Identification of factors that interact with the E1A-inducible adenovirus E3 promoter

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We have investigated the E1A-inducible E3 promoter of adenovirus type 5 with respect to its ability to bind specific nuclear proteins. Four distinct nucleoprotein-binding sites were detected, located between positions -7 to -33, -44 to -68, -81 to -103, and -154 to -183, relative to the E3 cap site. These sites contain sequences previously shown to be functionally important for efficient E3 transcription. No major qualitative or quantitative differences were found in the binding pattern between nucleoprotein extracts prepared from uninfected or adenovirus-infected HeLa cells. Competition experiments suggest that the factors binding to the -154 to -183 and -81 to -103 sites are the previously identified nucleoproteins, NFI and AP1, respectively. The factor binding to the -44 to -68 site, which we term ATF, also interacts with other E1A-inducible promoters and is very similar and probably identical to the factor that binds to the cAMP-responsive element of somatostatin. We have purified this factor, which is a protein of 43 kD in size.

[Key Words: Trans-activation; gel retardation; footprinting; nuclear factors]

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The control of the rate of transcription initiation is an important element in the regulation of eukaryotic gene expression. Consequently, considerable effort is being devoted to elucidate the mechanisms by which transcription initiation can be modulated. For a variety of reasons, transcriptional regulation by the adenovirus immediate early E1A gene represents a very good model system for investigating such mechanisms. The E1A gene can transactivate a number of viral and cellular genes, and in many cases the promoters of such genes have been analyzed genetically to locate sequences or elements essential for efficient transcription [for review, see Berk 1986]. The E1A gene can also repress expression of certain viral and cellular genes, specifically those dependent on cis-acting enhancer elements for efficient transcription [Borrelli et al. 1984; Hen et al. 1985; Velcich and Ziff 1985]. Transcription stimulation by the SV40, Py, and E1A [Borrelli et al. 1984; Velcich and Ziff 1985] enhancers, as well as the cellular immunoglobulin heavy chain [Hen et al. 1985] and insulin [Stein and Ziff 1987] enhancers, has been shown to be blocked by E1A protein.

These two activities of E1A appear to be functionally distinct. At early times of infection, the E1A gene produces two transcripts, 12S and 13S in size, which encode products of 243 and 289 amino acids, respectively [Berk and Sharp 1978; Chow et al. 1979; Perricaudet et al. 1979; Kitchingman and Westphal 1980]. Both products display regulatory activity; the product of 289 amino acids can trans-activate and repress gene transcription [Borrelli et al. 1984; Velcich and Ziff 1985], whereas the protein of 243 amino acids, although it represses very efficiently [Borrelli et al. 1984; Velcich and Ziff 1985], is generally found to be deficient in trans-activation [Carllock and Jones 1981; Montell et al. 1982; Svensson and Akusjarvi 1984; Winberg and Shenk 1984; Lillie et al. 1986; Moran et al. 1986; Wu et al. 1986a]. Furthermore, extensive mutational analysis of the E1A gene has identified domains that are required for repression but not for trans-activation [Lillie et al. 1986; Moran et al. 1986; Schneider et al. 1987]. Consequently, these two regulatory activities of E1A have been shown to be separable, but their mode of action remains unclear. It is very unlikely that either trans-activation or repression is accomplished by sequence-specific binding of E1A protein to cis-acting regulatory elements because purified E1A proteins do not bind directly to DNA [Ferguson et al. 1985]. In addition, there is only limited sequence homology between various promoters activated by E1A, and, except for the E1B promoter [Wu et al. 1987], mutational analysis has not led to the identification of sequences specifically required for E1A action; rather, it appears that it is the same promoter elements that are used both for basal level [uninduced] and E1A-induced transcription [for review, see Berk 1986]. Thus, E1A must act in a more indirect fashion, possible by modulating the level or activity of nuclear transcription factors that interact with E1A inducible promoters.

Understanding how E1A trans-activates will therefore require a comprehensive understanding of the E1A-re-
responsive promoters in terms of the DNA sequences required for transcription and the factors that interact with these sequences. With this goal in mind, we have been studying the promoter of the adenovirus early gene E3, which is strongly stimulated by E1A, both in the virus and in DNA transfection experiments. Moreover, promoter mutagenesis and gene transfer experiments have identified elements required for efficient E3 transcription (Leff et al. 1985; Weeks and Jones 1985) and shown that upstream regions of this promoter can confer E1A inducibility on normally insensitive proximal promoter regions (Weeks and Jones 1985; Left and Chambron 1986). In this report we describe four nuclear protein-binding sites, each of which corresponds to a region of the E3 promoter required for efficient transcription. These binding proteins are found in extracts prepared both from uninfected and adenovirus-infected cells. At least one of the factors appears to be very similar to a factor that also interacts with the E1A-inducible E2A and E4 promoters and to promoters that are activated by increasing cAMP levels. This factor has been purified to homogeneity and is a 43-kD protein which we term ATF.

Results

Detection of factors that bind to the E3 promoter

Gene-transfer experiments have shown that E3 promoter sequences extending from the transcriptional start site to the EcoRI site, 237 bp upstream, are sufficient for full promoter activity and induction by E1A protein (Weeks and Jones 1985). Furthermore, a number of important cis-acting regulatory elements have been identified within this region (Leff et al. 1985; Weeks and Jones 1985), the removal or alteration of which results in lower uninduced and E1A-induced levels of transcription. The critical nature of these sequences suggests that they serve as binding sites for sequence-specific DNA-binding proteins. We have analyzed the proteins that interact with the 5'-control region of the E3 gene using the DNase I footprinting [Galas and Schmitz 1978] assay. In these studies we used specific DNA probes that encompass the cis-acting regulatory regions. The precise origin of these probes is described in Materials and methods, but briefly, probe A contains sequences from −37 to −237 and, therefore, has all the sequences necessary for promoter function, except that TATA-box region. Probes B and C do include this region and extend from positions +25 to −85 and from +65 to −77, respectively.

