Pip, a novel IRF family member, is a lymphoid-specific, PU.1-dependent transcriptional activator

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The immunoglobulin light-chain gene enhancers Eκ3, Eκ2-4, and Eκ3-1 contain a conserved cell type-specific composite element essential for their activities. This element binds a B cell-specific heterodimeric protein complex that consists of the Ets family member PU.1 and a second factor (NF-EMS), whose participation in the formation of the complex is dependent on the presence of DNA-bound PU.1. In this report we describe the cloning and characterization of Pip (PU.1 interaction partner), a lymphoid-specific protein that is most likely NF-EMS. As expected, the Pip protein binds the composite element only in the presence of PU.1; furthermore, the formation of this ternary complex is critically dependent on phosphorylation of PU.1 at serine-148. The Pip gene is expressed specifically in lymphoid tissues in both B- and T-cell lines. When coexpressed in NIH-3T3 cells, Pip and PU.1 function as mutually dependent transcription activators of the composite element. The amino-terminal DNA-binding domain of Pip exhibits a high degree of homology to the DNA-binding domains of members of the interferon regulatory factor (IRF) family, which includes IRF-1, IRF-2, ICSBP, and ISGF3γ.

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The expression of immunoglobulin genes, which encode the heavy- and light-chain subunits of the antibody molecule, is tightly regulated in a cell type- and developmental stage-specific fashion. Expression of these genes in B cells is controlled both at the level of transcription and recombination. Multiple cis-regulatory elements, promoters, and enhancers appear to confer proper developmental regulation of transcription (Calame and Eaton 1988; Sen and Baltimore 1989; Staudt and Lenardo 1991). Immunoglobulin gene transcriptional enhancers also play a critical role in regulating V(D)J recombination (Ferrier et al. 1990; Engler et al. 1991; Takeda et al. 1993). Cell type-specific enhancers were first identified in the J-C introns of both the heavy (Banerji et al. 1983; Gillies et al. 1983; Neuberger 1983) and κ light chain (Queen and Baltimore 1983; Picard and Schaffner 1984) genes. More recently, additional enhancers have been identified 3′ of the constant region genes in both the heavy (Matthias and Baltimore 1993; Pettersson et al. 1990) and κ light chain (Meyer and Neuberger 1989) loci. The λ light-chain gene locus, which is organizationally distinct (Storb et al. 1989), possesses two transcriptional enhancers, each located 3′ of a Jκ-Cκ gene cluster (Hagman et al. 1990). These two enhancers, designated Eκ2-4 and Eκ3-1, are >90% homologous, are thought to have arisen by an evolutionary gene duplication event, and presumably function in a similar manner.

We have demonstrated previously that the Eκ2-4 enhancer contains two domains, AA and AB, which are essential for enhancer activity; mutation or deletion of either domain completely destroys enhancer function (Rudin and Storb 1992; Eisenbeis et al. 1993). Both AA and AB, which appear to be unrelated, have been shown to bind B cell-specific factors in nuclear extracts (Rudin and Storb 1992). The λB element consists of two juxtaposed but distinct transcription factor-binding sites (Eisenbeis et al. 1993). One site of the composite element is bound by PU.1, a member of the ETS family of transcription factors, that has been shown to be required for the development of lymphoid and myeloid lineages (Klemsz et al. 1990; Scott et al. 1994). The other site is likely bound by NF-EMS, a B cell-specific factor initially shown to form a ternary complex with PU.1 and a domain of the immunoglobulin Eκ3 enhancer (Pongubala et al. 1992). The λB and κ3′ domains are homologous composite elements. Both sites of the composite element are required for enhancer function (Jude and Max 1992; Pongubala et al. 1992; Eisenbeis et al. 1993). Intriguingly, whereas PU.1 can bind the composite element on its own, NF-EMS is unable to do so (Pongubala et al. 1992; Eisenbeis...
ET al. 1993). NF-EM5 is recruited onto the DNA by a specific interaction with PU.1. This protein-protein interaction is critically dependent on phosphorylation of PU.1 at serine residue 148 (Pongubala et al. 1993).

In this report we describe the cloning and characterization of a novel gene, Pip (PU.1 interaction partner) that encodes a PU.1-interacting protein with the properties expected of NF-EM5. PU.1 and Pip represent mutually dependent activators of the λB and κ3′ enhancer elements.

