Sequence-specific DNA binding of the B-cell-specific coactivator OCA-B

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B-cell-specific transcription of immunoglobulin genes is mediated by the interaction of a POU domain containing transcription factor Oct-1 or Oct-2, with the B-cell-specific coactivator OCA-B (Bob-1, OBF-1) and a prototype octamer element. We find that OCA-B binds DNA directly in the major groove between the two subdomains of the POU domain, requiring both an A at the fifth position of the octamer element and contact with the POU domain. An amino-terminal fragment of OCA-B binds the octamer site in the absence of a POU domain with the same sequence specificity. Coactivator OCA-B may undergo a POU-dependent conformational change that exposes its amino terminus, allowing it to recognize specific DNA sequences in the major groove within the binding site for Oct-1 or Oct-2. The recognition of both the POU domain and the octamer sequence by OCA-B provides a mechanism for differential regulation of octamer sites containing genes by the ubiquitous factor Oct-1.

[Key Words: OCA-B; Oct-1; Oct-2; octamer sequence; cell-type-specific transcriptional regulation; DNA binding]

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Specific gene expression is mediated by the interaction of transcription factors with DNA elements present in promoters and enhancers. However, many transcription factors and their DNA elements have widespread distribution, which leads to a fundamental question: How do ubiquitous components mediate selective gene activation? The octamer motif (5'-ATGCAAAT-3') represents a DNA element with widespread distribution. First identified as a conserved motif present in immunoglobulin [Ig] heavy and light chain promoters and enhancers (Falkner and Zachau 1984), the octamer motif was subsequently found in the promoters of many cellular genes including H2B (Sive et al. 1986; Fletcher et al. 1987) and snRNAs (Carbon et al. 1987; Tanaka et al. 1988; Murphy et al. 1989; Yuan and Reddy 1989) (see Fig. 6, below). Although present in diverse genes, a single copy of the octamer element is sufficient to mediate the B-cell-specific expression of heterologous genes expressed from minimal promoter elements [Wirth et al. 1987].

In B cells, the octamer element is bound by Oct-1 and Oct-2 (Singh et al. 1986; Staudt et al. 1986), which share a conserved DNA-binding domain, the POU domain [Herr and Cleary 1995]. The POU domain is composed of two structurally independent DNA-binding domains, the POU-specific domain [POUs] and the homeodomain [POUH], connected by a flexible linker. Both domains bind the major groove but on opposite sides of the DNA [Klemm et al. 1994], with the POUs domain contacting the 5'-ATGC subsite and the POUH binding the AAAT-3' subsite. The amino-terminal arm of the POUH contacts the minor groove at the center of the octamer element. No protein–protein contacts are made between the two POU subdomains, but they bind cooperatively even in the absence of the linker. The cooperativity is believed to be mediated partially by overlapping contacts with phosphate groups at the center of the octamer element [Klemm and Pabo 1996].

Because Oct-1 is expressed in a ubiquitous manner and Oct-2 is expressed predominantly in B cells, the latter initially was thought to be responsible for the B-cell-specific function of the octamer element. Subsequent experiments, however, showed that Oct-1 and Oct-2 could equivalently activate octamer containing promoters in vitro [LeBowitz et al. 1988; Pierani et al. 1990; Luo et al. 1992] and that cell lines and animals lacking Oct-2 could still produce Ig [Corcoran et al. 1993; Feldhaus et al. 1993]. The B-cell-specific activity of the octamer element was then proposed to be mediated by a B-cell-specific coactivator that interacted with both Oct-1 and Oct-2 and modulated their function [Pierani et al. 1990; Pfisterer et al. 1994].

The recently discovered B-cell-specific coactivator OCA-B binds the POU domain of Oct-1 and Oct-2 both in solution [Luo and Roeder 1995; Pfisterer et al. 1995] and as a complex with the prototype octamer sequence [Gstaiger et al. 1995; Luo and Roeder 1995; Pfisterer et al. 1995; Strubin et al. 1995]. OCA-B is expressed exclusively in cells of the B lymphocyte lineage. In B cells,
OCA-B is present as two polypeptides of 34 kD and 35 kD, probably attributable to a posttranslational modification, and is rich in proline, serine, and threonine residues. In vitro, OCA-B stimulates Ig but not H2B promoters [Luo and Roeder 1995] in spite of identical sequences for the octamer element, implying a role for promoter context in the mechanism of OCA-B stimulation. In vivo, OCA-B in the presence of Oct-1 stimulates the H2B promoter two-fold whereas it stimulates an Igk promoter — seven-fold [Strubin et al. 1995] and an IgH promoter fifteen-fold [Gstaiger et al. 1995].