32P-Labeled DNA and crude protein extracts from uninfected and Ads-infected HeLa cell nuclei were incubated in the presence of nonspecific competitor DNA to allow the formation of specific DNA–protein complexes, as described in Materials and methods. These complexes were treated briefly with DNase I, and the resulting digestion patterns were resolved on DNA sequencing gels alongside digestion tracks obtained with naked DNA. The results are summarized in Figure 1. Four distinct protein-binding sites (BS1–4) were detected as revealed by protection from digestion by the DNase I enzyme. The locations of the binding sites are as follows: (1) BS1 extends from −7 to −33 and includes the TATA-box region (−29 to −25) in the area of protection, which is seen using probe B and C labeled on the coding and noncoding strands, respectively. The protection was not as strong as that seen in other areas of the promoter but was observed consistently using both uninfected and infected extracts. (2) Probe A labeled at position −37 on either the coding or noncoding strand revealed three clear footprints. BS2 is located between positions −44 and −68, with the nucleotides at positions −68 and −69, often showing hypersensitivity to digestion. (3) BS3 is farther upstream between nucleotides −81 and −103, with the nucleotide at −104 being consistently hypersensitive to digestion. (4) BS4 is the most upstream site detected and extends from −154 to −183. The exact location of this site was demonstrated more easily using probe A labeled at the EcoRI site at position −237. The site is flanked on both strands by a number of nucleotides hypersensitive to digestion. Binding to this upstream site was especially easy to detect, which may reflect either a very strong DNA–protein interaction or a particularly abundant protein. The four sites all overlap with regions previously shown to be functionally important for transcription from the E3 cap site (see Discussion). The extent of the protected region and the degree of protection obtained was the same using several independent extracts prepared from both uninfected and adenovirus-infected cells and was not affected by the type of nonspecific competitor DNA [poly (dI/dC) or salmon sperm DNA] used.

The binding sites were analyzed further using the methylation protection assay. In these experiments, probe A was end-labeled with 32P on either the coding or noncoding strand, incubated with extracts from uninfected cells, and subsequently treated with the reagent dimethylsulfate (DMS), which methylates guanine residues at position N7 [Maxam and Gilbert 1980]. Those G residues in close proximity to the bound protein are protected from methylation. The DMS-treated DNA from incubations with [bound] and without [free] protein was treated with piperidine, which results in cleavage at methylated bases, and the cleaved DNA was resolved on denaturing acrylamide gels, as shown in Figure 2. Within BS2, G residues at positions −53, −55, and −58 on the coding strand and −57 on the noncoding strand were protected, as there was significantly less cleavage at these positions in the bound versus the free DNA. Similarly, G residues at −90 [coding] and −92 [noncoding] were protected within binding site BS5, and residues at −163, −164, −165, −168, and −170 [coding] and −166, −173, −174 [noncoding] were protected within BS4. The protected residues are indicated by dots above the bases within the binding-site sequences shown in Figure 1. We did not detect any “methylation protection” sites outside the binding sites delineated previously by DNase I footprinting. We have provisionally named the four factors that interact with the E3 promoter E3.F1, E3.F2, E3.F3, and E3.F4.

Nucleoprotein binding to E3 promoter
Different proteins interact with each of the E3-binding sites

Having identified four regions of the E3 promoter that interact with nuclear proteins, we prepared double-stranded DNA oligonucleotides of about 30 bp, containing individual sites BS2, BS3, and BS4 by chemical synthesis of the complementary single strands that were subsequently annealed [see Materials and methods]. In addition, a double-stranded control oligonucleotide (nonsense oligonucleotide) that contained the E3 promoter sequence located between BS3 and BS4 was similarly prepared. By the DNase I footprinting assay, this region did not interact with any nuclear protein. Each of the oligonucleotides was 32P-end-labeled and analyzed for its ability to bind specific nuclear proteins, as revealed by the gel retention assay [Fried and Crothers 1981; Garner and Revzin 1981]. In this assay, specific DNA–protein complexes are allowed to form in the presence of nonspecific competitor DNA, and the complexes are subsequently resolved from free DNA by non-denaturing gel electrophoresis. The BS2, BS3, and BS4 oligonucleotides all bound nuclear protein efficiently, as evidenced by the presence of retarded bands. The specificity of binding was demonstrated by competition experiments; binding to a labeled oligonucleotide was assayed in the presence of an excess of the identical unlabeled oligonucleotide, the nonsense oligonucleotide, or each of the oligonucleotides containing the other binding sites (Fig. 3). In all cases, binding to a labeled oligonucleotide was competed efficiently by an excess of itself but not at all by the nonsense oligonucleotide. Therefore, the binding detected is sequence specific. Oligonucleotides for BS2 and BS3 were also completely unable to compete for binding to BS4, and similarly BS4 did not compete for binding to BS2 and BS3. In contrast however, some competition was detected between BS2 and BS3. For example, unlabeled BS2 oligonucleotide could compete for binding to BS3, although less efficiently than the homologous BS3 oligonucleotide. The results indicate that some relationship between the binding-site specificities of factors E3.F2 and E3.F3 exists. However, the factors are almost certainly different because in both cases, cross-competition between the heterologous sites was not as efficient as competition with the homologous sites. Furthermore, the complexes formed with BS2 migrate slightly faster than the BS3 complex, again implying the involvement of two distinct factors.