Results

Isolation of the Pip gene

Of the many B-cell proteins that bind specifically to the λA and λB elements in vitro (Rudin and Storb 1992), only PU.1 represents a cloned transcription factor (Eisenbeis et al. 1993). In an effort to clone other factors that bind to the λB elements of the Ex2-4 enhancer, we generated a gtl 1 cDNA expression library from poly(A)+ RNA extracted from J558L, a murine λ1-producing myeloma cell line. We screened the library using an in situ filter-binding assay (Singh et al. 1988) and concatenated λB probes. A screen of 1.2 × 10^6 plaques identified three short clones (all <800 bp). Sequence analysis confirmed that the three clones, designated λ13, λ15, and λ37, contained extensively overlapping cDNA inserts that represented a portion of an incomplete open reading frame (ORF). The three clones were predicted to code for 134, 186, and 188 amino acids, respectively, with presumed DNA-binding activity.

To determine the DNA-binding characteristics of the cloned proteins, extracts were made from Escherichia coli lysogens harboring the λ13 and λ15 clones. These extracts were tested in electrophoretic mobility shift assays (EMSA) for their ability to bind to the λB site (Fig. 1A). Although there is no λB-binding activity present in a λgt1 control lysogen extract, the λ13 extract contains a protein that binds the λB element, forming a shifted complex in the gel. A similar result was obtained with the λ15 extract (not shown), confirming that these two phage clones contain cDNA inserts that code for a protein that binds the λB enhancer element in vitro.

Mapping the DNA binding site for Pip

To further define the binding site for the λ13-encoded protein, we tested the ability of the λ13 lysogen extract to bind a series of mutant λB probes in an EMSA (Fig. 1B, C). Not only does the protein encoded by the λ13 clone bind the wild-type λB element, but it also binds, albeit less tightly, a λB element containing a 4-base alteration in the core of the PU.1-binding site (Fig. 1B, C). This protein is unable, however, to bind to probes containing mutations in the region 3′ of the PU.1-binding site (Bm2 and Bm3, Fig. 1C) or to a shorter probe in which a portion of this 3′ region has been deleted (Ets, Fig. 1C). These results suggest that unlike PU.1, whose binding requires the AGGA in the Ets consensus sequence core but not the 3′ sequence that is mutated in the Bm2 and Bm3 probes (Eisenbeis et al. 1993), the λ13 protein binds the sequence 3′ of the PU.1 site.

To identify guanine residues involved in contacting the protein, a methylation interference assay was performed using the λ13 lysogen extract and a methylated λB site probe. We analyzed the top (Fig. 2A) and bottom (Fig. 2B) strands of the λB probe; the results are summarized in Fig. 2C. On the top strand, methylation of the first guanine 3′ of the AGGA Ets consensus sequence core partially interferes with complex formation (shown as an open circle in Fig. 2C). Methylation of the next guanine completely interferes with the ability of the λ13 protein to bind the λB DNA (shown as a solid circle, Fig. 2C), indicating that the λ13 protein comes into close contact with the major groove of the DNA at this position. On the bottom strand only two adjacent guanines exist for analysis, and these lie 3′ of the PU.1-binding site. Methylation of either of these G's abolished λ13

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Figure 1. The λ13-encoded protein binds λB specifically. (A) Gel mobility shift assay using the B probe and lysogen extracts from the parent vector [gtll] and the λ13 clone. (B) Gel mobility shift assay using the λ13 lysogen extract and a series of kB probes. (C) Summary of the binding characteristics of the λ13 clone. The sequences of the probes used in the gel shift analysis are shown. Dashes indicate residues that are identical to those at the analogous position in the B probe. The ability of the λ13-encoded protein Pip to bind to each probe is indicated.
protein binding, again suggesting that the λ13 protein makes intimate contacts in the major groove at these two positions. The interference patterns delimit a minimal binding site for the λ13 protein, which is adjacent to that determined previously for PU.1 and is identical to that determined previously for the 3′-binding factor in the JB2 complex (Fig. 2C; Eisenbeis et al. 1993).

These binding site experiments demonstrate that the protein encoded by the λ13 cDNA insert is capable of binding the AB element at a site adjacent to the PU1-binding site and may therefore be a component of the JB2 complex described previously (Eisenbeis et al. 1993), which contains a factor (NF-EM5) that binds this sequence. Interestingly, in EMSAs with J558L nuclear extract, we have not observed a binding activity that interacts with this site in the absence of simultaneously bound PU.1. These observations raised the possibility that whereas the truncated cloned protein could bind the AB element on its own, the full-length native molecule requires PU.1 for interaction with DNA.

