The Oct-2 protein binds cooperatively to adjacent octamer sites

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Recombinant proteins derived from the cloned human oct-2 gene were used to investigate cooperative binding by Oct-2 to adjacent DNA-binding sites. Oct-2, a B-cell-specific transcription factor, binds tightly to the octamer sequence in immunoglobulin promoters. A second apparently unrelated consensus sequence in heavy chain promoters, the heptamer site, also is recognized by the Oct-2 protein but with 1000-fold lower affinity. Simultaneous occupancy of both the octamer and heptamer sites is favored by cooperative interactions. The heptamer site is probably recognized by the same binding surface in the Oct-2 protein as the octamer site and thus is conserved as a lower-affinity binding site. This permits the immunoglobulin promoter to respond to a much broader range of levels of Oct-2 protein. Substitution of prototype octamer sequences for heptamer sequences yields a probe with two octamer sites spaced by 2 nucleotides, which also binds Oct-2 protein cooperatively. Only the POU domain in the Oct-2 protein is required for this cooperative interaction. Similar protein–protein interactions between bound Oct-2 proteins may promote promoter–enhancer synergism in the heavy chain gene.

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Cell-type-specific transcriptional control of gene expression generally is accomplished by the action of sequence-specific DNA-binding proteins that interact with promoters and enhancers [Maniatis et al. 1987; Jones et al. 1988]. The Oct-2 protein, a sequence-specific DNA-binding protein that binds with high affinity to the sequence 5'-AGCAAAT-3' [called the octamer site], activates transcription from immunoglobulin promoters both in vivo and in vitro [Landolfi et al. 1986; Staudt et al. 1986; Scheidereit et al. 1987; Wang et al. 1987; LeBowitz et al. 1988; Müller et al. 1988]. Although the Oct-2 protein is thought to be expressed only in lymphoid cells and mediate the cell-type-specific effects of the octamer site, a protein with identical DNA-binding specificity is constitutively synthesized in all cell types [Pruijn et al. 1986; Singh et al. 1986; Fletcher et al. 1987; Sturm et al. 1987, O'Neill and Kelly 1988]. This protein, Oct-1, probably mediates activation of a number of constitutively expressed promoters that contain octamer sites. For example, transcription of genes encoding histones and small nuclear RNAs (snRNAs) depends on sequences recognized by Oct-1 [Majtaj et al. 1985; Mangin et al. 1986; Sive and Roeder 1986; Sive et al. 1986; Ares et al. 1987; Bark et al. 1987; Bohmann et al. 1987; Carbon et al. 1987; Krol et al. 1987; Murphy et al. 1987, Roebuck et al. 1987; Das et al. 1988].

The octamer site is present in the immunoglobulin heavy-chain promoter, in the opposite orientation in the κ light-chain promoter and in the heavy-chain enhancer, which is located in a downstream intron [Banerji et al. 1983; Gillies et al. 1983; Falkner and Zachau 1984; Parslow et al. 1984; Grosschedl and Baltimore 1985; Mason et al. 1985]. Each of these octamer sites is an independent cis-activator of transcription in lymphoid cells, presumably due to interactions with the Oct-2 protein [Bergmann et al. 1984; Falkner and Zachau 1984; Foster et al. 1985; Gopal et al. 1985; Picard and Schaffner 1985; Mizushima-Sugano and Roeder 1986; Dreyfus et al. 1987, Lenardo et al. 1987, Wirth et al. 1987]. The octamer site is adjacent to the promoter of the heavy chain enhancer [Dreyfus et al. 1987, Wirth et al. 1987], whereas multiple sites are recognized by a variety of proteins that activate or suppress transcription in different cell types in the enhancer [Church et al. 1985; Ephrussi et al. 1985; Augereau and Chambon 1986; Schlokat et al. 1986; Sen and Baltimore 1986; Wasylyk and Wasylyk 1986; Weinberger et al. 1986, 1988, Gerster et al. 1987; Lenardo et al. 1987; Peterson and Calame 1987]. There may be a synergistic interaction between the heavy chain promoter and its cognate enhancer. Exchange of either promoter or enhancer in this set with other elements that function in B cells reduces the level of transcription significantly [Garcia et al. 1986].