Relationship of the E3-binding factors to factors that bind to heterologous promoters

We have examined the ability of other E1A-induced adenovirus early promoter sequences to compete for binding to the E3 promoter, or vice versa. Previous analysis of the E2A and E4 promoters indicated that similar factors might be interacting with specific regions of each promoter [Siva Raman et al. 1986, Lee and Green 1987]. Moreover, these same investigators also observed some slight competition with the E3 promoter, which we investigated further. Initially, we prepared a probe from the E2A promoter (−17 to −98) that contains a binding site for a factor E2A.EF, identified by Siva Raman et al. [1986], and used this in gel-retention assays in the presence of E3 oligonucleotides. The DNA probe used contains a binding site for a second factor E2F, in addition to E2A.EF. However, in this in vitro assay, binding of the second factor is dependent upon the use of salmon sperm DNA as the nonspecific competitor. Under the conditions we use [i.e., poly(dI/dC) as the competitor], only E2A.EF binding is detected [Siva Raman and Thimmappaya 1987; Yee et al. 1987]. As shown in Figure 4A, the BS2 oligonucleotide showed strong competition for binding to the E2A promoter, whereas the BS3 oligonucleotide competed only weakly and the BS4 and nonsense oligomers did not compete at all. This suggests that E3.F2 and E2A.EF are the same factor. Lee and Green [1987] have described a factor, E4.F1, that binds to at least three regions of the E4 promoter and is also competed by E2 promoter sequences. Subsequent competition assays strongly suggest that E3.F2 and E4.F1 are also the same factor [Lee et al. 1987a]. The possibility that all three factors are identical is based predominantly on the fact that all three bind an identical DNA sequence. Consequently, it cannot be ruled out that each represents a different member of a family of factors that have the same DNA-binding specificity. For the sake of discussion, however, we assume that the factors are identical and have renamed them ATF [Lee et al. 1987a].

By comparing the ATF-binding sites in the three adenovirus early promoters, a heptanucleotide consensus sequence T/G A/T CGTCA for ATF binding can be derived [Fig. 5A]. A number of other viral and cellular promoters also contain this same consensus sequence, including genes such as the somatostatin gene, whose expression is activated by raising cAMP levels [Montminy et al. 1986]. In the case of somatostatin, the con-

Figure 1. DNase footprinting of the E3 promoter. [For details of probes, see Materials and methods.] Binding reactions with DNA alone (tracks D) or with extracts derived from uninfected (U) or infected (I) cells were carried out as described in Materials and methods. The products of the DNase reactions were separated on sequencing gels, together with a Maxam and Gilbert A + C ladder for the particular probe. [A] Footprinting in the TATA-box region was examined using probes B (coding strand) and C (nons coding strand). In fact, BS1 and BS2 are present in these probes, but in this panel BS1 binding alone is indicated. [B] Footprinting using probe A filled in at the BamHI site (noncoding strand) covering the E3 promoter region between −37 and −237. [C] Similar to B, but the probe was labeled on the coding strand. [D] Footprinting in the −170 region using probe A labeled on the coding strand at the EcoRI site at −237. The sequences of each of the sites BS1–BS4 are shown. Arrows indicate those bases consistently found to be hypersensitive to DNase digestion. Within BS2–BS4 are G residues whose methylation by DMS was protected by factor binding [see Fig. 2]. These G residues are indicated by dots.
sensus is within an area of the promoter critical for expression and recently shown to bind a specific phosphoprotein (Montminy and Bilezikian 1987). We examined the relationship of this protein to ATF by performing a competition analysis similar to the one performed with the E2A promoter. An end-labeled probe derived from the rat somatostatin gene (extending from −10 to −70), which contains the ATF consensus sequence, was incubated with crude HeLa nuclear extract, and complex formation analyzed by the gel retention assay (Fig. 4B). A single retarded band was resolved that was competed away with the BS2 (ATF) oligomer very efficiently but not with the BS3, BS4, or nonsense oligomers. Therefore, the factor that binds to the somatostatin promoter is very closely related and probably identical to ATF.

The importance to binding of individual nucleotides within the ATF consensus sequence was examined by synthesizing a series of oligomers containing mutations within this sequence and assessing the ability of these mutant oligomers to compete for binding to the wild-type probe in gel-retention assays (Fig. 6). Oligomers containing mutations at positions 3, 4, 5, or 7 within the heptanucleotide consensus all competed less efficiently, showing the critical importance of these sequences in
Figure 3. Competition analysis. (Top) A schematic diagram of the factor-binding sites within the E3 promoter. Each of the double-stranded DNA oligomers was labeled by filling in and was incubated (0.1 ng) with 5 μg of crude nuclear extract under standard conditions (see Materials and methods). For competition analysis, 500-fold molar excess of an unlabeled oligonucleotide was added to each incubation, as indicated. Samples were run on an 8% native gel. (BS2) Binding site 2 oligomer; (BS3) binding site 3 oligomer; (BS4) binding site 4 oligomer; (N) nonsense oligomer.

binding ATF. Surprisingly, mutations at positions 2 and 6 had less of an effect.