Isolating a complete Pip ORF

To investigate the possibility of isolating a complete ORF, we screened the same J558L cDNA library with the λ13 cDNA insert as a probe to obtain the complete cDNA sequence. A screen of 1×10⁶ plaques identified ~200 clones containing inserts with homology to λ13. Twenty of these were purified, and the four clones with the largest cDNA inserts were subcloned for sequencing. The longest of the clones, λ13.9, had an insert of 1783 bp (sequence submitted to GenBank), which contained a 64-bp 5′-untranslated region (UTR), a long ORF of 1353 bp (65–1417), and a 365-bp 3′ UTR. Because no poly(A) addition signal was found, this is unlikely to be the complete 3′ UTR. The deduced ORF codes for a protein of 450 amino acids with a predicted molecular mass of 51.5 kD. We have named this protein Pip (see below). All of the sequenced clones (including the short λ13, λ15, and λ37 clones) share nearly identical 5′ ends. Thus, we believe that the 5′ end of the sequence is at or near the cap site. The sequence surrounding the putative initiator methionine is consistent with the loosely defined consensus sequence for eukaryotic translational initiation (Kozak 1987). In addition, the 64 bp of 5′ UTR contains two translational stops in-frame with the coding sequence, suggesting that this is the true start of the ORF. It should be noted, however, that a second methionine occurs as the fourteenth amino acid in this sequence (Fig. 3A), and we have not ruled out the possibility that this may be an alternative translational start.

The shortest of the original clones, λ13, codes for the first 134 amino acids of the Pip protein. Because this clone was identified by its ability to bind specifically to the AB element in vitro, it localizes the DNA-binding domain to the first 134 amino acids of the Pip protein. Interestingly, this portion of the Pip protein sequence exhibits a considerable degree of sequence homology to the DNA-binding domains of the interferon consensus sequence-binding protein (ICSBP, Driggers et al. 1990) and other, related members of the interferon regulatory factor family (IRF) (Fig. 3A). In addition, Pip shares a series of small blocks of homology with ICSBP at similar positions in the carboxy-terminal half of each protein (Fig. 3B).

The amino-terminal half of the Pip protein is highly hydrophilic and is separated from the carboxy-terminal half by a potential PEST domain (a region rich in proline, glutamate, serine, and threonine residues whose presence in intracellular proteins correlates with a short half-life; Rogers et al. 1986). This 31-amino-acid domain (208–238) contains 45% P, E, S, and T and is flanked by basic amino acids. The region of the Pip protein that

Figure 2. Methylation interference analysis of the λ13-encoded DNA-binding protein. Analysis of the upper (A) and lower (B) strands of the B probe is shown. The λ13 lysogen complex was isolated and analyzed. (B) The complexed probe; (F) the free probe recovered from the native gel. Corresponding sequences are shown for reference. (C) Summary of the methylation interference analysis of the λ13 complex in comparison to methylation interference results for the JB2 and PU.1 complexes (Eisenbeis et al. 1993). (Pip) The λ13 complex. Sites of close contact between protein and DNA upon complex formation are indicated. (●) G residues whose methylation results in a complete loss of complex formation; (○) Residues whose methylation results in a substantial decrease in complex formation; (□) residues whose methylation results in a modest decrease in complex formation. Protein binding sites are boxed.
contains this putative PEST domain is in general proline-rich (151–237, 20% proline), representing a potential transcriptional activation domain (Mermod et al. 1989). In addition, there is a region in the carboxyl terminus of Pip that is rich in glutamine residues (354–419, 15% glutamine). Glutamine-rich domains have also been shown to function as transcriptional activation domains (Courey and Tjian 1988).

Full-length Pip does not bind AB independently of PU.1

To test the binding activity of the full-length Pip protein, we subcloned the complete ORF (from amino acid 2) and 218 bp of 3′-untranslated sequence into pBSATG, which provides a strong initiator codon for in vitro translation (Baldwin et al. 1990). Recombinant Pip protein was produced and analyzed by EMSA for its ability to bind to the kB element (Fig. 4A, B). Although Pip was initially cloned by virtue of its ability to bind kB, the full-length protein produced in vitro (Fig. 4B) does not display detectable binding activity in the EMSA (Fig. 4A). However, if recombinant PU.1 is supplied along with Pip, a complex migrating slower than the PU.1/kB complex appears in the gel. This complex, which has an electrophoretic mobility identical to that of JB2, appears only when PU.1 and Pip are present in the reaction together and is dependent on the presence of both the PU.1-binding site and the Pip-binding site, as mutation of either (Bm1 and Bm2, see Fig. 1C) abolishes its formation. In addition, the formation of the Pip/PU.1 complex is dependent on the presence of a serine residue at amino acid position 148 of the PU.1 protein, as mutation of this serine to alanine leads to a protein that can no longer recruit Pip to the kB site, even though its own DNA-binding ability is unaltered (Fig. 4A, S>A). This serine has been shown to be phosphorylated both in reticulocyte lysate-produced PU.1 and in PU.1 produced in vivo and, in its phosphorylated form, is required for the essential interaction between PU.1 and NF-EM5, which recruits NF-EM5 to its binding site on the K3′ enhancer DNA (Pongubala et al. 1993). Therefore, phosphorylation of Ser-148 in PU.1 is also required for interaction with Pip.