In the immunoglobulin heavy-chain promoter there is an additional conserved sequence, referred to as the heptamer site 5'-CTCATGA-3' [Ballard and Bothwell 1986; Eaton and Calame 1987; Landolfi et al. 1988]. The position of the heptamer site varies in different heavy chain promoters, ranging from 2 to 22 nucleotides upstream of the heptamer site is probably recognized by the same binding surface in the Oct-2 protein as the octamer site and thus is conserved as a lower-affinity binding site. This permits the immunoglobulin promoter to respond to a much broader range of levels of Oct-2 protein. Substitution of prototype octamer sequences for heptamer sequences yields a probe with two octamer sites spaced by 2 nucleotides, which also binds Oct-2 protein cooperatively. Only the POU domain in the Oct-2 protein is required for this cooperative interaction. Similar protein–protein interactions between bound Oct-2 proteins may promote promoter–enhancer synergism in the heavy chain gene.
the octamer site. The heptamer site has been shown to contribute two- to fivefold to the transcriptional activity of the heavy chain promoter in vivo and in vitro [Eaton and Calame 1987; Poellinger et al. 1989]. Although the heptamer site is not closely related in sequence to the prototype octamer site, recent evidence suggests that this site may be recognized by the Oct-2 and Oct-1 proteins. Octamer-binding proteins purified from extracts of B cells or HeLa cells [OTF-2 = Oct-2, OTF-1 = Oct-1] specifically bind both octamer and heptamer sequences [Kemler et al. 1989; Poellinger and Roeder 1988; Poellinger et al. 1989]. Interestingly, occupancy of the heptamer site by these two protein preparations did not occur if the adjacent octamer site was altered by mutation. This suggested a cooperative interaction between proteins bound to the adjacent two types of sites.

Recently, the genes encoding the Oct-1 and Oct-2 proteins have been cloned [Clerc et al. 1988; Müller et al. 1988; Scheidereit et al. 1988; Staudt et al. 1988; Sturm et al. 1988]. Although they are distinct genes, comparison of the derived amino acid sequences of these two proteins has revealed a region of remarkable structural similarity. The DNA-binding domains of both Oct-1 and Oct-2 are related to the homeo box consensus and are identical at 55 of 62 amino acid residues. An adjacent region amino-terminal to the homeo box is also highly conserved between Oct-1 and Oct-2, with 74 of 75 amino acid residues being identical. This second homology has been extended to at least two other genes: the rat pit-1 gene, a rat pituitary-specific transcription factor, and unc-86, a Caenorhabditis elegans gene implicated in determining cell lineage and differentiation. In each of these proteins, this second homology resides adjacent to a well-conserved homeo box. The second conserved sequence, together with the adjacent homeo box, has been dubbed the POU domain [Herr et al. 1988]. The POU domain is said to contain a POU box and a POU homeo box. The POU homeo box is believed to be responsible for the specific DNA binding of these proteins [Clerc et al. 1988; Ko et al. 1988; Garcia-Blanco et al. 1989]. The POU box appears to stabilize high-affinity DNA binding [Sturm and Herr 1988]; however, it is clearly dispensable for specific DNA binding, because one of the original Oct-2 cDNA clones isolated on the basis of specific binding to the octamer site contained less than half of the POU box [Staudt et al. 1988; Ko et al. 1988].

We used recombinant proteins derived from the Oct-2 cDNA to examine quantitatively the cooperative interaction between Oct-2 molecules bound to heptamer and octamer sites and to two octamer sites. The results indicate that Oct-2 binds cooperatively to both sets of sites and that the only protein determinant required for the cooperative interaction between adjacent Oct-2 molecules is the highly conserved POU domain.

**Results**

Previous studies have shown that Oct-2 protein sequences containing the POU box bind with high specificity and affinity to the octamer site in immunoglobulin promoters. Large amounts of a sequence-specific binding protein containing the POU box were isolated from bacteria expressing a Staphylococcus aureus protein A—Oct-2 fusion gene. The protein A sequence was joined to a partial Oct-2 cDNA sequence [plasmid 3-1; Clerc et al. 1988] at the EcoRI site, immediately upstream of the POU box sequences. Approximately 40 mg of the fusion protein [PAOct-2] was purified from bacteria by sequential chromatography on IgG-Sepharose and DNA-cellulose [see Materials and methods]. The purified PAOct-2 protein resolved into two major bands with molecular weights of 57,000 and 51,000, respectively, upon analysis by electrophoresis in SDS-polyacrylamide gels. The larger polypeptide corresponds to the predicted molecular weight of the fusion protein. The faster migrating species probably represents a proteolytic fragment. Recovery and renaturation of protein from either mobility species yielded a protein that specifically recognized the octamer sequence with high affinity.

Two different sources of proteins related to Oct-2 were used during the course of this study. The primary source was the purified PAOct-2 protein, which is free of other mammalian cellular proteins. Thus, the binding characteristics of this protein represent the activities of the cDNA-encoded sequences. All previous studies of binding by Oct-2 protein were performed with protein purified from B-cell extracts or reticulocyte lysates primed with RNA transcribed in vitro. These preparations could have been contaminated by other mammalian factors. The other source of Oct-2 protein was material translated in vitro in reticulocyte lysates. RNAs encoding portions of the Oct-2 protein were transcribed from the cDNA segment and added to the translation reaction. The latter source of protein permitted convenient manipulation of the Oct-2 sequences but yielded a product contaminated by reticulocyte material.

