Structure of an Ebf1:DNA complex reveals unusual DNA recognition and structural homology with Rel proteins

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Early B-cell factor 1 (Ebf1) is a key transcriptional determinant of B-lymphocyte differentiation whose DNA-binding domain has no sequence similarity to other transcription factor families. Here we report the crystal structure of an Ebf1 dimer bound to its palindromic recognition site. The DNA-binding domain adopts a pseudo-immunoglobulin-like fold with novel topology, but is structurally similar to the Rel homology domains of NFAT and NF-κB. Ebf1 contacts the DNA with two loop-based modules and a unique Zn coordination motif whereby each Ebf1 monomer interacts with both palindromic half-sites. This unusual mode of DNA recognition generates an extended contact area that may be crucial for the function of Ebf1 in chromatin.

Supplemental material is available at http://www.genesdev.org.

Received August 2, 2010; revised version accepted August 23, 2010.

Early B-cell factor 1 (Ebf1) is the best-studied member of the EBF family (also known as the CO/E family) of transcription factors, which is common to all metazoans [Liberg et al. 2002]. The EBF family includes four members [Ebf1–4] in mammals, one member [knot/collier] in Drosophila [Krzemien et al. 2007], and one member [unc3] in Caenorhabditis elegans [Kim et al. 2005]. EBF family proteins consist of an N-terminal DNA-binding domain (DBD) of ~225 amino acids, a transcription factor immunoglobulin (TIG/IPT) domain, and a helix–loop–helix (HLH) domain, followed by a C-terminal transactivation domain, which is predicted to be unstructured [Hagman et al. 1995]. They form stable dimers via the HLH domain [Hagman et al. 1995] and bind to the palindromic consensus motif 5′-TCCCCNGGGA-3′ [Travis et al. 1993; Treiber et al. 2010].

All EBF proteins characterized to date play important roles in developmental processes, including cell fate decisions, cell differentiation, and cell migration [Liberg et al. 2002]. Ebf1 is essential for the differentiation of B lymphocytes, as a loss of gene activity leads to a complete block at the pre-pro-B-cell stage [Lin and Grosschedl 1995]. Conversely, forced expression of Ebf1 in hematopoietic stem cells or multilineage progenitors leads to enhanced formation of B lymphocytes at the expense of other lineages, demonstrating the instructive capacity of Ebf1 [Zhang et al. 2003; Medina et al. 2004; Pongubala et al. 2008]. In addition, Ebf1 regulates the differentiation of adipocytes [Jimenez et al. 2007] and sensory neurons [Wang et al. 1997; Garel et al. 1999]. Finally, a role for EBF proteins as tumor suppressors is beginning to emerge [Liao 2009].

Results and Discussion

Structure of Ebf1 bound to DNA

To gain insight into the mechanism of DNA recognition by EBF proteins, we recombinantly expressed the DBD of murine Ebf1 (amino acids 26–240) and co-crystallized it with a DNA duplex containing the Ebf1 consensus binding site. The structure, determined at 2.4 Å, shows two DBD molecules bound as a symmetric dimer to the palindromic DNA duplex [Fig. 1A], and an additional Ebf1 monomer not bound to DNA (Supplemental Fig. S1). The DBD folds into a β sandwich, in which a four-stranded β sheet (consisting of strands A, B, G, and F) packs against a five-stranded sheet formed by strands C, D, E, H, and I. The N terminus forms an α helix (helix 1) and packs against the bottom of the β structure. The apical part of the domain is formed by three additional short β strands [X, Y, and Z] and extensive loops that constitute the DNA-binding surface. The unusual Zn knuckle, which does not fit into any described class of zinc fingers [Schwabe and Klug 1994], is formed by a loop and three short helices [helices 2, 3, and 4] that are stabilized by a centrally coordinated zinc ion [Supplemental Fig. S2]. The Zn knuckle protrudes from one side of the domain and also participates in DNA recognition. The two protein monomers contact each other at a small interface of 240 Å², formed by a loop and a 310 helix. The C-terminal loops of
Crystal structure of Ebf1 bound to DNA. (A) Structure model of a dimer of Ebf1 DBDs bound to DNA. β-Sandwich assembly are shaded in yellow and orange and are labeled A-I. Additional strands are colored green and are labeled X–Z. Helices are shown in blue and are numbered. Zinc ions are represented by pink spheres. (B) Structure model of dimeric Ebf1 26–423 bound to DNA. DBDs are shaded as in A, TIG domains are shown in light and dark green, and HLH domains are blue and purple. Unstructured loops and linkers are indicated as dashed lines.