The Pip protein binds specifically to the kB element, but only after interacting directly with a PU.1 protein whose intact binding site is nearby. This interaction requires serine at position 148 of the PU.1 protein and results in the formation of a ternary complex with a mobility identical to that of the previously described complex JB2. It is likely that Pip is the second component of the JB2 complex. It is the correct size and displays all of the unique DNA-binding characteristics that have been described for that factor. In addition, because the NF-EM5 protein displays the same set of characteristics and
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Figure 4. Binding characteristics of the Pip and ICSBP proteins. (A) In a gel mobility shift assay, the full-length Pip protein binds to the B probe in a PU.1-dependent fashion. (B, Bm1, and Bm2) Probes used in the assay. (J) J558L nuclear extract, [PU.1, S>A, and Pip] recombinant proteins. (B) SDS-PAGE analysis of [35S]methionine-labeled recombinant proteins. Protein size markers are indicated. (C) ICSBP interacts with PU.1 to form a ternary complex on the B probe. A gel mobility shift assay using the B probe and recombinant ICSBP and PU.1 proteins is shown. (IBP) Recombinant ICSBP protein. The mobility of the PU.1 and JB2 complexes is indicated.

participates with PU.1 to form a λB complex with a mobility identical to that of JB2, it is likely that Pip and NF-EM5 are the same protein.

ICSBP binds λB in conjunction with PU.1

Because ICSBP is the protein related most closely to Pip with a nearly identical DNA-binding domain, we tested its ability to bind the λB element in the gel shift assay. ICSBP protein produced in the reticulocyte lysate system was unable to bind λB independently but was able to interact with PU.1 to form a complex that has a mobility identical to that of the Pip/PU.1 and JB2 complexes. (The size of the ICSBP protein produced in vitro is indistinguishable by SDS-PAGE from that of the Pip protein, C.F. Eisenbeis, unpubl.). The formation of the ICSBP/PU.1 complex is also dependent on the presence of serine at position 148 of the PU.1 protein [Fig. 4C]. Thus, ICSBP and Pip are identical in terms of their size and DNA-binding characteristics. Experiments with anti-ICSBP antibodies, however, indicate that ICSBP is not a major component of the JB2 complex [C.F. Eisenbeis, unpubl.]. This suggests that ICSBP is not NF-EM5.

The expression of Pip mRNA is tissue restricted

To determine the pattern of Pip gene expression, we performed Northern blot analysis and ribonuclease protection assays [RPAs] on a variety of RNAs from murine tissues and cultured cell lines. Northern blot analysis of polyA+ RNA from cultured murine cell lines [Fig. 5A] reveals that a 5-kb message is present in all of the B-cell lines examined (J558L, WEHI 231, and lipopolysaccharide [LPS]-stimulated 70Z/3), but not in an erythroid line [myeloid erythroleukemia (MEL)]. An additional, abundant message of 2.5 kb and a less abundant message of 1.5 kb are present only in J558L, a myeloma line that expresses λ1 light chains. The significance of these additional transcripts is unclear at the present time. Clearly, the complete Pip message is much longer (5 kb) than the largest cDNA clone that we have obtained (1.8 kb). Because numerous cDNA clones have a common 5' end, the Pip message appears to contain a long 3' UTR.

Analysis of total RNA from cultured murine cell lines reveals a similar pattern of expression (Fig. 5B): A 5-kb message is present in all B-cell lines analyzed, whereas additional 2.5- and 1.5-kb messages are present in the myeloma lines only (J558L, S194, and MOPC315). No Pip message was detectable by Northern blotting in EL4 (a T-cell lymphoma) or NIH-3T3 (a fibroblast) cell lines. To achieve greater sensitivity, further analysis of Pip gene expression was performed using an RPA. The Pip riboprobe used in this assay is 400 nucleotides in length, 350 of which will be protected by specific hybridization to Pip mRNA. A mouse β-actin probe was employed to control for RNA loading. Analysis of tissue RNAs shows that the Pip gene is expressed in murine spleen, thymus, and bone marrow but not liver, kidney, brain, or testes [Fig. 5C]. Analysis of cultured cell line RNAs confirms that while expression of Pip RNA is greatest in the three myelomas, there is expression of the Pip gene in all B-cell types tested. The Pip message is also present, al-
Figure 5. Analysis of the expression of Pip mRNA. A Northern blot analysis of 5 μg of polyA⁺ RNA [A] and 10 μg of total RNA [B] from cultured cell lines is shown. Blots were probed with a 679-bp NcoI fragment corresponding to the carboxy-terminal half of the Pip-coding sequence. Blots were rehybridized with a GAPDH probe to control for differences in sample loading. RNA size markers are indicated. An RPA of 5 μg of total RNA from tissues [C] and cultured cell lines [D] is shown. A β-actin riboprobe was included in the RPA to control for RNA quantity. The time of exposure of each set of samples is indicated.

beit at a much lower level than in any of the B-cell lines, in two T-cell lines, EL4 and PGL2 (Fig. 5D). There is no detectable expression of Pip RNA in MEL or NIH-3T3 (Fig 5D). Preliminary evidence also suggests that the Pip gene is not expressed in macrophage lines (A. Brass, unpubl.).

