 (Ibanez and Lipsick 1988, 1990)].

A model consistent with our results would include a c-Myb carboxy-terminal interaction mediated by one part of the Myb transcriptional activation sequence and
the “poisoning” of another part responsible for activation activity. In the case of v-Myb–VP16, the adjacent VP16 activation domain would be suppressed in an analogous way. Apparently, this process can tolerate the three amino acid changes present in the c-Myb activation domain of LexA–Cd. The carboxy-terminal sequences that perform these binding and poisoning functions would be “pseudosubstrates,” resembling the sequences on the protein molecules, such as adaptors or transcription machinery, that normally mediate transcriptional activation by Myb in the absence of negative regulation. Alternatively, the carboxy terminus of c-Myb may suppress multiple classes of transcriptional activation domain by a general mechanism such as steric hinderance. In any case, the results presented here and previously (Ibanez and Lipsick 1988, 1990) imply that the structure–function relationships of the Myb transcriptional activation domain may also be complex.

Interestingly, Grasser et al. (1991) observed suppression of v-Myb transformation by the c-Myb carboxy terminus in cis. The data reported in the present paper suggest that suppression in trans may have the same effect, a possibility supported by the fact that cell lines transformed by v-Myb or by truncated forms of c-Myb express little or no detectable full-length c-Myb (Lipsick 1987; Ramsay et al. 1989; Grasser et al. 1991). Thus, suppression of v-Myb activity by the c-Myb protein might constrain transformation to those precursor cells whose differentiation has progressed to the point that c-Myb expression has been shut off. The presence of an additional carboxy-terminal exon in some forms of murine and human c-Myb (for review, see Shen-Ong 1990) might alter their suppression activity or specificity. Human A- and B-Myb proteins that contain sequences homologous to this alternative exon have been found coexpressed with the normal c-Myb protein (Nomura et al. 1988). These classes of cellular Myb could interact with each other to suppress activity under different conditions of growth or differentiation.

In this paper we show that the c-Myb carboxy terminus contains functionally complex sequences capable of negatively regulating Myb transcriptional activity both in cis and in trans. Regulation in cis could actually be an intermolecular reaction in which two Myb molecules bind together and mutually inhibit transcriptional activation. Regulation in trans, or suppression, does not require DNA binding by the carboxy-terminal sequence and probably involves specific protein–protein interactions. The Myb DNA-binding domain is not required on either the suppressor or target protein. Our results do not distinguish between alternative direct or indirect interaction (Fig. 7). The heptad leucine repeat structure is apparently not involved in suppression, but it is possible that the use of a GAL4–c-Myb fusion precludes detection of the cis participation of this motif. The fact that the v-Myb target sequence is necessary for suppression of a heterologous activation domain suggests that transcriptional activation and suppression are specified by somewhat different amino acid sequences. Suppression represents a novel form of transcriptional regulation in trans and may also operate within other classes of transcriptional activator.

Materials and methods

Plasmid constructions

DNA restriction and modifying enzymes were purchased from New England Biolabs [Beverly, MA]. Oligonucleotide adaptors and primers were made on an Applied Biosystems [Foster City, CA] model 380B synthesizer. Recombinant DNA manipulations were carried out using standard techniques (Maniatis et al. 1982). Retroviral constructions that express v-Myb [dGE], c-Myb, and d-Myb truncations have been described previously (Ibanez and Lipsick 1988; Grasser et al. 1991). Single and double truncations of c-Myb are named according to the sequences present [C] or absent [d] relative to the v-Myb-coding sequence. Thus, the c-Myb proteins deleted of the amino-terminal, carboxy-terminal, or both sequences missing in v-Myb are referred to as dCC, C, or dCd, respectively. Expression of the v-Myb–VP16 fusion protein [VPV] used in this study has been described previously (Ibanez and Lipsick 1990).