The cooperative assembly of multiprotein complexes involving distinct combinations of transcription factors and/or cofactors on adjacent sequence elements is a common theme in differential gene regulation. For example, induction of the immediate early genes of herpes simplex virus [HSV] is mediated by the formation of a complex, termed C1, composed of Oct-1, the viral regulatory protein VP16 (aTIF, Vmw65), and the cellular protein HCF [C1] [Kristie and Roizman 1984; Gerster and Roeder 1988; Kristie et al. 1989; Xiao and Capone 1990]. Formation of the C1 complex requires a specific DNA sequence (5'-ATGCAAAT-3') that differs from the prototypic octamer site in two ways. There is a T at position 5 and an additional 3'-flanking element, the GARAT motif (5'-GATATT-3'), immediately adjacent to the octamer site.

The precise assembly of this complex on the HSV immediate early promoters is characterized by multiple protein–protein and protein–DNA interactions. The GARAT element is recognized by VP16 but the affinity of this interaction is sufficiently low that assembly of the complex requires additional interactions with both Oct-1 and HCF [Kristie and Sharp 1990]. Four residues on the surface of the Oct-1 homeodomain are critical for formation of the C1 complex. Their importance is highlighted by the fact that the closely related Oct-2 POU homeo domain, which only differs by one of these and six additional residues, cannot form the C1 complex [Pomerantz et al. 1992]. The other essential cellular protein HCF [C1] probably does not contact DNA but interacts with both VP16 and Oct-1. HCF is a large complex protein that is post-translationally modified by proteolytic cleavage and by phosphorylation [Kristie et al. 1995; Wilson et al. 1995]. These modifications may provide further contributions to specific interactions within the C1 complex or affect the functioning of the C1 complex.

Thus, the octamer element is involved in the formation of at least two multicomponent, transcriptionally active complexes. Sequences within or adjacent to the octamer motif may contribute to the differential activity of the octamer element in different cell types. Although the recent discovery of OCA-B [Gstaiger et al. 1995; Luo and Roeder 1995; Pfisterer et al. 1995; Strubin et al. 1995] has indicated how a ubiquitous promoter element can be activated in a tissue-specific manner, further levels of control must exist as not all genes containing octamer sites are activated in B cells. We show that OCA-B is a unique type of coactivator that binds DNA in a sequence-specific fashion and interacts with the POU domain of Oct-1 or Oct-2. The ability of OCA-B to simultaneously contact protein and DNA provides a further level of control which may be a general mechanism of coactivator function.

Results

OCA-B binds only a subset of octamer elements on the basis of the identity of position 5 and not the sequence of the flanking bases

The B-cell-specific coactivator OCA-B binds the POU domain of Oct-1 and Oct-2 on the prototype octamer sequence [Gstaiger et al. 1995; Luo and Roeder 1995; Pfisterer et al. 1995; Strubin et al. 1995]. However, in a mobility shift analysis using a fragment of the HSV αIE element containing a nonprototype octamer element and additional flanking sequences (5'-A1TGCTAATGATA TT-3'), an Oct-1 POU domain/OCA-B/DNA complex could not be detected [Fig. 1A, lanes 1–4]. This αIE element has been extensively studied and supports the assembly of a multiprotein complex requiring interactions between Oct-1, the viral aTIF [VP16], and an additional cellular factor C1 [HCF] to form the C1 complex [Kristie et al. 1989]. To determine whether the nonprototype octamer element bearing a T at position 5 or the flanking sequences interfered with the formation of the complex, a probe [αIE-half] containing sequences known to be recognized by only Oct-1 (5'-A1TGCTAATGATA TT-3') was constructed. Although an Oct-1 POU complex formed with this sequence, it did not stably interact with OCA-B [Fig. 1A, lanes 9–12]. The αIE octamer sequence differs from the prototype at position 5, T in place of A. To assess the effect of the nonprototype T at position 5 on binding, the αIE-wt probe was mutated from T to A at position 5 (5'-ATGCAAATGATATT-3'). The αIE(T/A) probe efficiently formed the Oct-1 POU domain/DNA complex, and this complex bound OCA-B resulting in an additional ternary complex of slower mobility [Fig. 1A, lanes 5–8]. The same results were obtained using full length Oct-2, OCA-B bound to the Oct-2/DNA complex only on the αIE(T/A) probe [data not shown]. Thus, formation of a multiprotein/DNA complex involving an octamer site, the POU domain from Oct-1 or Oct-2, and OCA-B requires a specific DNA sequence at the center of the octamer element (5'-ATGCAAATGATATT-3'). These data suggest that having an A at position 5 as contrasted to T either confers a specific conformation on the POU domain that is competent for OCA-B binding or that OCA-B directly contacts DNA.