**PAOct-2 binds cooperatively to adjacent heptamer and octamer sites**

It has been suggested that the Oct-2 protein binds to the heptamer site [5'-CTCATGA-3'] when positioned adjacent to an octamer site [Kemler et al. 1989; Poellinger and Roeder 1989]. Such a potential cooperative interaction between bound proteins can be demonstrated in a gel shift assay by comparison of the DNA–protein complexes formed with a series of mutant and wild-type probes. The oligonucleotide probes used in these studies were derived from the immunoglobulin heavy-chain promoter V<sub>17,2,25</sub> [Grosschedl and Baltimore 1985]. Probe H*O* contained the wild-type promoter sequence [positions −69 to −47] in which the heptamer site is separated by 2 nucleotides from the octamer site [Table 1]. In probe H*O*, the heptamer site was specifically mutated by incorporating 5 nucleotide changes, whereas the octamer site and flanking sequences remained intact. Probe H*ΔO contains the wild-type heptamer sequence and a deletion of the octamer site. Comparison of the
concentrations of PAOct-2 protein required to bind to the heptamer site in probes H\(^+\)O\(^+\) and H\(^+\)AO permitted estimation of the cooperative interactions between bound Oct-2 proteins.

Titration of added PAOct-2 protein revealed three sets of complexes with the H\(^+\)O\(^+\) probe [Fig. 1], B1 complex, which forms at low protein concentrations, primarily reflects binding of PAOct-2 to the octamer site. This assignment has been confirmed by methylation interference and copper/phenanthroline footprint analysis (Figs. 2B and 5A). Complex B2 had a slower mobility and only formed at higher protein concentrations. Appearance of B2 required an intact heptamer sequence (Fig. 1A, cf. lanes designated 500, H\(^+\)O\(^+\) and H\(^+\)O\(^+\)). The slower mobility of this complex, together with the observation that its formation depended on the presence of the heptamer sequence, suggested that B2 was the result of simultaneous binding of PAOct-2 proteins to the octamer and heptamer sites. That complex B2 contained two protein molecules bound to the DNA was confirmed by detection of a heterodimer-type complex with two different sized Oct-2-related proteins produced by translation in vitro were added to the reaction simultaneously (data not shown). Also, methylation interference analysis of the B2 complex generated by PAOct-2 confirmed the occupancy of both the octamer site and heptamer sites [Fig. 2B]. The B3 complex only formed at high protein concentrations and required that the probe contain both an intact heptamer and octamer site. The specific sequences bound in this complex have not been determined, and it is unclear whether they represent the binding of three or more proteins.

A titration of the binding of the PAOct-2 protein to the H\(^+\)AO probe was generated also [Fig. 1]. A complex with a mobility similar to that of B1 was only detected with this probe at high protein concentrations [Fig. 1A]. This complex was shown to be due to specific interaction of the protein with the heptamer site by methylation interference analysis [Fig. 2A] and because the complex could be specifically competed by fragments that contained either the heptamer or octamer sites (data not shown).

Is the binding of PAOct-2 to the adjacent octamer and heptamer sites cooperative? Cooperativity can be assessed by comparing the protein concentrations required to generate equivalent occupancy of the heptamer site in the H\(^+\)O\(^+\) and H\(^+\)AO probes. Because the affinity of PAOct-2 for the heptamer site is much less than that for the octamer site [see below], fractional occupancy of the heptamer site in probe H\(^+\)O\(^+\) can be approximated by levels of the doubly liganded H\(^+\)O\(^+\) probe [complex B2]. The fraction of probe H\(^+\)O\(^+\) in the doubly liganded complex B2 was compared with amounts of H\(^+\)AO complex in Figure 1B. About 50- to 100-fold higher levels of protein were required to generate equivalent levels of occupancy of the heptamer site in the latter probe, indicating a 50- to 100-fold increase in the affinity for binding to the heptamer site when the adjacent octamer site is occupied. As discussed later, the association constant for PAOct-2 binding to the octamer site is \(\sim 6 \times 10^9/M\). Because equivalent levels of binding to the heptamer site alone require a 10-fold higher protein concentration [Fig. 1B], the association constant for this site can be estimated to be on the order of \(10^9-10^{10}/M\).

Equivalent levels of binding of the protein to the heptamer site when adjacent to an occupied octamer site can be detected at a protein concentration \(\sim 10\)-fold higher than that necessary to bind to the octamer site. Thus, cooperative interactions between the PAOct-2 proteins bound to the heptamer and octamer sites must account for the 50- to 100-fold higher affinity. This is likely an overestimate of the cooperativity, because the low affinity of the protein for binding to the single heptamer site may result in an underestimate of the level of this complex by the gel shift assay.

The results shown in Figure 1 were not analyzed in a more quantitative fashion because even at the highest protein concentrations, a significant fraction of the H\(^+\)O\(^+\) probe was recovered in the B1 complex. This suggests that there is a limitation in the binding assay that prevents quantitation of complex B2, the complex with both the heptamer and octamer sites occupied. This limitation may reflect dissociation of the protein from the heptamer site during the gel assay. Alternatively, it may reflect dissociation of the protein from the heptamer site in solution because we have also been unable to generate a complete protection of the heptamer site in a DNase I footprinting experiment. The same limitation may also be present in the detection of the protein—heptamer complex with the H\(^+\)AO probe. Although the quantitation of the results may be qualified, the binding is cooperative. This conclusion is strengthened by the coopera-