The Pip gene appears to be expressed exclusively in lymphoid cells. In addition, it appears to be expressed at a higher level in B cells than in T cells. The comparable levels of expression in the spleen and thymus could either be attributable to high levels of Pip RNA in thymocytes or, more likely, contaminating B cells in the tissue preparation (the thymus RNA sample had large amounts of κ RNA). Because PU.1 is restricted in its expression to the B-cell and macrophage lineages, the PU.1/Pip complex will only form in B cells, the only cell type in which the expression of both genes appears to overlap. Thus, this complex may play a critical role in determining the B-cell specificity of the λ and κ3' enhancers.

In conjunction with PU.1, Pip functions as a transcriptional activator

We have reported here that Pip and PU.1 interact to form a complex on the λB element of the Ex2.4 enhancer. We had shown previously that when multimerized, this element was sufficient to drive transcription of a reporter gene transiently transfected into the terminally differentiated B-cell line, J558L [Eisenbeis et al. 1993]. To determine whether the complex formed by Pip and PU.1 has transcriptional activating potential, we cotransfected expression plasmids containing PU.1 and Pip cDNAs into NIH-3T3 cells (which express neither protein) along with a chloramphenicol acetyltransferase [CAT] reporter construct containing the thymidine kinase [TK] promoter and a tetramer array of λB sites (Fig. 6). The λB sites function as cell type-specific activation elements, as they promote transcription in J558L B cells but not in NIH-3T3 fibroblasts. Whereas expression of Pip or PU.1 alone had no effect on CAT expression, the simultaneous expression of Pip and PU.1 in 3T3 cells was sufficient to drive transcription of the reporter. Coexpression of Pip with the S148A mutant of PU.1 did not lead to an appreciable increase in CAT expression, suggesting that the direct interaction between these two proteins is important in vivo as well as in vitro. These results suggest that Pip acts as a transcriptional activator in conjunction with PU.1, which is required for Pip to bind to the λB element. Interestingly, PU.1, which has been shown by others to be a transcriptional activator [Klemsz et al. 1990], cannot activate transcription alone through the λB tetramer.

Discussion

We have identified Pip, a novel transcription factor, which interacts with the Ets protein, PU.1, to form an activating complex on the λB motif of the Ex2.4 enhancer. Pip does not bind λB autonomously but relies on DNA-bound PU.1 to facilitate its binding to this DNA element. Pip is restricted in its expression to B and T cells; because PU.1 is not expressed in the T lineage, the Pip/PU.1 complex is B cell specific and may thus be an important determinant of the B cell specificity of the mu-
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The IRF-1, IRF-2, ICSBP, and ISGF3γ proteins are all transcription factors implicated in mediating transcriptional responses of cells to interferon stimulation (for review, see Williams 1991; Pellegrini and Schindler 1993). ISGF3γ is one of the subunits of the multimeric transcription factor ISGF3 (Darnell et al. 1994) whose nuclear localization is regulated by interferon signaling. Whereas ISGF3 and IRF-1 function as activators of transcription (Fujita et al. 1989; Kessler et al. 1990), ICSBP and IRF-2 function as transcriptional repressors (Harada et al. 1989; Weisz et al. 1992, Nelson et al. 1993).

DNA binding by Pip

The Pip protein is able to bind specifically to sequences essential for the activities of the murine E2A-4 (this paper) and Eκ3′ (C.F. Eisenbeis and A. Brass, unpubl.) enhancers. In each case, Pip binding requires that PU.1 be bound at its adjacent binding site. We have shown that the Pip DNA-binding domain, itself, can bind to these sites independently, suggesting the presence of an inhibitory domain within Pip that prevents DNA binding by the full-length protein. We postulate that interaction with PU.1 alters the conformation of Pip to unmask its DNA-binding domain and allow it to interact with the kB sequence. The interaction between Pip and PU.1 is reminiscent of a similar interaction between the Ets family members Elk-1 and SAP-1, which interact with serum response factor (SRF) to form a ternary complex on the bipartite binding site of the serum response element (SRE) of the c-fos promoter (Hipskind et al. 1991; Dalton and Treisman 1992).