In transfection experiments, OCA-B has been shown to activate transcription from the Igk, IgH, and H2B promoters [Gstaiger et al. 1995; Strubin et al. 1995] [albeit with different strengths] that have prototype octamer sequences. Probes were prepared containing the prototype element as well as 5' and 3' flanking regions of both the IgH and H2B promoters. The Oct-1 POU domain efficiently bound OCA-B on both the IgH-wt and H2B-wt probes [Fig. 1B, lanes 1–4 and 9–12]. When position 5
The B-cell-specific coactivator OCA-B contacts DNA 

OCA-B can bind to an octamer element with separated subsites for the POU$_5$ and POU$_H$ subdomains

The importance of position 5 of the octamer element is apparent in the crystal structure of the Oct-1 POU domain–octamer complex as it is contacted by both the POU$_5$ and POU$_H$ domains [Klemm et al. 1994]. Each of the POU subdomains can independently bind a subsite of the octamer element flanking position 5. Although the two POU subdomains do not make protein–protein contacts with one another, they are connected by a flexible linker and bind DNA cooperatively [Klemm and Pabo 1996] [see Fig. 3]. The cooperativity is believed to be partially mediated by overlapping contacts with phosphate groups at position 5, the same position found to be important for the formation of the Oct-2 or Oct-1 POU/OCA-B/DNA complex.

Separation of the subsites by insertion of two base pairs in the middle of the prototype octamer sequence, in a way that retains the base immediately flanking each subsite, lowers the affinity of the POU domain by ~30-fold and abolishes the cooperativity of the binding of the subdomains [Klemm and Pabo 1996]. A ten-fold higher concentration of Oct-1 POU was used to generate a significant level of Oct-1 complexed with DNA on the H2B sequence with separated subsites [5'-ATGCAACAAT-3']. Under these conditions, OCA-B was able to bind and produce an Oct-1 POU/OCA-B/DNA complex [Fig. 4A, lanes 4 and 6]. This suggests that the contacts needed for OCA-B binding are present even when the POU subdomain requires a specific DNA sequence at the center of the octamer element [5'-ATGCAAAT-3'].

Figure 1. OCA-B distinguishes between octamer elements on the basis of the identity of position 5 and not the sequence of the flanking bases. (A) For EMSA, full-length OCA-B was added at 100 or 200 ng to DNA-binding reactions containing the probe listed at the top of each group of lanes in the presence of the POU domain of Oct-1 (137.5 pg/20 μl reaction). The probes used contained either the HSV alE wt sequence (5'-GTGCATGCTAATGATATTCTTTGG-3') [Kristie et al. 1989] (lanes 1–4), a T to A mutation at position 5 (alE-(T/A) = 5'-GTGCATGCAAATGATATTCTTTGG-3') (lanes 5–8), or a deletion at the 3'-end (alE-half = 5'-GTGCATGCTAATGAT-3') (lanes 9–12). The position of the various protein–DNA complexes are indicated by arrows. (B) Using the same protein concentrations as above, EMSA was performed on parallel DNA-binding reactions containing the probes listed at the top of each group of lanes. The probes contained the following sequences: Ig-wt = 5'-AATTCACCCTGTCTCATGAATATGCAAATCAGGTGAGTCTATG-3' Ig-(A/T) = 5'-AATTCACCCTGTCTCATGAATATGCTAATCAGGTGAGTCTATG-3' H2B-wt = 5'-GATCCTAGAATCGCTTATGCAAATAAGGTGAAGAGTTGAAGTCG-3' H2B-(A/T) = 5'-GATCCTAGAATCGCTTATGCTAATAAGGTGAAGAGTTGAAGTCG-3'. The sequence of these probes contains the octamer site and immediate flanking regions of the endogenous IgH [Landolfi et al. 1986] and H2B [Pierani et al. 1990] gene promoters. Within the octamer element of each of these probes was mutated from A to T, OCA-B did not bind the Oct-1/DNA complex [Fig. 1B, lanes 5–8 and 13–16]. As anticipated, the POU domain of Oct-1 bound both sites with similar affinities. The same results were obtained when Oct-2 was tested with these probes [Fig. 2, and data not shown for IgH]. Thus, formation of a multiprotein Oct-1 or Oct-2/OCA-B/DNA complex involving an octamer site requires a specific DNA sequence at the center of the octamer element [5'-ATGCAAAT-3'].