### Table 1. DNA probes used in the DNA-binding assays

| Probe designation | Orientation | Sequence* |
|------------------|-------------|-----------|
| H\(^+\)O\(^+\) | →⁻⁻⁻ | AATTCTGCCTCATGAGTATGCAAATCATG |
| H\(^+\)AO | →⁻⁻⁻ | AATTCTGCCCTCATGAGT |
| H\(^+\)O | →⁻⁻⁻ | AATTCTGAGGTGAGTATGCAAATCATG |
| 2xOA | ⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻‥ |

* See Materials and methods.
Methylation interference analysis of PAOct-2 binding to heptamer site

Methylation interference data for the PAOct-2-heptamer complex and complexes B1 and B2 are shown in Figure 2. The pattern of interference over the heptamer site is almost symmetric for the heptamer complex. The heptamer sequence, together with the adjacent guanine residue from the immunoglobulin promoter, constitutes an 8-nucleotide sequence with dyad symmetry: CTCTATGAG. This symmetry is probably fortuitous, as it is not a conserved feature of the heptamer sequence in immunoglobulin promoters. The two guanine residues on each strand of this sequence as well as one additional guanine residue, interfere with binding when methylated (Fig. 2A). Interaction of PAOct-2 with the octamer site is not symmetric with five sites on one strand and with one site on the other strand showing strong interferences (Fig. 2B). The affinity of the PAOct-2 protein for
Oct-2 cooperative binding

Figure 2. [A] Methylation interference analysis for binding of PAOct-2 to the H+O probe. Methylation interference analysis was carried out as described in Materials and methods. Lanes from a DNA sequencing gel are shown. [Left] The four lanes are derived from probe labeled at the HindIII end; [right] the four lanes are derived from probe labeled at the EcoRI end. [Lanes A + G] Reference ladders of cleavages at A and G residues; [lanes F] the cleavage pattern of free DNA; [lanes B] the cleavage pattern of DNA isolated from the bound complex. The relevant sequences are aligned with the A + G ladders. Arrowheads indicate the position of interfering residues denoted on the adjacent sequence by • (strong interferences) or O (weak interferences). 

[B] Methylation interference analysis for binding of PAOct-2 to the H+O* probe. Methylation interference analysis was carried out as described above. [Left] The six lanes are derived from probe labeled at the EcoRI end; and [right] the five lanes are derived from probe labeled at the HindIII end. [Lanes B1] The cleavage pattern of DNA isolated from the singly liganded complex, [lanes B2] the cleavage pattern of DNA isolated from the doubly liganded complex. The relevant sequences are aligned with the A + G ladders. Arrowheads indicate the position of interfering residues in the B2 complex denoted on the adjacent sequence by • (strong interferences) or O (weak interferences). 

[C] Summary of methylation interference contacts for probe H+O* complexes B1 and B2 and for probe H+O complex B1.

these two sites differs by 1000-fold. Given this weak binding to the heptamer site, the interaction of PAOct-2 with the heptamer sequence may be more sensitive to modifications that only slightly reduce the binding affinity. The complex B2, representing simultaneous occupancy of the octamer and heptamer sites, has the same methylation interference pattern observed for complex B1 but with one additional site on each strand where methylation of guanine interferes with binding (Fig. 2B). However, when the methylation interference contacts for the heptamer site complex, octamer site complex...
The loss of detection of interference at these sites becomes weaker in the doubly liganded complex or may reflect an alternate conformation of binding by the protein in the doubly liganded complex or may reflect an increased stability of these protein-DNA complexes due to cooperative interaction of the adjacent protein molecules.

The Oct-2 POU domain is sufficient for cooperative binding to adjacent heptamer and octamer sites

Having established that the PAOct-2 protein bound cooperatively to the heptamer and octamer sites, it was important to rule out the possibility that protein A sequences in the fusion protein mediated the cooperative interaction. This was accomplished by assaying truncated forms of the Oct-2 protein synthesized in vitro from RNAs transcribed from the Oct-2 cDNA (Fig. 3). The Oct-2 protein, designated P-H-C, extends from the POU box through the POU homeo box to the carboxyl terminus of the protein. This protein shares the same segment of the oct-2-coding sequences with PAOct-2. When assayed by mobility shift assay, P-H-C (which contained residues spanning the POU box, the POU homeo box, and through the carboxyl terminus) produced B1 and B2 complexes with the H+O+ probe. Formation of complex B2 depended on the presence of the intact heptamer site because it was not detected with the H-+O- probe. Furthermore, no complex was detected with the H+ΔO probe, indicating that at the protein concentrations assayed, P-H-C was capable of occupying the heptamer site only when the adjacent octamer site was occupied. Therefore, Oct-2 sequences contained within P-H-C specify cooperative binding to the heptamer site.

To determine whether Oct-2 protein sequences downstream of the POU homeo domain were required for the cooperative interaction, an additional protein synthesized in vitro was assayed. P-H, a protein of 157 amino acid residues, contains only the POU box and POU homeo box of the Oct-2 protein. When assayed in the mobility shift assay with the three probes described above, this protein behaved identically to P-H-C: Formation of complex B1 and B2 was observed with the H+O+ probe, formation of only complex B1 was observed with the H-+O- probe, and no complex was detected with the H+ΔO probe. Therefore, the POU box and POU homeo box are sufficient to allow cooperative binding of Oct-2 to adjacent octamer and heptamer sites.