The ATF consensus is very similar to the consensus binding site [T G/T AGTCA G/C] for the nuclear factor AP1 [Angel et al. 1987; Lee et al. 1987b]. This factor binds specifically to a number of regulatory regions, including the SV40 and Py enhancers, the human metallothionein IIA (MT IIA) and collagenase promoters, and the rat stromelysin promoter [Angel et al. 1987; Lee et al. 1987b]. Comparison of the ATF- and AP1-binding sites reveals a clear difference at position 3, which is a C residue in the ATF site and an A residue in the AP1 site. One of the mutant BS2 oligomers that we synthesized contains this C → A transversion [BS2/3]; as a result, the mutant oligomer competed for ATF binding poorly, although some competition was still evident. This suggests that AP1 and ATF are different factors. The BS3 sequence, however, does contain the AP1 consensus [see Fig. 5B], and the competition results between BS2 and BS3 are similar to those obtained with the BS2/BS3 mutant and BS2 wild-type oligomers. This raised the possibility that the E3.F3 factor (which binds to BS3) is AP1. This possibility was explored further by additional competition analysis. A double-stranded DNA oligonucleotide containing the SV40 enhancer AP1-binding site [Lee et al. 1987b] was synthesized and tested for its ability to compete for binding to the E3 BS2 and BS3 sites. The results are shown in Figure 7, where the specific complexes are indicated by arrows. (We often detected a faster migrating complex, not competed in a specific fashion, which probably represents nonspecific binding to the oligomer ends.) The competing oligonucleotides were all present at a 100-fold molar excess over the probe. Binding to the BS2 oligomer was competed efficiently by itself but only poorly by the BS3, BS2/3, or AP1 oligomers. In contrast, binding to the BS3 oligomer was competed efficiently by itself, by the BS2/3, and by the AP1 oligomers but not by the BS2 oligomer. Therefore, we conclude that the E3.F3 factor is very closely related and probably identical to the AP1 factor and that AP1 and ATF, although they have related binding sites, are distinct factors.

The E3 BS4 sequence contains homology to the binding site for the nuclear factor NF1, which has the consensus TGGN6_TGCCAA [Leegwater et al. 1985]. A 24-bp oligonucleotide containing the region of the Ad2-inverted terminal repeat that is protected from DNase I digestion by the NF1 factor [Leegwater et al. 1985] was synthesized and tested for competition for binding to the BS4 oligonucleotide. The NF1 oligonucleotide competed very efficiently [data not shown]; therefore, we conclude that the E3.F4 factor is NF1. Furthermore, affinity-purified NF1 [Jones et al. 1987] binds to the BS4 region very efficiently [T. Williams, unpubl.].

The level of the E3-specific factors does not change following adenovirus infection

All four factors that interact with the E3 promoter were clearly present in extracts from uninfected HeLa cells. Nevertheless, their concentration might be altered upon infection by adenovirus, and if one or more of the factors were limiting, an increase in their concentration could be a contributing factor to the activation of this pro-
infected extract was prepared from HeLa cells 6 hr postinfection. If extracts were prepared at later times (20 hr postinfection), three- to fivefold lower levels of the factors binding to the BS2, BS3, and BS4 sites were seen (data not shown). This has been reported previously with respect to binding of ATF to the E2A promoter (Siva Raman et al. 1986) and is probably due to sequestration of factors by the very high copy number of adenovirus DNA template generated by viral DNA replication. Interestingly, binding to the BS1 region was the same between uninfected and late-infected extracts. This may reflect either a higher abundance of this factor at late times of infection or more efficient dissociation from its template during the extraction procedure.

| Promoter   | Position | Sequence          |
|------------|----------|-------------------|
| E3.BS2     | -60 to -53 | TTCGTCAC          |
| E2A        | -70 to -77 | TAAGTCAT          |
| E4         | -50 to -42 | TAAGTCAT          |
| Hsp 70     | -30 to -37 | GTGTCGTCAC        |
| Somatostatin | -42 to -49 | GACGTCAG          |
| c-Fos      | -60 to -67 | TAAGTCAC          |
| **Consensus** |          | TTCGTCGA          |

| Promoter   | Position | Sequence          |
|------------|----------|-------------------|
| E3.BS3     | -87 to -94 | TTAAGTCAT         |
| SV40       | 120 to 113 | TTAAGTCAG         |
| Polyoma    | 5120 to 5112 | TTAAGTCAC       |
| Collagenase| -72 to -65 | TGAGTCAG          |
| Stromolysin| -71 to -66 | TGAGTCAG          |
| MT IIA     | -98 to -105 | TGAGTCAC         |
| **Consensus** |          | TTAAGTCAG         |

Figure 4. Comparison with other promoters. (A) Approximately 1.0 ng of the E2A probe (see Materials and methods) was incubated with 5 μg of nuclear extract from uninfected cells and 1 μg poly[dI/dC]. For competition analysis, 135-fold molar excess of unlabeled oligonucleotides was included as indicated, and the binding reactions were run on an 8% gel. (B) Approximately 0.2 ng of the somatostatin probe (see Materials and methods) were incubated with 5 μg of nuclear extract and 1 μg poly [dI/dC]. For competition analysis, a 250-fold molar excess of unlabeled oligonucleotides was added to the incubations as indicated. The samples were run on an 8% gel. (See legend to Fig. 3 for abbreviations.)