Figure 2. OCA-B forms an Oct-2/OCA-B/DNA complex on octamer sites containing an A but not a T at position 5. DNA–protein binding reactions were carried out as described in Materials and methods. The sequences of the probe DNAs used are listed in the legend to Figure 1B. Full-length Oct-2 was isolated from the human B-cell line, BJAB. Full-length OCA-B containing a six-histidine tag at the carboxyl terminus was used at 100 ng [lanes 3 and 7] and 200 ng [lanes 4 and 8] per reaction. The position of the Oct-2/DNA and Oct-2/OCA-B/DNA complexes are indicated with arrows. Free probe migrated to the bottom of the gel.
Figure 3. Structure of the Oct-1 POU domain–octamer complex illustrating the A–T base pair at position 5. The crystal structure of the Oct-1 POU domain–octamer DNA complex is shown (Klemm et al. 1994). The DNA is colored blue and the Van der Waals surface of the A–T base pair at position 5 is highlighted in purple. The POU<sub>5</sub> is shown as a red ribbon, the POU<sub>4</sub> as a yellow ribbon, and the linker connecting the carboxyl terminus of the POU<sub>4</sub> to the amino terminus of the POU<sub>H</sub> is shown in gray. The linker sequence was not ordered in the crystal structure but has been modeled here and is shown to emphasize that the two domains are linked. One strand of the octamer sequence is labeled. This figure was generated with INSIGHT II (Biosym Technologies). Adapted from Klemm and Pabo (1996).

The specificity for OCA-B binding A at position 5 and the structure of the POU domain/DNA complex (Fig. 3) led to the prediction that OCA-B contacts DNA in the major groove. To test this hypothesis, two octamer elements containing DNA probes from the H2B gene were synthesized with modified bases at position 5. It has been shown that substitution of the A–T base pair at position 5 with A–U results in a four-fold reduction in the binding of the POU domain caused by the loss of a contact between the methyl group in the major groove of DNA and the exocyclic amino group at position 4 of T and Leu55 in the POU<sub>H</sub> domain (Klemm and Pabo 1996). Analysis of the position 5 A–U substitution confirmed the reduced affinity of the POU domain; however, when the concentration of POU was increased so that a significant amount of POU domain/DNA complex was formed, addition of OCA-B at the same concentration as with the prototype element produced a supershifted complex (Fig. 4C). Thus, loss of the major groove methyl group at position 5 by replacing the T with a U did not inhibit OCA-B binding.

Additional modifications substituted the wild-type A–T base pair with an Inosine (I)–5 methyl (Me) C base pair. This substitution alters the topology of the major groove of DNA without changing the minor groove. In an A–T base pair, the major groove is characterized by a phosphate group at position 4 of A and thymine with the exocyclic amino group at position 6 of adenine, its substitution with a carboxyl group results in a three-fold lower than the wild-type H2B probe (data not shown). Addition of OCA-B to these reactions did not generate a slower mobility complex indicating that this sequence was a very low affinity binding site for OCA-B (Fig. 4A, lanes 7–12). Also, the amino-terminal 118 amino acid fragment of OCA-B did not bind the I–5MeC probe (Fig. 4A, lanes 8 and 11, and Fig. 4B, lanes 10–12). Thus, elimination of the exocyclic amino group at position 6 of adenine, its substitution with a carboxyl group at position 6 of inosine, or the exchange of the carboxyl group at position 4 of thymine with the exocyclic amino group movement of A–T base pair at position 5 with A–U results in a four-fold reduction in the binding of the POU domain caused by the loss of a contact between the methyl group in the major groove of DNA and the exocyclic amino group at position 4 of T and Leu55 in the POU<sub>H</sub> domain (Klemm and Pabo 1996). Analysis of the position 5 A–U substitution confirmed the reduced affinity of the POU domain; however, when the concentration of POU was increased so that a significant amount of POU domain/DNA complex was formed, addition of OCA-B at the same concentration as with the prototype element produced a supershifted complex (Fig. 4C). Thus, loss of the major groove methyl group at position 5 by replacing the T with a U did not inhibit OCA-B binding.