Binding of PAOct-2 to adjacent octamer sites

Is the heptamer site simply a low-affinity octamer site? As it was difficult to align the heptamer sequence with the octamer sequence, we wondered whether the heptamer site was contacted by a distinct DNA-binding domain of the Oct-2 protein. Cooperative binding by the Oct-2 protein could be dependent on the presence of this unique site adjacent to an octamer site. Alternatively, the cooperative interactions could reflect an inherent property of the Oct-2 protein; thus, substitution of a second octamer site for the heptamer site should also yield cooperative binding. Four probes were used to test the possibility that Oct-2 could bind to two adjacent octamer sites cooperatively (Fig. 5). Probes 2xOA, 2xOB, and 2xOD (Table 1) contained the three possible orientations of two octamer sites separated by 2 nucleotides [as are heptamer and octamer in H+O+]. Probe 2xOC contained two octamer sites in inverted orientation with a 2-nucleotide overlap. This probe was designed to test the possibility that the consensus heptamer site represented an overlapping 5/8 match with the octamer site (cf. probes H+O+ and 2xOC in Table 1). Titrations of PAOct-2 with the four probes indicated that probes 2xOA and 2xOD formed doubly liganded complexes at relatively low protein concentrations (Fig. 4). Probes 2xOB and 2xOC proved refractory to formation of doubly liganded complexes (Fig. 4), except at very high protein concentrations. Identical results were obtained with the in vitro-translated Oct-2 P-H-C protein [data not shown]. Confirmation that the complexes labeled B2 were, in fact, doubly liganded complexes was provided by copper/phenanthroline footprinting, which revealed that both octamer sites in probes 2xOA (Fig. 5A) and 2xOD [data not shown] were occupied. Additionally, an experiment where two different molecular-weight
Oct-2-related proteins were mixed and confirmed that two-protein molecules were present in the B2 complexes [data not shown]. Finally, the B2 complexes formed with probes 2xOB and 2xOC are likely to represent specific doubly liganded complexes, because an equivalent amount of protein did not yield a second complex when assayed on probe H^-O^+ [see Fig. 1A].

Examination of DNase I and copper/phenanthroline footprints of the PAOct-2 octamer site complex reveals an asymmetric protection that extends 3' further of the octamer sequence than 5' [Fig. 5A]. This asymmetry predicts that when two octamer sites are oriented in a convergent pattern (2xOB probe), occupation of both sites necessitates considerable overlap of the footprints [Fig. 5B]. On the other hand, when the sites are oriented in a divergent pattern (2xOA probe), occupation of both sites requires a minimal overlap of the footprints [Fig. 5B]. The parallel orientation of probe 2xOD falls somewhere between these two extremes [some overlap of the footprints would be expected]. The difficulty in generating a doubly liganded complex on probe 2xOB is consistent with a steric hindrance, due to an extensive overlap of the binding sites.

Quantitative analysis of cooperative binding to adjacent octamer sites

Probes 2xOA and 2xOD were readily capable of forming doubly liganded complexes. Quantitative analysis of the binding of PAOct-2 protein to these two probes was performed on titration data over the entire range of the transition. This analysis makes the assumption that the gel shift assay accurately reports the fraction of molecules in the unbound, singly bound, and doubly bound forms. The titration curves for the interaction of PAOct-2 and probe 2xOA are shown in Figure 6A. The data were analyzed subject to the constraint that the microscopic binding constants describing the interaction of protein with either site in the probe are identical \( k_1 = k_2 \) [see Materials and methods]. Copper/phenanthroline footprint analysis of singly liganded DNA complex B1 showed no apparent bias toward occupancy of either site in probe 2xOA, in agreement with the above constraint [Fig. 5A]. This constraint allows the determination of \( k_1 \) and \( k_{12} \), the cooperativity constant, from the binding titration of probes with two sites [M. Brenowitz, unpubl. numerical analysis]. The agreement of the curves predicted by the best-fit binding constants [Table 2] with the data points over the entire range of the titration suggests that this model is adequate to describe the results. This quantitative analysis indicates that binding of PAOct-2 to the two adjacent octamer sites in 2xOA occurs with ~10-fold cooperativity [Table 2].