Interaction between Pip and PU.1

The binding of Pip to the kB element is dependent on direct interaction between the Pip and PU.1 proteins and requires the presence of a serine residue at position 148 of the PU.1 protein. Phosphorylation of S148 occurs both in vivo and during translation of PU.1 in the reticulocyte lysate system and is required for the interaction between PU.1 and NF-EM5 (Pongubala et al. 1993). The fact that this phosphorylation is also required for the PU.1/Pip interaction further supports the contention that Pip is NF-EM5. This phosphorylation event could represent a regulated step in the control of immunoglobulin κ and λ light-chain gene expression. Interestingly, kB/PU.1 can also form a trimolecular complex with ICSBP, which also requires phosphorylation of S148 of PU.1.

The carboxy-terminal region of the Pip protein, which appears to be required for interaction with PU.1 (C.F. Eisenbeis, unpubl.) is highly conserved in the ICSBP protein (Fig. 3B), representing a possible shared interaction domain. Recently, it has been discovered that the activity of ICSBP may also be regulated by association with related IRF family members IRF-1 and IRF-2 to form complexes that bind strongly to the α, β interferon (IFN-α,β)-stimulated response element (ISRE) (Bovolenta et al. 1994). It will be interesting to determine whether Pip can also interact with IRF-1 and IRF-2.

Pip is a novel member of the IRF family, particularly related to ICSBP

The amino-terminal DNA-binding domain of the Pip protein displays a strikingly high degree of homology to the DNA-binding domains of the IRF family, including IRF-1, IRF-2, ICSBP, and ISGF3γ (Fig. 3A). The Pip protein is 80% identical to ICSBP across this region (amino acids 24–130 of Pip). In addition, Pip and ICSBP share many contiguous segments of homology in their carboxy-terminal halves (see Fig. 3B). The two proteins are 48% homologous over a 160-amino-acid region of the Pip protein (Pip amino acids 254–413). Thus, Pip represents a new member of the IRF gene family that is related most closely to ICSBP.

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Figure 6. Functional analysis of the Pip protein. Ectopic expression of Pip and PU.1 in cultured NIH-3T3 fibroblasts with a CAT reporter gene whose expression is under the control of a tetramer of kB elements. PU.1, Pip, and PU.1 S148A cDNAs are cloned into a eukaryotic expression vector containing the CMV promoter and enhancer. The reporter is the B4TKCAT construct (Eisenbeis et al. 1993). The indicated constructs were transiently cotransfected into NIH-3T3 cells. After 46 hr, cells were harvested and extracts were analyzed for CAT activity. The results are reported as an average of at least three independent transfections. The activity of the reporter alone has been set at 1.0.

rime immunoglobulin λ, as well as κ3′, enhancers. The Pip/PU.1 complex is identical in every respect to the NF-EM5/PU.1 and JB2 complexes described previously.
Transcriptional activation by PU.1/Pip

PU.1 has been shown to be a transcriptional activator [Klemsz et al. 1990] which possesses a transcriptional activation domain in the amino-terminal 75 amino acids [Hagemeier et al. 1993]. However, when expressed alone in NIH-3T3 cells, PU.1 is incapable of stimulating the expression of a cotransfected CAT reporter whose transcription is driven by a tetramer of kB elements [Fig. 6]. Perhaps the affinity of PU.1 for the kB site is lower than that for other sites, and an array of four sites is insufficient to achieve an activating potential. In this scenario, the Pip-dependent activation of the kB element by PU.1 could be attributable to cooperative binding [Fig. 4; Eisenbeis et al. 1993]. In addition, Pip may contain a transcriptional activation domain that enhances the activation potential of the complex.

The Pip protein contains a glutamine-rich domain at its carboxyl terminus and a proline-rich domain in the center of the protein sequence. Glutamine- and proline-rich sequences have been shown to function as transcriptional activation domains [Courey and Tjian 1988; Mermod et al. 1989]. In the Pip protein an 87-amino-acid region (151–237) contains 20% proline residues, and a 66-amino-acid region (354–419) contains 15% glutamine residues. Within these domains, the proline and glutamine residues are clustered, so that the P-rich domain consists of three P-rich modules (151–162, 33% P; 172–200, 24% P; 225–237, 38% P), and the Q-rich domain consists of two Q-rich modules (354–369, 25% Q, 395–419, 20% Q). Recent reports suggest that some transcriptional activation domains have a modular structure composed of 10- to 20-amino-acid subdomains with little transactivational activity, which cooperate to form a highly functional activation domain when present in combination [Seipel et al. 1992; Sutherland et al. 1992]. In addition, analysis of Oct-2 and the Fos:Jun heterodimer has demonstrated that different types of activation domains may exist within a single transcription factor and that these domains cooperate to provide a greater activation potential [Sutherland et al. 1992; Tanaka et al. 1994].