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An amino-terminal 118 amino acid fragment of OCA-B can bind DNA in the absence of the POU domain

The amino-terminal 118 amino acids of OCA-B are fully active in generation of the Oct-1 POU/OCA-B/DNA complex (Strubin et al. 1995). Surprisingly, in the absence of the Oct-1 POU domain, GST–OCA-B 118N bound DNA in a sequence specific manner that required an A at position 5, even when the subsites were separated (Fig. 4B, Fig. 4A, lanes 2 and 8). A titration of OCA-B 118N into DNA-binding reactions containing the H2B probe used in Figure 4B, in the presence or absence of the POU domain of Oct-1, revealed that the binary complex of OCA-B 118N and DNA formed at the same concentration of OCA-B 118N as the Oct-1 POU/OCA-B 118N/DNA complex (data not shown). This indicates that a major contribution to the affinity of OCA-B 118N in the trimolecular complex may result from its interaction with DNA. In addition, protein–protein interactions occur between the amino-terminal domain of OCA-B and the POU domain (Pfisterer et al. 1995; Gstaiger et al. 1996). These interactions may stabilize OCA-B binding in the trimolecular complex. This is supported by the difference in intensity of the OCA-B 118N/DNA complex (Fig. 4A, lane 2) compared to the stronger Oct-1 POU/OCA-B 118N/DNA complex (Fig. 4A, lane 5). Importantly, full-length OCA-B did not bind the octamer element in the absence of the POU domain (Fig. 4A, lanes 3 and 9; data not shown for Ig-wt and Ig-[A/T]). This suggested that the carboxyl terminus may shield the amino terminus from interaction with DNA.
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Figure 4. OCA-B binding requires base contacts in the major groove in the center of the octamer element. An amino-terminal 118 amino acid fragment of OCA-B can bind this site in the absence of the POU domain. [A] For EMSA, DNA-binding reactions were carried out in 20 µl and included the probes labeled at the top of each group of lanes as well as the Oct-1 POU domain (2.75 ng for H2B (+2), and 275 pg for 5 Me C-I), and full-length GST–OCA-B (250 ng), or GST–OCA-B 118N (500 ng) as indicated. The H2B probes used here are based on those constructed by Klemm et al. (1994) and have different flanking sequences and lengths than those used in Figure 1. The 84 base long sequence of one strand of the H2B-wt probe is 5'-AATTCCTGGATCAAGATCTGGTCACCCCATGGGCTAGCGCAGGCCCAAGGCTGTATGCAAATAAGGACGCGTTCGCGAGGGCCCG-3', and the other strand is complementary. The sequence of the H2B (+2) probe is similar to the H2B-wt probe except that two base pairs have been inserted into the middle of the octamer site (5'-ATGCACAAAT-3'). The sequence of one strand of the H2B derived 5-methyl C-I probe is 5'-TTTTTGCGCCCATGGGCTAGCGCCTGCCCAAGGCTGTATGCAAATAAGGACGCGTTCGCGAGGGCCCG-3', and the other strand is 5'-GGATCCCCGCGGGCCCTCGCGAACGCGTCCTTATTUGCATACAGCCTTGGGCATGCGCTAGCCCATGGGC-3' such that the probe length is 47 base pairs. [B] For EMSA, DNA-binding reactions were carried out in 20 µl and included the probes labeled at the top of each group of lanes as well as GST–OCA-B 118N (400 or 800 ng). (*) Position of the free probe. Note that in data not shown, removal of GST from OCA-B 118N had no effect on its sequence-specific binding. The sequence of the 70 base long H2B A/T probe is 5'-AATTCCTGGGTCCATGGGCTATGCCCAAGGCTGTATGCTAATAAGGACGCGTTTCGCGAGGGCCCGGATCC-3'. (C) EMSA was performed as above (275 pg Oct-1 POU and 100 or 200 ng OCA-B) with a synthetic DNA probe containing an A–U base pair at position 5. The sequence of one strand of the probe is 5'-TTTTTGCGCCCATGGGCTAGCGCCTGCCCAAGGCTGTATGCAAATAAGGACGCGTTCGCGAGGGCCCG-3' and the sequence of the other strand is 5'-GGATCCCCGCGGGCCCTCGCGAACGCGTCCTTATTUGCATACAGCCTTGGGCATGCGCTAGCCCATGGGC-3' such that the probe length is 84 base pairs.

group at position 4 of 5-methyl cytosine dramatically decreased the binding of both full length OCA-B in an Oct-1 POU/OCA-B/DNA complex, and OCA-B 118N in an OCA-B 118N/DNA complex.