Binding of PAOct-2 to a probe containing a single octamer site [probe H^-O^+] was analyzed quantitatively also [Fig. 6B]. The value of \( k_1 \) determined from this experiment was identical, within experimental error, to that determined from probe 2xOA [Table 2]. As another, but related, test of the validity of the model of cooperative interactions of protein binding to two identical octamer sites, values for \( k_1 \) and \( k_{12} \) can be determined by simultaneous analysis of total data from both the single-site probe experiment and the 2xOA probe experiment [Brenowitz et al. 1986; Senear et al. 1986]. The results of
this analysis are completely consistent with the analysis of each probe alone (Table 2) and support the conclusion that Oct-2 binding to two adjacent octamer sites show 10-fold cooperativity. A similar analysis was performed with the data from titrations with probe 2xOD (data not shown). In this case, the two octamer sites are aligned in parallel with a spacing of 2 nucleotides. The results in Figure 4 show that both sites in probe 2xOD can be occupied by the protein at reasonably low concentrations. Quantitative analysis of the data using a pool of two titrations yields a value of 1.3 ± 0.6 for $k_{12}$, the cooperativity parameter (Table 2, see Materials and methods). This result suggests that the two proteins bind with little or no cooperativity to two sites in parallel alignment spaced by 2 nucleotides.

Titrations of the binding of the P-H form of the Oct-2 protein to the probe 2xOA with divergent octamer sites were also analyzed. The absolute concentration of P-H protein was not known in this titration because the binding material was synthesized by translation in vitro. Thus, relative values of the concentration of P-H were used in analysis of the titration curves, and a value for $k_{12}$ was determined (Fig. 6C). The value of $k_{12}$ obtained from this analysis, 4.3 ± 1.5, is lower than that reported above for PAOct-2. However, given that P-H clearly binds cooperatively to the H+O+ probe, this result suggests that the P-H protein also binds cooperatively to...
two octamer sites when they are in a divergent orientation.

Discussion

The major transcriptional determinant in the immunoglobulin promoter is the octamer site that is recognized by the B-cell-specific protein Oct-2 (Dreyfus et al. 1987; Wirth et al. 1987). Promoters of immunoglobulin heavy chain genes also have a conserved heptamer sequence, spaced either 2, 14, or 22 bp upstream of the octamer site (Ballard and Bothwell 1986; Eaton and Calame 1987). Mutation of this conserved sequence reduced three- to fivefold the level of transcription from the promoter in B cells (Eaton and Calame 1987). We demonstrated that a protein synthesized in *Escherichia coli* from a plasmid containing part of the Oct-2 cDNA binds specifically to the heptamer site, even though this sequence is not closely related to the octamer sequence. Furthermore, the Oct-2-related protein interacts cooperatively when bound to both the octamer and heptamer sites and thus increases the occupancy of the heptamer site by ~50 to 100-fold. The presence of the heptamer site greatly expands the concentration range over which the heavy chain promoter is sensitive to levels of Oct-2. At low Oct-2 concentrations, the octamer site alone would be partially occupied. As this site becomes occupied with increasing Oct-2 concentration, a second molecule of Oct-2 would bind to the heptamer site. The binding of this second molecule would account for the observed three- to fivefold dependence of transcription on the

| Probe   | $k_1/M$          | $k_{12}$ |
|---------|------------------|----------|
| H-O+    | $5.8 \pm 1.0 \times 10^8$ | —        |
| 2xOA    | $5.3 \pm 1.1 \times 10^8$ | $10.7 \pm 3.4$ |
| Simultaneous* | $5.4 \pm 1.0 \times 10^8$ | $10.2 \pm 3.5$ |
| 2xOD    | $18.1 \pm 5.0 \times 10^8$ | $1.3 \pm 0.6$ |

*Simultaneous analysis of the two probes H-O+ and 2xOA.
heptamer sequence. This increase in transcription, due to binding to the heptamer site, could potentially be greater in some cells, as it is not clear that the B-cell tumor lines that were tested for heptamer activity express adequate concentrations of Oct-2 to saturate the heptamer site [Eaton and Calame 1987].

Oct-2-derived proteins containing only the POU domains are fully active in binding to the octamer site and in cooperatively binding to the heptamer site. This 162-amino-acid sequence contains two evolutionary conserved regions, the POU box and POU homeo box [Clerc et al. 1988; Herr et al. 1988; Sturm et al. 1988]. Because the sequence of the heptamer site does not closely resemble the octamer site, it was possible that the POU box specified two distinct DNA sequence recognition surfaces. We do not believe this is the case but rather propose that the same protein surface is responsible for recognition of both the octamer and heptamer sites. A DNA probe containing two prototype octamer sites, oriented in a divergent pattern and spaced by 2 nucleotides, also binds the Oct-2-related protein in a cooperative fashion. The degree of cooperativity is 10-fold. These results suggest that the heptamer sequence in the heavy chain gene is conserved as an intermediate affinity Oct-2-binding site so that the promoter responds to a broader range of levels of the B-cell-specific protein.