The role of IFNs in immunoglobulin gene expression

The relationship of Pip to ICSBP and the IRF family has interesting implications for the regulation of immunoglobulin light-chain gene expression. The IRF family members are regulated by extracellular actions of IFNs and the consequently activated signaling pathways. Whereas IFNs have been shown to play important roles in the regulation of the murine immune system [Sen and Lengyel 1992], it is not known at the present time whether the expression of Pip, like ICSBP, is regulated by IFNs.

It is intriguing that the kb and k3' composite elements are highly related to the ISRE. The ISRE consensus sequence is A/GNGAAANNGAAACT [Darnell et al. 1994]. This is identical to the kb site, except at the highlighted positions [Fig. 1C]. The kb and k3' elements may therefore function as ISREs, and expression of κ- and λ-genes may be positively regulated by IFN-α,β signaling.

IFN-γ, on the other hand, stimulates the expression of ICSBP, which is most closely related to Pip. ICSBP antagonizes activation by IRF-1 and ISGF3. Because ICSBP can also interact with PU.1 it may be a negative regulator of the λ and k3' enhancers.

The expression of immunoglobulin light-chain genes is regulated at several levels. In late pre-B cells, the κ and λ light-chain loci become accessible to the action of the V(D)J recombinase [Doglio et al. 1994]. Germ-line transcripts of κ and λ genes can be detected at the time of rearrangement and are presumed to precede the joining events. It is not known, however, how the activation of these loci is controlled. The proportion of rearranged κ genes greatly exceeds that of λ genes, and this discrepancy may be attributable to differential activation of the two loci. κ genes are under the control of the κ-intron enhancer [Eκi] and the k3' enhancer [Eκ3'], and only the latter is dependent on PU.1–NF–EM5/Pip. At the time of κ-gene rearrangement, only the intron enhancer appears to be active [Chen et al. 1994; Klug et al. 1994]. The λ locus, on the other hand, is regulated only by enhancers dependent on the PU.1–NF–EM5/Pip complex. Thus, it is conceivable that at the early stage of B-cell development when the Eκi is active, and Eκ3' is inactive, the λ enhancers are also inactive. This would allow preferential accessibility of the κ locus to the V(D)J recombinase, whereas λ remains inaccessible. It will be interesting to determine the roles of Pip/ICSBP and their control by interferons in the differential expression of κ and λ genes.

Materials and methods

λgt11 library construction and screening

RNA was isolated from J558L, a murine myeloma cell line, by the guanidinium isothiocyanate method. Poly(A)+ RNA was generated by two rounds of chromatography on an oligo(dT)–cellulose column [Pharmacia]. Using this poly(A)+ RNA, a custom cDNA library was made in the λgt1 vector by Stratagene. cDNA was synthesized by priming the reverse transcription reaction with both oligo(dT) and random primers. Primary plaques (3.9×10⁶) were obtained, and ~1×10⁶ of these were used to amplify the library. By use of a tetramer of λB sites, ~1.2×10⁶ plaques were screened from the amplified library [Singh et al. 1988]. The probe was isolated as a restriction fragment from a plasmid into which four directly repeated copies of the λB site had been cloned and labeled by filling in the overhanging ends with Klenow fragment in the presence of 100 μCi each of [³²P]dATP and [³²P]dCTP. Putative positive clones were purified through at least five rounds of plaque purification. Specific binding affinity for the λB probe was determined by probing each purified phage stock with a tetramer of λA sites in a similar in situ-binding assay. High titer stocks of recombinant phage were generated by the plate lysis method [Sambrook et al. 1989].

Generation of lysogen extract

E. coli lysogens harboring recombinant λgt11 phage were isolated as described [Singh 1991]. After inducing for expression of
β-gal fusion proteins, lysogenized cells were used to prepare soluble extracts for DNA-binding assays.

In vitro transcription and translation

Plasmids containing cDNA inserts were linearized 3' of the coding sequence. Capped mRNA was synthesized by in vitro transcription of 1 µg of linearized plasmid in the presence of 10 mM 7mG(5')ppp(5')G (New England Biolabs) and T3 or T7 RNA polymerase (Promega). After treatment with RNase-free DNase I for 1 hr. In some cases, parallel reactions were prepared in which the proteins were labeled by inclusion of [35S]methionine in the reaction mix. When proteins were labeled with [35S] they were analyzed by SDS-PAGE and autoradiography.