OCA-B selectively activates promoters containing an A vs. T at position 5 of the octamer prototype in vivo

To determine whether OCA-B could distinguish A- from T-containing sites in vivo, transient transfection experiments were performed with 293 cells [Fig. 5]. Exponentially growing cells were cotransfected with OCA-B and a reporter construct containing two copies of either the prototype octamer site or the position 5 A to T mutant upstream of an IgH core promoter followed by the luciferase gene and the SV40 enhancer. Under these conditions in vivo, endogenous Oct-1 protein should permit formation of the Oct-1/OCA-B/DNA complex. Cotransfection of the cells with OCA-B and the A containing reporter resulted in a 4.4-fold increase in luciferase activity over transfection of the reporter in the absence of OCA-B. Neither the T-containing reporter nor an octamer negative reporter were stimulated by cotransfection of OCA-B. These results are consistent with the gel shift analyses and suggest that whereas Oct-1 can bind to octamer sites containing either an A or a T at position 5 in vitro, OCA-B only binds to and activates transcription from Oct-1 bound to the A containing prototype octamer site.

Discussion

OCA-B is a novel type of coactivator in that it contacts DNA, yet its function as a transcriptional coactivator is
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Figure 5. OCA-B distinguishes the presence of A vs. T at position 5 of the octamer prototype in vivo. 293 cells were cotransfected with 5 μg of reporter vector, 0 (white), 3 (gray) or 6 μg (black) of OCA-B expression vector [pCATCH Bob-1 (Gstaiger et al. 1995)], 1 μg of pCMV β-gal used as an internal control, and additional empty expression vector, pRCMV [Invitrogen], to a total of 20 μg. Reporter vectors were unaltered or contained two tandem copies of either the octamer element (5′-ATGCAAAT-3′) or the A to T mutation at position 5 (5′-ATGCTAAT-3′) followed by the core promoter (49′-1) from the V17.2.25 IgH gene [Grosschedl and Baltimore 1985], the luciferase gene, and the SV40 enhancer. Fold activation was calculated by dividing the amount of luciferase activity obtained in the presence of OCA-B by the amount of activity obtained in the absence of OCA-B, normalized to β-galactosidase production, for each reporter construct. Error bars indicate one standard deviation from the mean when the fold activation in three experiments was averaged. Each measurement was performed in duplicate.

Two separate lines of evidence strongly imply that OCA-B binds to specific DNA sequences, and a third experiment supports this conclusion. First, an amino-terminal 118 amino acid fragment of OCA-B binds DNA in the absence of Oct-1. Like the interaction of full-length OCA-B with the Oct-1/DNA complex, this binding is dependent upon an A in the fifth position of the octamer element and more specifically on the atomic structure of the major groove at this site. Second, an alteration of the major groove at position 5 of the octamer element from an A-T base pair to an I-5MeC base pair does not alter contacts between the Oct-1 POU domain and DNA [Klemm et al. 1994]; however, it abolishes the formation of an Oct-1 POU/OCA-B/DNA trimolecular complex. Third, both full-length OCA-B and OCA-B I18N can bind to an Oct-1 POU/DNA complex formed on an octamer site containing a two-base-pair insertion at its center which separates the subsites for the POU<sub>δ</sub> and POU<sub>α</sub> domains. If OCA-B binds exclusively to the POU domain and not DNA, it is surprising that it is able to bind to the complex of the Oct-1 POU domain on an octamer element where the separation of subsites is expected to alter the relative orientation of the two POU subdomains. This configuration of the POU subdomains would not be expected to support a fixed interaction with OCA-B unless either OCA-B makes direct contacts with the DNA or OCA-B binding requires interaction with only one of the POU subdomains. In addition, OCA-B I18N binds to this site in the absence of the Oct-1 POU domain, reinforcing the idea that it makes contacts with the DNA.