The DBDs run through the major groove, leaving the domain opposite of the dimerization interface. A detailed assignment of secondary structure elements within the DBD is shown in Supplemental Figure S3. A second crystal form with only one Ebf1:DNA complex in the asymmetric unit yielded an identical model at 3.0 Å resolution [data not shown].

Purification and crystallization of a complex consisting of a longer Ebf1 protein fragment (amino acids 26–422) and a cognate oligonucleotide provided additional structural information about the TIG domain and the HLH dimerization domain at 2.8 Å resolution [Fig. 1B]. The structure models for the DNA-bound DBDs from the three crystal forms at different pHs are essentially identical, with a Cα RMSD [root-mean-square deviation] between 0.45 Å and 0.62 Å for the individual chains, corroborating the presented structure. The C termini of the DBDs are connected to the TIG domains by a linker, which is visible in only one of the monomers. Due to the length (~25 Å) and flexibility of the linker, the orientation of the TIG domains relative to the DBDs is probably determined by crystal packing. As predicted from sequence comparison, the TIG domains adopt an Ig-like fold. Although they form a considerable dimer interface (490 Å²), the major part of the contact area is contributed to by the HLH domains, which are necessary for dimerization in the absence of DNA [Hagman et al. 1995]. In the structure, the electron density for the HLH domain was only weakly defined. However, four helices that form a helix bundle reminiscent of dimerized HLH domains, such as those found in MyoD (Ma et al. 1994), were discernable in one of the two Ebf1 dimers of the asymmetric unit. In all vertebrate EBF proteins, the second helix of the HLH motif is duplicated, which has been interpreted to suggest that the dimerization domains of these factors are considerably divergent from canonical HLH domains. No additional electron density for the duplicated third helix could be detected, arguing against a compact inclusion of this helix in the HLH domain.

Within the EBF family of transcription factors, the domain architecture of the individual members is strictly conserved (Liberg et al. 2002). Moreover, the three structurally investigated domains exhibit high sequence conservation, making it unlikely that major deviations in structure will be found in other EBF family members.

Ebf1 is structurally related to Rel family proteins

To explore the structural relationship of Ebf1 with other proteins, we searched for structures with homology with the TIG domain and the DBD using DALI [Holm et al. 2008]. As predicted by sequence comparison, the TIG domain is similar to the homologous domains of NFAT and NF-κB that constitute the C-terminal half of the Rel homology domain [RHD] (Supplemental Fig. S4). For the DBD, modest structural similarities to the Ig-fold DBDs of STAT, p53, and the N-terminal half of the RHD of the NF-κB and NFAT families were found (Supplemental Table S3). For both domains, NFAT1 (Giffin et al. 2003) shows the highest structural similarity to Ebf1. Moreover, the overall arrangement of the TIG-binding domains and DBDs in the DNA-bound dimers is similar, raising the intriguing possibility that the EBF proteins structurally represent a branch of the Rel superfamily [Fig. 2A].

The TIG domains mediate dimerization in all Rel family members. With the exception of an asymmetric NFAT1 dimer [Giffin et al. 2003], the interfaces in the reported structures are symmetric and consist of a hydrophobic patch surrounded by residues forming H bonds and salt bridges [Ghosh et al. 1995; Muller et al. 1995; Stroud et al. 2002]. The contact areas range from 650 Å² to 740 Å². The TIG domains of Ebf1 dimerize at the homologous surface (Supplemental Fig. S4), but the interface is markedly smaller (490 Å²) and slightly tilted compared with the Rel proteins. In addition, it lacks important residues, especially most of those mediating the polar interactions [Supplemental Fig. S4]. These differences with Rel family members probably necessitate the presence of the HLH domain for stable dimerization. The TIG domain apparently represents a genetic module that has been evolutionarily adapted by the Rel and EBF families. However, its main functions of dimerization in the Rel family and dimerization and DNA binding in NF-κB are not obvious in EBF proteins, and the role of this domain still remains elusive.