EMSAs and methylation interference

EMSAs were performed as described [Singh et al. 1986; Eisenbeis et al. 1993]. Probes were generated by digestion of plasmid DNAs containing the sequence of interest and labeled by filling in the ends with Klenow in the presence of [32P]dCTP. Methylation interference analysis was performed as described [Baldwin 1988; Eisenbeis et al. 1993]. Lysogen extract from the λ13 cDNA clone was incubated with methylated λB site probe and electrophoresed on a 4% polyacrylamide gel to isolate free and complexed probe. Chemical cleavage products were electrophoresed on 5% polyacrylamide/7 M urea gels. Autoradiography was performed on dried gels.

Subcloning and sequencing

The cDNA inserts from the λgt11 clones were subcloned as Ncol–Ndel fragments into the EcoRV site of pBluescript II (SK+) to generate λ3N/N, λ15N/N, λ37N/N, λ37.5N/N, and λ37.9N/N. In the case of λ37.9, which contains two Ncol sites within its coding sequence, the full-length insert was isolated from a partial Ncol–Ndel digest. In addition, the Pip coding sequence from a partial Ncol–Ndel digest was subcloned as a Mlu–Bcll fragment into the EcoRV site of pBluescript II (SK+) to generate 9(M/B). The entire λ37.9N/N cDNA insert was sequenced completely on both strands by generation of a set of nested exonuclease III deletions from each end of the insert [Slatko et al. 1991], followed by sequencing with T3 and T7 primers, dideoxynucleotide chain terminators, and Sequenase enzyme. The Pip ORF was subcloned into the pBSATG vector to generate 9ATG by engineering a HindIII site just upstream of the second amino acid of the Pip protein by polymerase chain reaction (PCR) using the 9ATG primer (5'-GGGGAAGCTTAACTTGGAGACGG-GCGA-3') in conjunction with the T7 primer and 9(M/B) as the template DNA. The Pip sequence was then isolated as a HindIII fragment, blunted by filling in with Klenow, and subcloned into Smal-cut pBSATG [Baldwin et al. 1990]. This construction places the Pip gene in the context of a strong translation initiation site and produces a protein in vitro that has the initial methionine and 8 amino acids provided by the vector and artificial restriction site, linked to the Pip ORF beginning with the second amino acid.

Northern analysis

Mouse tissue and cultured cell RNA was prepared by the guanidinium thiocyanate method. Poly[A]+ RNAs were fractionated on an oligo(dT)-cellulose column. For Northern analysis, 10 µg of tissue RNA, 5 µg of poly[A]+ RNA, or 10 µg of cultured cell RNA was electrophoresed on a 1% agarose/formaldehyde gel and blotted onto Hybond N nylon membrane [Amersham]. Blots were hybridized for ~16 hr at 40°C in 50% formamide, 5× Denhardt's solution, 5× SSC, 0.1% SDS, 10 µg/ml of sonicated salmon sperm DNA, and ~106 cpm/ml probe. After washing at high stringency the blots were subjected to autoradiography.

RPA

RPAs were performed using the RPA II kit [Ambion]. The Pip antisense riboprobe was generated by digestion of the 9(M/B) Pip plasmid clone with Nhel and transcription of the linearized plasmid in vitro using T7 RNA polymerase and 50 µCi of [32P]CTP. A β-actin control probe was synthesized in a similar manner using T3 RNA polymerase and the control plasmid DNA provided with the kit. Riboprobes were gel purified on 5% polyacrylamide/7 M urea gels before use. Hybridization reactions contained 10 µg of tissue RNA or 5 µg of cultured cell RNA and 5× 106 cpm of each riboprobe, and were incubated for 18 hr at 45°C before digestion with RNase. Protected fragments were analyzed by electrophoresis on 5% polyacrylamide/7 M urea gels, followed by autoradiography with an intensifying screen at ~70°C.

Transient transfection and CAT assay

The CAT reporter construct, B7TKCAT has been described previously [Eisenbeis et al. 1993]. To create Pip/CMV, PU.1/CMV, and S148A/CMV, the Pip, PU.1, and S148A cDNAs were subcloned as HindIII–Xbal fragments into the HindIII and Xbal sites in the polylinker of pRC/CMV [Invitrogen], which places the cDNA inserts under the control of the CMV promoter. NIH-3T3 cells were transfected by the DEAE-dextran protocol [Selden 1987] using a total of 25 µg of DNA. Transfection involved 5 µg of reporter construct and 10 µg of the expression constructs, with the balance made up with empty expression vector. After 46 hr at 37°C, cell lysates were prepared and CAT assays performed using 160 µl of lysate as described previously [Eisenbeis et al. 1993]. TLC plates were analyzed on a Molecular Dynamics PhosphorImager; the results shown here represent an average of at least three independent transfections.

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