We attempted to investigate the affinity of OCA-B I18N for an octamer site and for the Oct-1/POU/DNA complex. Direct measurement of the affinity of either complex was hampered by the inability to detect a high level of binding in the electrophoresis mobility shift assays, possibly attributable to dissociation of the complexes during electrophoresis. Nevertheless, the formation of a shifted OCA-B containing protein-DNA complex was saturable. If the saturated level in the gel reflects binding of all of the probe in the binding reaction before electrophoresis, and the amount of probe bound in solution for each amount of OCA-B is proportional to the amount detected in the gel assay, the affinity of OCA-B I18N for the octamer site can be estimated to be 500 nM. As the activity of the OCA-B protein diminished with time, our calculation may be an overestimate of the concentration of active protein in the binding reactions, and thus an underestimate of the affinity.

Qualitatively, we are able to make several observations concerning the relative affinities of OCA-B I18N for DNA alone and for the POU domain/DNA complex. It is clear that the full-length OCA-B protein has a much greater affinity for the POU domain/DNA complex than for the DNA alone. Binding of the full-length protein to an octamer site was undetectable. Similarly GST–
OCA-B 118N showed strong preferential binding to the POU domain–DNA complex over the octamer element alone. In contrast, binding of OCA-B 118N was less discriminating. Its preference for the POU domain DNA complex over DNA alone was revealed in gel electrophoresis mobility assays by the formation of equal or more complex over DNA alone was revealed in gel electrophoresis. Analysis of the Oct-1 POU domain and direct binding of OCA-B to both the POU domain and the DNA, it is not known if POU domain/DNA contacts are unchanged after binding of OCA-B. In fact, the amino terminus of OCA-B may interact with residues in the POU specific domain known to contact DNA [Gstaiger et al. 1996]. Thus, the contacts that the POU domain makes with DNA in a POU/DNA complex may be different from those made in a POU/OCA-B/DNA complex. A loss of DNA contacts by the POU-specific domain may be compensated by additional contacts to OCA-B such that the off rate of Oct-1 from DNA is not altered. The precise mode of OCA-B binding cannot yet be understood through relation to any known families of DNA-binding proteins by sequence homology. Thus, OCA-B might represent a novel motif for protein–DNA interactions.

The conservation of adenosine at position 5 of the octamer element and the B-cell-specific expression of the OCA-B coactivator represent complementary aspects of B-cell-specific gene activation. The octamer elements of immunoglobulin genes universally conserve the adenosine at position 5 [Fig. 6], allowing high levels of OCA-B dependent expression of these genes in B cells. Octamer elements with T at the 5th position are not to our knowledge present in immunoglobulin genes even though this sequence binds the Oct-1 and Oct-2 proteins with high affinity. A similar adenosine also is found in the octamer elements associated with ubiquitously expressed genes such as those encoding histone H2B and snRNAs. Expression of these genes in non-B cells and their correct temporal expression in both B and non-B cells may require additional coactivator proteins, as has been suggested [Luo and Roeder 1995]. In contrast, octamer elements with a T at position 5 are found in viral genes, snRNAs, and origins of DNA replication, allowing these elements to exploit the activities of Oct-1 or Oct-2 but evade the regulatory effects of OCA-B in B cells.

It is well established that a complex of Oct-1, HSV αTIF and a cellular factor (HCF) recognizes a subset of octamer sites dependent on specific flanking sequences to activate the transcription of HSV immediate early genes [McKnight et al. 1987; Gerster and Roeder 1988; Preston et al. 1988; Kristie et al. 1989; Kristie and Sharp 1990]. Through its interactions with both Oct-1 and the octamer site, OCA-B represents a cellular example of a similar strategy. The OCA-B protein also simultaneously recognizes DNA sequences and makes specific interactions with the POU domain, however, in this case the sequence requirements are particularly compact. OCA-B recognizes sequences between the two subdomains of the POU domain. The ability of a transcriptional coactivator to contact simultaneously specific DNA sequences and a transcription factor may provide a widespread mechanism by which highly related cis-regulatory sites can exist in many genes and still mediate tissue-specific gene activation.
Materials and methods

Production of fusion proteins

Full-length OCA-B was produced as a glutathione S-transferase (GST)-OCA-B-His fusion protein by adding a six-histidine tag to the carboxy-terminus of OCA-B with the polymerase chain reaction (PCR) and cloning in-frame into the pGEX-5X-3 vector [Pharmacia]. The fusion protein was expressed using standard methods in TOPP3 cells [Stratagene] and purified on glutathione-Sepharose 4B [Pharmacia]. Free OCA-B-His was cleaved from the glutathione beads with Factor Xa protease [Promega] and full-length OCA-B containing the His tag at the carboxy-terminus was purified on Ni²⁺ agarose [Qiagen]. The protein was eluted with 250 mM imidazole in 20 mM HEPES, 5 mM 2-Me, 100 mM KCl, 0.1% NP40, and 20% glycerol.