Figure 5. Comparison of factor-binding sites. (A) The heptanucleotide core sequence within BS2 is shown compared to homologous sequences found within factor-binding sites characterized for other promoters: E4 (Lee and Green 1987), E2A (Siva Raman et al. 1986), Hsp 70 (Wu et al. 1986b), somatostatin (Montminy et al. 1986), c-fos (Gilman et al. 1986). Comparison of these sequences leads to a consensus ATF-binding site, as shown. (B) Comparison of the core sequences within known AP1-binding sites and the E3 BS3 site. SV40 and metallothionein IIA (MT IIA) (Angel et al. 1987; Lee et al. 1987b); polyoma, collagenase, and stromolysin (Angel et al. 1987). Where binding factors have been mapped to these sequences using methylation protection analysis, the protected Gs are shown by dots. Gs protected on the opposite strand are indicated by a dot below the pairing C residue.
BS2 oligonucleotide as probe. Peak binding fractions were analyzed further by DNase I footprinting on probe A, which showed the gradual loss of BS3 (AP1)- and BS4 (NF1)-binding activities, whereas the BS2 (ATF) activity was retained and remained the only E3-binding activity in the affinity column eluant (data not shown). A portion of the peak fraction from the oligonucleotide affinity column was subjected to SDS gel electrophoresis, and separated proteins were revealed by silver staining. As shown in Figure 9B, the affinity column eluant contained a major band at 43 kD and an additional series of bands between 60 and 65 kD. These latter bands were detected in all the tracks, including the control lane that contained sample buffer only. Therefore, we assume that these bands are nonspecific artifacts. The protein running as a 43-kD species is therefore likely to be ATF. To determine unambiguously that indeed this species is the active binding protein, we subjected some of the affinity-purified protein to preparative SDS gel electrophoresis. Regions containing the 43-kD and 60 to 65-kD species were excised and eluted, and the resulting renatured proteins tested for the ability to bind to the BS2 oligonucleotide.
Figure 8. Titration of uninfected and infected cell extracts. Each panel shows a gel-retention assay with an oligomer probe representing one of the E3 factor-binding sites, titrated with nuclear extracts derived from uninfected (lanes 1–5) and Ad5 early infected (lanes 6–10) cells. (A,C). Incubations in lanes 1, 2, 3, 4, and 5 and lanes 6, 7, 8, 9, and 10 contained 2, 4, 6, 8, and 10 μg crude nuclear protein, respectively. (B,D). Incubations in lanes 1, 2, 3, 4, and 5 and lanes 6, 7, 8, 9, and 10 contained 1, 2, 3, 4, and 5 μg crude nuclear protein, respectively. All incubations included 0.1 ng of the relevant end-labeled oligomer and poly(dI/dC) at a ratio of 1 μg/2.5 μg of crude protein. Samples were run on 8% native gels.

oligomer. As shown in Figure 9C, incubation with the 43-kD protein resulted in retardation of the BS2 probe. We therefore conclude that the 43-kD protein is the ATF factor.

Discussion

We have investigated the DNA sequences upstream of the E3 transcriptional start site for their capacity to interact specifically with nuclear proteins, using the DNase I footprinting technique. The results summarized in Figure 10 show that there are four distinct nuclear protein-binding sites. Each of these binding sites encompasses sequences that previously had been shown to be important for efficient transcription initiation from this promoter: (1) The most proximal binding site (−7 to −33) contains the TATA-box sequences. A deletion of this region extending from nucleotide −15 to −35 was found to lower E3 transcription approximately fivefold [Leff et al. 1985]. However, we have no direct evidence to suggest that the factor BS1, which we detect binding in this region, specifically recognizes the TATA box itself. (2) The next proximal binding site (−44 to −68) encompasses the nucleotides −55 to −57 found to be critical for promoter functions; a deletion from −39 to −55 was found to have relatively little effect on transcription, whereas a deletion from −39 to −57 reduced transcription significantly [Leff et al. 1985]. Consistent with these results, we found that a mutation at position 56 (BS2/BS5, Fig. 6) severely affected binding of ATF to this region of the promoter. Surprisingly, however, a mutation at position 54 (BS2/BS7) also diminished binding, although not quite as severely; this residue would be removed in the −39/−55 deletion mutant.
Nucleoprotein binding to E3 promoter

Figure 9. Purification of ATF. [A] Scheme used for the purification of ATF. Precise details of the preparation and use of the columns are given in Materials and methods. [B-70] Protein from the peak fractions of the Biorex-70 column; [DNA] peak fraction from the nonspecific DNA column; [affinity] protein after one pass over the BS2 oligomer affinity column; [C] control lane containing sample buffer only. [C] Gel retention analysis of ~100 ng of affinity-purified proteins eluted from the 43 and 60K regions of a preparative gel after denaturation and renaturation (see Materials and methods). Incubations contained 0.1 ng of end-labeled BS2. Lane C had 5 μg crude extract and 1 μg poly[dI/dC]. For each protein sample, lanes 1–3 represent one-tenth the volume of successive fractions (150 μL) collected from the renaturation gel filtration column. Samples were run on an 8% gel.

Figure 10. Summary of nuclear factors interacting with the E3 promoter. Shown is a schematic representation of the site of interaction of four nuclear factors that we have found to recognize the adenovirus E3 promoter. Two of the factors, namely AP1 and ATF, are implicated as mediators of inducers of gene transcription. The relevance of these inducers (TPA and cAMP) to E3 transcription and E1A transactivation remains to be established.

Additional experiments that fully explore the effect of point mutations on ATF binding and E3 transcription are in progress. [3] The binding site located between −81 and −102 coincides almost exactly with a functional promoter element defined by 5′-deletion analysis of the promoter [Weeks and Jones 1985]. Wild-type levels of transcription were obtained with upstream sequences extending to the EcoRI restriction site at position −237. Deletions 5′ to position −105 resulted in a small (two- to fourfold) reduction in transcription efficiency, but a further deletion to position −82 lowered transcription drastically, indicating the presence of a minor promoter element between −105 and −228 and a major promoter.
element between −82 and −105. The major element thus coincides with BS3. [4] BS4 probably coincides with the minor promoter element upstream of position −105, although the functionally important sequences in this region have not been defined precisely as yet. No proteins were found to interact with sequences that were neutral with respect to promoter function. Both qualitatively and quantitatively, the same nuclear protein–DNA interactions were observed regardless of whether the nuclear protein extracts were isolated from uninfected or adenovirus-infected HeLa cells. It is unlikely, therefore, that E1A stimulates E3 transcription by increasing the levels of limiting factors.

In all of these analyses it has been noted that deletion of the functional element did not abolish stimulation by the E1A protein, but rather the deletions lowered both uninduced and E1A-induced transcription, suggesting that the same elements and therefore presumably the same nuclear proteins are required for basal and induced E3 expression. However, the degree of stimulation by E1A was lower when either BS2 or BS3 was removed (Leff et al. 1985; Weeks and Jones 1985), possibly indicating that cooperation between the factors that interact with these two sites is necessary for full stimulation.