It is interesting to picture two Oct-2 proteins bound to two octamer sites, spaced by 2 bp oriented in a divergent pattern. In this configuration, the copper/phenanthroline footprints of the two proteins abut precisely, and the proteins should be aligned on the same side of the duplex helix and related by an approximate twofold center of rotation. Binding to the two octamer sites in this divergent configuration was clearly cooperative. Cooperative interactions between Oct-2 proteins bound to octamer sites with spacings of 14 and 22 bp would also be expected because the heptamer and octamer sites are separated by these distances in some immunoglobulin promoters. Interestingly, the relative orientations of the heptamer and octamer sites are conserved in all heavy-chain promoters examined, suggesting that this orientation may be essential for cooperative binding. Two octamer sites oriented in a parallel fashion and spaced by 2 bp can also bind Oct-2 proteins simultaneously. In this case, the binding is apparently noncooperative. This is somewhat surprising, as the copper/phenanthroline footprints should overlap by 1 bp in this orientation. It is possible that a potential steric hindrance due to this overlap is compensated by positive cooperative interactions. If this were the case, parallel binding sites with other spacings may display cooperative binding. However, this would also infer that two Oct-2 proteins could cooperatively interact in two different relative orientations. Competitive binding for the two octamer sites was clearly observed when two octamer sites were convergently oriented and spaced by 2 nucleotides. This was not unexpected, as the asymmetric footprints would have to overlap extensively when two proteins bind to this probe simultaneously.

Cooperative binding to two octamer sites also only requires protein sequences within the POU domain. The ability to specifically recognize another protein bound to adjacent sequences may reflect the function of the POU box within the POU domain. The other region within the POU domain, the homeo box, is clearly involved in DNA binding and must make direct contacts with the major groove of the binding site. The POU box, on the other hand, is not essential for specific DNA binding, because one of the original Oct-2 cDNA clones, isolated on the basis of binding to the octamer site, contained less than half of the POU box. However, the POU box is important for DNA binding, because deletions or mutations of this region significantly lower the affinity of Oct-2 for the octamer site. There are several indications that proteins possessing the POU domain interact cooperatively with other proteins. First, it is highly likely that Oct-1, as well as Oct-2, binds cooperatively to the heptamer–octamer pair of sites [Poellinger and Roeder 1989]. Second, the Oct-1 protein is thought to interact with the herpes simplex virus virion protein aTIF [ICP25/VP16/VMW65] and another cellular protein in cooperatively binding and regulating the immediate early promoters of the virus [Gerster and Roeder 1988; T.M. Kristie et al., in prep.]. These types of cooperative interactions have not been described for previously characterized proteins containing homeo box domains and thus may reflect the contribution of the POU box.

Perhaps the best-characterized system displaying cooperative interactions is the binding of bacteriophage λcl repressor protein to the operators O 1, Y 2, and Y 3 of the A P 1 promoter [Johnson et al. 1979]. This system is highly cooperative with values for the cooperativity constants reported ~26 [Akers et al. 1982]. Interestingly, although the physiologically relevant configuration of cl-binding sites is one where the sites are adjacent, when two operator sites are separated by six helical turns, i.e., 60 bp, cooperative binding of cl repressor to the two sites is still observed [Hochschild and Ptashne 1986]. This cooperative interaction presumably occurs via a looping of the DNA.

Regulation of gene expression through the cooperative interaction of DNA-binding proteins also has been observed with mammalian proteins [Topol et al. 1985; Davidson et al. 1988; Tsai et al. 1989]. The ability of Oct-2 molecules to interact cooperatively when binding to octamer sites may account for the observed synergistic transcriptional activity of the immunoglobulin heavy-chain promoter and enhancer [Garcia et al. 1986]. Protein–protein interactions, via DNA looping, between Oct-2 molecules bound to the octamer site in the heavy-chain enhancer and the octamer and heptamer sites in the heavy-chain promoter would bring the enhancer more stably into close proximity with the promoter. This, in turn, might enhance the ability of factors bound to the enhancer to interact with general transcription factors on the promoter. Currently we are examining the possibility that Oct-2 can bind to octamer sites separated by large distances through a looping mechanism.
Materials and methods

Plasmids and constructions

Plasmid 3-1 contained the 1.1-kb cDNA fragment with EcoRI termini isolated from a λgt11 recombinant and ligated into the unique EcoRI site of pKS+ (Stratagene). The insert corresponded to nucleotide positions 655–1713 in the oct-2-coding sequence [Clerc et al. 1988].

Plasmid pRIT2T (Pharmacia) is designed to give temperature-regulated overexpression of recombinant proteins resulting from fusion of coding sequence of interest downstream of the amino-terminal portion of the S. aureus protein-A gene. Plasmid oct-2/pRIT2T was constructed by isolating the 1.1-kb EcoRI fragment from plasmid 3-1, filling in the EcoRI termini with the Klenow fragment of DNA polymerase I (Boehringer–Mannheim), and ligating the resultant blunt-end fragment into the Smal site of plasmid pRIT2T with T4 DNA ligase (Boehringer–Mannheim). The ligation mixture was transfected into E. coli strain X90, a lysogenic strain with cI+ phenotype (gift of R. Sauer). Recombinants with an in-frame fusion with protein A were isolated. One such isolate was designated oct-2/pRIT2T.

Plasmid pBS-ATG-oct-2 has been described previously [Clerc et al. 1988].