Recombinant Oct-1-POU was produced as a GST–fusion [Klemm et al. 1994]. Briefly, following affinity chromatography on glutathione–Sepharose and cleavage from the matrix, the Oct-1 POU domain was further purified on a DNA–cellulose column in 0.05 M Tris at pH 8.0, 0.15 M NaCl, 1 mM EDTA, and 5 mM dithiothreitol (DTT). The column was washed with 0.05 M HEPES at pH 7.5, 0.05 M NaCl, and 0.01 M DTT, and the protein was eluted in the same buffer with 0.25 M NaCl. The protein was then dialyzed against double-distilled water and concentrated by lyophilization.

A vector expressing GST–OCA-B 118N was constructed by inserting a PCR product encoding the first 118 amino acids of OCA-B into a modified pGEX-5X-3 vector [Pharmacia] such that sequences encoding the initiator methionine of OCA-B were juxtaposed directly with sequences encoding the four amino acids of the Factor Xa cleavage site. The fusion protein was expressed in BL21 cells grown in minimal media at room temperature, and purified on glutathione-Sepharose 4B [Pharmacia]. GST–OCA-B 118N was eluted with 10 mM reduced glutathione in 50 mM Tris at pH 8.0, 1 mM DTT, 1 mM EDTA, 100 mM KCl, and 10% glycerol.

Electrophoretic mobility shift assays

DNA–protein binding reactions contained 0.5–1.0 ng of DNA probe, 400 ng of poly [dI-C]/poly[dI-C], 50 ng salmon sperm DNA, 10 mM HEPES at pH 7.9, 0.5 mM EDTA, 56.7 mM KCl, 0.75 mM DTT, 4% Ficoll-400, 300 μg/ml BSA and the appropriate amount of purified proteins listed in the figure legends in a total volume of 20 μl. Reactions lacking the DNA probe were incubated for 15 min at room temperature prior to addition of the probe and subsequent incubation for an additional 20 min before the complexes were resolved on 4% (Oct-2), 6% (Oct-1 POU), or 11% (OCA-B 118N) nondenaturing polyacrylamide gels using Tris-glycine electrophoresis buffer.

Transient transfection

293 cells were transfected by calcium phosphate precipitation with a glycerol shock using standard procedures [Ausubel et al. 1994]. Reporter vectors were constructed by first cloning DNA fragments containing KpnI and XhoI restriction sites and two copies of the indicated octamer sequence separated by a PstI site into the pGL2 enhancer vector [Promega]. Next, the V172.25 IgH gene core promoter was inserted downstream of the octamer sequences by cloning into the XhoI and HindIII sites. Luciferase activity was measured at 48 hr with 40 μl of a 100 μl total extract per 10-cm plate using 150 μl Luciferase Assay Reagent [Promega] and a ML2250 Luminometer [Dynatech Laboratories, Chantilly, VA] with the enhanced flash program and integration for 20 s with no delay. β-gal production was measured by incubating 2 μl of cell extract with the CPRG substrate [Boehringer Mannheim] in NaPO₄ buffer in the presence of MgCl₂ and β-mercaptoethanol for 10 min at 37°C, stopping the reaction with 0.055 M Na₂CO₃, and determining the OD at 574 nm of the resulting solution.

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Note added in review

During review of this manuscript, Gstaiger et al. [1996] independently reported that formation of an Oct-1 or Oct-2 POU domain/OCA-B/DNA ternary complex occurs when an A but not a T is present at position 5 of the octamer element. In addition, they report that a fragment of OCA-B containing the amino-terminal 65 amino acids of OCA-B interacts with the POU domain via protein–protein interactions. They identify residues in the POU-specific domain close to the major groove at position 5 of the octamer site which, when mutated eliminate formation of the ternary complex in a gel shift assay. They suggest that loss of the ternary complex occurs because these mutations abolish protein–protein contacts between OCA-B and the POU domain, however, this need not be the case. It remains possible that the critical mutations in the POU-specific domain alter more than protein–protein interactions. In particular, conformational alterations may occur in the POU domain because of the mutational changes such that different DNA moieties are exposed in the major groove at the center of the octamer element. Such alterations could abolish formation of the ternary complex by obscuring moieties used by OCA-B to contact DNA.

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