The factor that binds to the −44 to −68 region of E3, which we now call ATF, also binds to critically important regions of the E2A and E4 promoters (Lee et al. 1987a). It is possible therefore that this factor plays a crucial role in E1A trans-activation of these early genes. The human Hsp70 promoter is also induced by E1A (Wu et al. 1986a), and examination of the promoter sequence reveals a region just upstream of the TATA box that is identical to the ATF consensus sequence. Therefore, it is interesting to speculate that this site also binds the same factor, and indeed binding of a nuclear protein to this site has been detected (Greene et al. 1987). A number of other cellular and viral promoters contain the same consensus sequence (Fig. 5A). The c-fos promoter contains the consensus, and a protein that binds to this sequence has been detected (Gilman et al. 1986). Importantly the close contact points (as defined by methylation protection assays) within the c-fos-binding site are identical to those found within the E3 BS2, making it very likely that the same protein is binding to each site. The site is also found in a functionally critical region of the somatostatin gene, which appears to be important for its inducibility by cAMP (Montminy et al. 1986). We show here that the BS2 oligomer competes efficiently for binding of a factor to this region of the somatostatin promoter, again indicating the very close relationship of this factor and ATF. This conclusion is substantiated further by our purification of ATF, revealing it to be a 43-kD protein, identical in size to the phosphoprotein purified from the neuronal cell line PC12, which binds to the somatostatin sequence (Montminy and Bilezikjian 1987). A number of other genes inducible by cAMP contain the same consensus sequence. The interesting possibility that emerges, therefore, is that a common factor is at least partially mediating activation by cAMP and by E1A. Initial experiments indicate that raising cAMP levels by treating HeLa cells with forskolin does not lead to activation of the E3 promoter (J. Jordan, H. Hurst, and N. Jones, unpubl.). This might not be too unexpected, however, because it appears that although the consensus sequence is certainly a component of the cAMP response of the somatostatin gene, it is not in itself sufficient for cAMP inducibility (Montminy et al. 1986). It has been shown recently that increasing cAMP levels result in an increase in the phosphorylation of the 43-kD factor that interacts with the somatostatin promoter (Montminy and Bilezikjian 1987). It would be of obvious interest to determine whether such an increase also occurs in response to the action of E1A.

The competition experiments strongly indicate that the factor that binds to the E3 BS3 is the previously identified factor AP1, and affinity-purified AP1 (Lee et al. 1987b) binds efficiently to this site (T. Williams, unpubl.). The AP1-binding site is found in a number of promoters that can be induced by phorbol esters such as 12-O-tetradecanoylphorbol-13-acetate (TPA), and a number of lines of evidence suggest that AP1 mediates this TPA response. For example, synthetic copies of the AP1-binding site can confer TPA inducibility upon heterologous promoters (Angel et al. 1987; Lee et al. 1987b).

Our initial results indicate that the E3 promoter can be activated by TPA, although to a lesser degree than with E1A ([J. Jordan, H. Hurst, and N. Jones, unpubl.]), we have not shown that this TPA response requires that E3.F3 (AP1) factor, but this would seem likely to be the case. The mechanism by which AP1 mediates this response to TPA is not clear, but because TPA activates protein kinase C it is tempting to speculate that as a result of TPA action, the phosphorylation state of the AP1 factor is increased. Likewise, it is possible that an increase in phosphorylation could also result from E1A action, a possibility that is currently being tested.

Both the ATF and AP1 factors appear to be highly conserved because we have found that factors with identical DNA-binding specificities are present in the fission yeast, Schizosaccharomyces pombe (R. Jones and N. Jones, unpubl.). Moreover, promoters such as E2A and E3, which normally require E1A transactivation to function efficiently, are active in S. pombe cells in the absence of E1A. It is interesting to speculate, therefore, that if E1A is mediating its response through specific activation of these factors, the factors are constitutively present in their “active” state in S. pombe. In contrast, the E3.F4 factor, which we have identified as being NF1, does not appear to be present in these yeast cells.

The mechanism by which E1A protein trans-activates is still unclear. It is very unlikely to act directly because isolated E1A protein lacks the ability to bind to DNA (Ferguson et al. 1985). It is more likely that it acts through the cellular transcription factors that interact with the inducible promoters. One possibility is that E1A induces a modification of these factors to increase their capacity for promoting transcription. Alternatively, E1A could result in an increase in the amount of a particular factor, and if that factor happened to be limiting, an increase in transcription would result. The
early gene E2A promoter has been examined in some detail. In addition to ATF, a second factor, E2F, also interacts with critically important sequences of the E2A promoter (Kovesdi et al. 1986). This factor can be detected by in vitro-binding assays when salmon sperm DNA is used as the nonspecific DNA. When poly(dI/dC) is used instead, ATF binding and not E2F is predominantly detected (Siva Raman and Thimmappaya 1987; Yee et al. 1987). Interestingly, E2F DNA-binding activity is markedly increased upon infection with adenovirus, suggesting that activation of the E2A promoter by E1A is at least partially due to activation of the E2F factor (Kovesdi et al. 1986). It is not clear at present whether this activation is due to an increase in the actual level of E2F or a modification of a preexisting factor that markedly alters its DNA-binding properties. Recent evidence demonstrating the activation of E2A transcription in vitro by the addition of purified E1A protein suggests that the latter possibility is more likely (Spangler et al. 1987). The E2F factor appears to be unique among the factors that interact with the early promoters, because all the others, including those that bind to the E3 promoter, can be readily detected in extracts from uninfected cells. In these cases, it is possible that E1A-induced modification could affect some activity of the factor other than DNA binding. This may involve the physical relocation of factors within the nucleus, as has been suggested by Lee and Green (1987) who find that in vitro extracts from uninfected cells are as active in E4 transcription as infected whole cells. Alternatively, factor modification may alter interactions with other transcription factors and RNA polymerase II, so leading to the formation of stable initiation complexes. Obviously, a greater understanding of the functions of transcription factors, in general, and particularly those that interact with E1A-inducible promoters is required to elucidate fully this phenomenon. Such an understanding can only come from the purification and subsequent biochemical characterization of these factors.