Deletions within the carboxy terminus of c-Myb were constructed by blunt-end ligation of the appropriately restricted and Klenow-treated DNA where such treatment would give in-frame deletions [Table 1]. For the two deletions where this was not possible (BglII–NsiI, NsiI–NheI), advantage was taken of the fact that both of these constructions used restriction enzymes resulting in one 5′-sticky and one 3′-sticky end. A single-stranded DNA adaptor was therefore synthesized for the BglII–NsiI (5′-dCATCAGTGA) or NsiI–NheI (5′-dCTAGCTGCA) deletion that was complementary to both overhangs, and included an additional nucleotide necessary to give the correct reading frame and regenerate one of the original restriction sites [Table 1]. The appropriate 5′-phosphorylated adaptor was added to the purified large BglII–NsiI or NsiI–NheI resistant fragment, ligated using T4 DNA ligase, and transformed into Escherichia coli strain DH5αF′. Resulting transformants were screened with diagnostic restriction endonucleases, and the sequences of positive clones were confirmed by the dideoxy chain-termination method [Chen and Seeburg 1985]. The double deletion d425–464 was made by ligating a BstXI–BglII fragment that contained the d425–464 deletion to an NsiI–BstXI fragment in the presence of the BglII–NsiI adaptor. The resulting double deletion within the carboxy terminus was transferred into the GAL4–CC expression plasmid on an XmaI–BamHI fragment, and protein of the expected molecular size was detected upon transfection and immunoblotting as described. Deletions are named according to the chicken c-Myb amino acids removed in each case [Table 1].

Plasmids encoding the GAL4(DBD) and LexA were kind gifts from Stan Fields and Katherine Martin, respectively. Expression of these proteins and their fusions was controlled by the SV40 promoter and polyA/spllice site in the plasmid pECE [Ellis et al. 1986]. Fusions of chicken c-Myb with either GAL4 or LexA were made by ligating the blunt Smal site at c-Myb amino acid 233 to the upstream Smal in pSG424 [Sadowski and Ptashne 1989] or the filled EcoRI of pBXL1 [Martin et al. 1990]. The GAL4–CC fusion protein includes c-Myb amino acids 233–641, whereas LexA–CC encodes c-Myb amino acids 234–641. Carboxy-terminally truncated versions of both fusions (GAL4–Cd and LexA–Cd) were created by cleavage at the MscI site in c-Myb [amino acid 427] and an XbaI site in the distal untranslated region, filling in with Klenow, and religating [Table 1]. Reporter plasmids bearing nine Myb-, five GAL4-, or six LexA-binding sites were derived from E1b–CAT, which contains a
Deletions v-Myb-dH and VVP-dTA were made by ligating filled BstElI--XmaI and Smal--HincII ends to give in-frame deletions of v-Myb amino acids 134--167 and 169--239, respectively. These mutant proteins migrated with the expected mobility on SDS-PAGE relative to their respective parent proteins. Numbering of v-Myb residues begins with the initiating methionine encoded by the fused retroviral gag sequence at number one (Lane et al. 1990).

Cells and media
Quail QT6 fibroblasts were grown in Dulbecco's modified essential medium supplemented with 5% fetal calf serum, 4.5 grams/liter of glucose, nonessential amino acids, L-glutamine, sodium pyruvate, 100 µg/ml of streptomycin, and 100 U/ml of penicillin in a humidified 10% CO2/90% air incubator at 37°C. Chicken HD-11 macrophage and BK3A lymphocyte cells were grown in the same media with the addition of 5% heat-inactivated [1 hr at 56°C] chicken serum.

DNA transfections, immunoblotting, and assays for CAT activity
Transient transfections into QT6 quail fibroblasts and HD-11 macrophages were performed using a modification of the calcium phosphate procedure of Chen and Okayama [1987] as described previously [Ibanez and Lipsick 1990]. Introduction of DNAs into BK3A B-cells was done using the DEAE-dextran method [Choi and Engel 1988]. Transfections included 3--5 µg of activator plasmids that express proteins without the VP16 activation domain [v-Myb, CCC, dCd, GAL4--CC, etc.], from 0.1 to 0.5 µg of the activators encoding VP16 fusion proteins (VVP, VVP--dTA), and 3--6 µg of inhibitor plasmid when present. Activator plasmids were cotransfected with CAT reporter DNA [1 µg], tRNA [3.8 µg], and 0.5 µg of a plasmid expressing β-galactosidase (β-gal) from the cytomegalovirus (CMV) promoter as an internal control for transfection efficiency. All experiments were performed at least twice, and representative results are shown. Specific details are given in the legend of each figure.

Half of the cells from each transfection plate were solubilized by boiling for 4 min in SDS-PAGE loading dyes. Normalized volumes of each were then subjected to SDS-PAGE through 15% acrylamide gels. Normalized volumes of each were then subjected to SDS-PAGE through 15% acrylamide gels. The phase extraction method was employed to determine the CAT activity in normalized volumes of extract as described previously [Seed and Sheen 1988]. Appropriate dilutions were assayed in those cases where the level of CAT activity indicated that substrate was limiting. A background value, generally 100--300 cpm, was obtained by extracting CAT assay mix to which no cell extract had been added. Activities shown in the figures have been corrected for background.

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