DNA probes

Complementary synthetic oligonucleotides with an EcoRI site compatible sequence on one strand and a BamHI compatible sequence on the other strand were annealed and ligated into pUC18 or pUC19, cut with EcoRI and BamHI. One exception to this is the oligonucleotide for probe 1202O, which is self-complementary with BamHI overhangs. This was cloned directly into the BamHI site of pUC19. The resultant series of plasmids were the source of DNA probes used in these studies and were excised by digestion with EcoRI and HindIII, except where noted. One strand of the oligonucleotide pairs and their designations are listed in Table 1.

Induction and purification of PAOct-2

The PAOct-2 fusion protein was purified from E. coli strain N4830-1 [F-, su-, his-, ilv-, galKΔ8, N+ (Boehringer–Mannheim), harboring the plasmid oct-2/pRIT2T. A 150-liter culture of cells was grown in L broth, containing 1 g/liter of cerelose and 100 μg/ml of ampicillin in a New Brunswick fermentation fermentor at 30°C. Cells were grown with the Klenow fragment of DNA polymerase I (Boehringer–Mannheim), and ligating the resultant blunt-end fragment into the Smal site of plasmid pRIT2T with T4 DNA ligase (Boehringer–Mannheim). The ligation mixture was transfected into E. coli strain X90, a lysogenic strain with cI+ phenotype (gift of R. Sauer). Recombinants with an in-frame fusion with protein A were isolated. One such isolate was designated oct-2/pRIT2T.

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DNA-binding assays

Reactions for gel mobility shift assays [Fried and Crothers 1981; Garner and Revzin 1981] were assembled at room temperature in a final volume of 15 μl and contained 50 mM Tris-HCl (pH 7.5), 60 mM NaCl, 4% Ficoll, 1 mM DTT, 150 μg/ml bovine serum albumin, and 5 × 10^{-10} M DNA-binding-site probe. Nonspecific competitor DNAs used in the assays were 26 μg/ml of poly(dIdC-dIdC) for assays of the purified PAOct-2 protein and a mixture of 25 μg/ml of sonicated denatured calf thymus DNA and 2.5 μg/ml of sonicated native calf thymus DNA for assays of in vitro synthesized proteins. Assays of in vitro synthesized protein also contained 1 mM EDTA. DNA probes were prepared as described [LeBowitz et al. 1988]. Complexes were resolved on 4% [40 : 1 acrylamide/bis-acrylamide] native acrylamide gels, using a Tris-glycine buffer: 25 mM Tris, 190 mM glycine, 1 mM EDTA (pH 8.3).

Methylation interference assays were performed essentially as described [Kristie and Roizman 1988], with minor modifications. DNA probes were end-labeled at either the EcoRI or HindIII site by filling in the 5’-overhang with the Klenow fragment of DNA polymerase I in the presence of [α-32P]dATP [New England Nuclear]. These probes were methylated at purine residues (less than one methylation event per DNA molecule) with dimethylsulfate [Aldrich], as described [Kristie and Roizman 1988]. DNA-binding reactions with methylated DNA probes were assembled, and nucleoprotein complexes were resolved as described above. Nucleoprotein complexes and free probe were isolated from the gel by electric transfer to NA45 paper [Schleicher and Schuell]. Species visualized by autoradiography of the NA45 paper were eluted by incubation in 10 mM Tris-HCl (pH 7.5), 1 M NaCl, and 1 mM EDTA at 65°C for 60 min. The eluted material was extracted sequentially with phenol/chloroform [1 : 1] and chloroform and then was ethanol-precipitated in the presence of 30 μg of carrier tRNA. Is-
lated DNAs were cleaved at methylated purines, as described, and resolved on 10% sequencing gels.

Copper/phenanthroline footprints were performed in situ by incubating mobility shift gels directly in the copper/phenanthroline cleavage reagents as described (Kuwabara and Sigman 1987). DNA-binding reactions with end-labeled probes were assembled and complexes resolved as described above. Cleaved nucleoprotein complexes were isolated by electric transfer to NA45 paper and purified as described above. Isolated DNAs were resolved on 10% sequencing gels. End-labeled probes were prepared as described above.

DNase I footprinting was performed essentially as described (Galas and Schmitz 1978). DNA-binding reactions with end-labeled probes were assembled as described above. DNase I (Worthington) (200 ng) was added plus sufficient magnesium chloride to achieve a final concentration of 1 mM. After a 20-sec incubation at room temperature, the reaction was terminated by the addition of 200 µl of stop solution (10 mM EDTA, 0.14 mg/ml tRNA, 2.67 M ammonium acetate). Reactions were extracted sequentially with phenol/chloroform [1:1], and chloroform and were then ethanol-precipitated. Footprint patterns were revealed by electrophoresis on 10% sequencing gels.

Quantitative analysis

The densitometric analysis of mobility-shift titration autoradiograms was conducted using the methods developed to analyze footprint titration autoradiograms [Brenowitz et al. 1986; Brenowitz and Senear 1989]. The model used to describe binding of the Oct-2 fusion protein to DNA containing two binding sites, Equations la-c, were fit to the titration data for all the bands in the lane. For the probes containing two binding sites, Equations la–c, were fit to the titration data for free, singly, and doubly liganded molecules, respectively. The details of this analytical procedure will be published elsewhere.

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