Materials and methods

Cells and extracts

HeLa cells were grown in suspension in 2% RPMI with 5% fetal calf serum and harvested at 5 × 10⁸ to 8 × 10⁸ cells/ml. Cells for infection were concentrated 10-fold and incubated with wild-type Ad5 at 20 pfu/cell for 1 hr at 37°C before dilution back to 5 × 10⁸ cells/ml for a further 6 hr (early infection) or 18 hr (late infection) before harvesting.

Nuclear extracts were prepared from pelleted cells by the method of Dignam et al. (1983). The proteins were usually concentrated by addition of ammonium sulfate to 0.33 g/ml, with slow stirring for 30 min at 4°C. The precipitate was collected by centrifugation at 25,000g and resuspended at 10–25 mg/ml before dialysis versus three changes of dialysis buffer: 20 mM HEPES–KOH (pH 7.9), 20% glycerol, 20 mM KCl, 1 mM MgCl₂, 0.5 mM dithiothreitol (DTT), 0.5 mM PMSF. Extracts were stored in liquid nitrogen, and protein concentrations estimated using the BioRad assay kit versus plasma γ-globulin standards. Extracts prepared from adenovirus-infected cells were checked for the presence of E1A protein using Western blotting analysis (Feldman et al. 1982), with an E1A polyclonal rabbit antiserum raised against E1A expressed in Escherichia coli (Ferguson et al. 1984) and the Amersham horseradish peroxidase detection system.

Nucleoprotein binding to E3 promoter

E3 probes A, B, and C were derived from subclones of E3 promoter sequences. The 230-bp Smal fragment between –37 and –268 was cloned into the Smal site of pUC8. This clone was cleaved at the BamHI site (proximal to –37) within the pUC8 polylinker and either incubated with [α-32P]dATP and reverse transcriptase to label the noncoding strand or treated with alkaline phosphatase followed by T4 kinase and [γ-32P]ATP to label the coding strand. Subsequent cleavage with EcoRI released the 200-bp fragment designated probe A. Alternatively, the probe was labeled at the EcoRI site by filling in. Probes B and C were derived from an E3 promoter deletion mutant described by Weeks and Jones (1985), which was prepared by converting the Ddel (–85)–SalI (–65) restriction fragments into a BamHI fragment with linkers. Therefore, BamHI cleavage and incubation with reverse transcriptase and [α-32P]dATP generated either probe B (labeled on the coding strand) after Sall (–25) digestion or probe C (labeled on the noncoding strand) after Hpal (–77) digestion.

The E2A probe was obtained as described by Siva Raman et al. (1986) by labeling at the BssHII site (at –17) using reverse transcriptase and [α-32P]dCTP and subsequent digestion with NarI (at –98). The rat somatostatin probe (Tavianini et al. 1984) was obtained by labeling at the BgllI site (at –70) using reverse transcriptase and [α-32P]dATP followed by Sall (–10) digestion to release a 60-bp probe. All of the end-labeled probes were purified by acrylamide gel electrophoresis and electrolution. The isolated probes were ethanol-precipitated without the addition of carrier and recovered by ultracentrifugation at 100,000g.

A + G marker tracks were prepared by treating labeled probes with 3% formic acid followed by piperidine cleavage (Maxam and Gilbert 1980).

DNase footprinting and methylation protection assays

The exact conditions for DNase footprinting were determined empirically for each probe and each extract examined. Binding reactions of 45 µl in DNase buffer [20 mM HEPES–KOH (pH 7.9), 50 mM KCl, 20% glycerol, 2 mM DTT, 2 mM MgCl₂] contained 150–200 µg of crude nuclear extract, 2–4 µg nonspecific competitor, and –0.3 ng end-labeled DNA probe. Incubations containing probe alone were also prepared. After 20 min at room temperature, 3–5 µl of DNase I [Amersham, diluted to 5 µg/ml in DNase buffer, was added for 30–60 sec. For “DNA only” incubations, DNase I was diluted to 0.1 µg/ml. The reaction was stopped by addition of 50 µl of proteinase K stop [buffer] [100 mM Tris–Cl (pH 7.5), 2% SDS, 20 mM EDTA, 400 µg/ml proteinase K] for 10 min at 37°C. Samples were extracted once with 100 µl phenol mix [phenol, chloroform, isoamyl alcohol at 25 : 24 : 1] and precipitated with tRNA carrier. The dried pellets were Chervenkov counted, and equal counts loaded in all tracks on a standard 50% urea, 10% acrylamide sequencing gel, together with an appropriate A + G marker lane.

For methylation protection assays, 45-µl binding incubations were set up using probe A as for DNase footprinting, and after 20 min, each was treated with 2 µl DMS (diluted fivefold in water) for 45–60 sec. The reaction was stopped by the addition
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of 1 μl β-mercaptoethanol and 50 μl of proteinase K stop buffer, as above. Methylation of free probe was carried out according to Maxam and Gilbert (1980). The labeled DNA recovered from the protein bound and free probe samples was then cleaved by piperidine, and equal Cherenkov counts from bound and free samples run side by side on a standard 50% urea, 10% acrylamide sequencing gel.

Gel retardation

Binding reactions between nuclear extracts and end-labeled probes were carried out in 20- to 25 μl mixes in gel-retention buffer (GRB): 25 mM HEPES–KOH (pH 7.9), 1 mM EDTA, 5 mM DTT, 10% glycerol, 150 mM NaCl. Typically, these incubations contained ~0.1 ng probe, 5–10 μg crude protein extract, and 1–2 μg of nonspecific DNA competitor: either poly (dI/dC) (Pharmacia) or sonicated salmon sperm DNA (Sigma). Binding mixes were incubated for 20 min at 20°C and loaded directly onto 6–8% polyacrylamide gels (acyr : bis = 44 : 0.8) and run in 0.5 x TBE for 90 min at 200 V. Gels were then fixed and dried for autoradiography.

Oligonucleotides

We are grateful to Iain Goldsmith for preparing a series of oligonucleotides for us. The deprotected oligonucleotides were ethanol-precipitated, and the dried pellets resuspended in 98% formamide, 10 mM EDTA, and incubated for 2 min at 90°C, and loaded onto a gel 1.5 mM thick, containing 50% urea, and 10% acrylamide. After electrophoresis at 60 mA in 1 x TBE the full-length oligonucleotides were located by UV shadowing, excised from the gel, and eluted overnight (Maxam and Gilbert 1980). The purified oligomers were annealed as described (Kadonaga and Tjian 1986) to generate short double-stranded DNA regions containing a nucleoprotein-binding site.

Sequences contained within each double-stranded DNA oligomer, with respect to the E3 promoter, were as follows: BS1 oligomer (-32 to -7); BS2 oligomer (-44 to -67); BS3 oligomer (-82 to -103); BS4 oligomer (-157 to -180); nonsense oligomer (-107 to -128). All were constructed to give BamHI overhangs to the double-stranded DNA oligonucleotides. Incubations containing [α-32P]dATP and reverse transcriptase were used to label the oligomers for use as probes.

Preparation of nonspecific and specific DNA affinity columns

In each case, DNA was coupled to CNBr-activated Sepharose 4B (Pharmacia), which had been hydrated and washed extensively in 1 mM HCl. Coupling was carried out overnight at room temperature in 10 mM Na phosphate buffer (pH 8.0) on a rotary shaker. After coupling, residual active groups in the resin were deactivated by incubation for 2–3 hr at room temperature with 1 M ethanolamine–HCl (pH 9.0). Finally, the coupled resin was washed successively with 10 mM Na phosphate (pH 8.0), 1 M KCl, and oligonucleotide column storage buffer [Kadonaga and Tjian 1986: 10 mM Tris (pH 7.5), 0.3 M NaCl, 1 mM EDTA, 0.02% azide].

DNA for the nonspecific column was derived as follows: Calf thymus double-stranded DNA (Sigma) was dissolved in water at 10 mg/ml, sonicated briefly, and passed successively through 19 and 21-gauge needles. The DNA was then boiled for 10 min and quick-cooled on ice. Next, 20 mg of this DNA was coupled per 5 grams of resin, and the efficiency was 70–80%, giving a final concentration of ~0.75 mg DNA/ml of resin.

Oligonucleotide was prepared for the affinity column, largely as described by Kadonaga and Tjian (1986). End-labeled, ligated, double-stranded BS2 oligomer (150μg) was incubated with 6 ml of resin. After coupling, the residual counts in the supernatant gave an estimated coupling efficiency of 80%, that is, 20 μg double-stranded oligomer/ml of gel.

Purification of ATF

Each stage of the purification was monitored by gel-retention assay of the fractions, using BS2 oligomer as probe to determine the peak of binding activity. Cleared crude extract (10 ml at 10 mg/ml protein) was chromatographed over a 50-ml Biorex-70 column [200–400 mesh, BioRad], equilibrated with 0.2 M KCl CB [column buffer: 20 mM HEPES (pH 7.9), 20% glycerol, 1 mM EDTA, 0.1% NP40, 0.5 mM DTT, 0.5 mM PMSF, 0.1 mg/ml leupeptin], and washed extensively with this buffer. The column was developed with successive step increases in KCl concentration to 0.3 M, 0.45 M, 0.6 M, and 0.75 M. BS2 activity eluted at 0.45 M KCl, with an approximate fivefold increase in specific activity. The pooled fractions were dialyzed to 0.2 M KCl CB and applied to a 35-ml DNA–Sepharose column equilibrated with the same buffer. After extensive washing, ATF was eluted in one step with 0.4 M KCl CB, with a further increase in specific activity of three- to fourfold. These fractions were diluted to 0.2 M KCl CB and applied to a 5-ml BS2 oligomer affinity column equilibrated with the same buffer also containing 0.1 μM ZnCl2 [Lee et al. 1987b]. The bound factor was washed and then eluted in one step with 0.6 M CB. Flow of the column was interrupted for 10 min to allow dissociation before collection of the elution fractions.

Fractions from each step of the purification were run on standard 9% acrylamide–protein gels. Elution fractions were precipitated with 20% trichloroacetic acid (TCA) and resuspended in sample buffer [31.25 mM Tris–Cl (pH 6.8), 1% SDS, 5% glycerol, 2.5% β-mercaptoethanol, 0.0005% Pyronin Y]. Silver staining was performed as described by Sammons et al. (1981), and estimates of the concentration of ATF gave an approximate 1500-fold increase in specific activity for the affinity column steps, that is, over 20,000-fold purification over all.

Renaturation of ATF-binding activity from a preparative SDS gel was carried out as described by Lee et al. [1987b], except that the denaturation and renaturation buffer was CB with 0.15 M KCl. The net recovery of specific binding activity was ~3%.

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