Both the DBD of Ebf1 and the N-terminal part of the RHD use a β sandwich as a scaffold for the DNA-binding loops. For Ebf1 and NFAT1, this central structure can be
The Ebf1 DBD resembles an RHD with a novel pseudo-Ig-like fold. (A) Surface representations of Ebf1 and NFAT1 (PDB: 1P7H) bound to their cognate DNA-binding sites. (B) Structure model of an Ebf1 DBD and an NFAT1 DBD shown after superposition of an NFAT1 monomer onto the second Ebf1 subunit of a DNA-bound dimer. The DNA of the superposed structures assumes an equivalent position. Regions that overlay with a Ca deviation below 3 Å are colored according to Figure 1A; structurally divergent regions are shown in gray. (C) Topology plots of the DBDs of Ebf1 and NFAT1.

superposed with a Ca RMSD of 2.14 Å [98 aligned residues] [Fig. 2B]. However, no significant sequence similarity is detected in the structurally equivalent regions [Supplemental Fig. S5], showing that the DBD of EBF proteins represents an evolutionarily independent domain. Moreover, the β strands are arranged in a different topology. While NFAT1 and all other analyzed Ig-like DBDs [see Supplemental Table S3 for a complete list and references] adopt a canonical “Greek-key” Ig-like fold, Ebf1 shows an unprecedented pseudo-Ig-like fold in which the β strands are connected in a unique manner [Fig. 2C].

The use of a different topological arrangement of the β sheets in Ebf1 and Ig-like DBDs represents an interesting example of convergent evolution.

**DNA binding by Ebf1**

Ebf1 contacts the cocrystallized DNA duplex over a region of 18 base pairs [bp], which is consistent with the size of Ebf1 footprints in DNase I protection experiments [Hagman et al. 1991]. The major and minor grooves of the DNA are contacted with three distinct modules [Fig. 3A]. A central moiety that comprises the loop between β strands X and B and the C-terminal loop recognizes bases of one half-site of the palindromic binding site, whereas the Zn knuckle contacts the other half-site in the minor groove. Unlike other dimeric transcription factors in which each monomer recognizes only one half-site of the recognition sequence, the Ebf1 dimer forms a unique symmetric clamp over the entire binding site in which each monomer contacts both half-sites. This assembly explains the strict requirement for a 2-nucleotide (nt) spacer between the half-sites, as has been noticed in mutation analyses [Travis et al. 1993]. In addition, the large loop between strands G and H (termed the GH loop) protrudes into the minor groove outside of the conserved recognition motif [Fig. 3A]. The importance of this contact is corroborated by a methylation interference assay that showed reduced binding of Ebf1 to DNA methylated in these positions [Hagman et al. 1991]. Three residues in the GH loop [N197, G203, and N204] form H bonds to the DNA backbone and bases [Fig. 3B]. While the side chain of N197 binds to two phosphate groups of the DNA backbone, the peptide nitrogen of G203 and the side chain of N204 form H bonds to bases T18 and A20, respectively. However, the contacted positions (O2 in T18, and N3 in A20) are occupied by a hydrogen acceptor, regardless of the type of base. Thus, no sequence specificity is generated by these protein:base contacts. The major contribution to specific DNA recognition is provided by the central module, which fills the major groove over one half-site of the palindrome [Fig. 3C] and contacts the nucleotides that had been found to define the Ebf recognition motif [Travis et al. 1993; Treiber et al. 2010]. It comprises the R63–F67 loop and the C-terminal loop [amino acids 234–239], which is held in position by H235. Four amino acids from this module form hydrogen bonds with the invariant bases of the recognition site: R63 reaches deep into the groove and contacts nucleotides C9 and G13. From the C-terminal loop, the backbone carbonyl of N236 and the side chains of S238 and K239 form hydrogen bonds with DNA bases contacting C8, G14, and G15.

The Zn knuckle [amino acids 157–175], which contacts the half-site from the other side in the minor groove, forms extensive hydrogen bonds with the sugar-phosphate backbone at the flanks of the groove. R163 and N172, which have been shown previously to be involved in DNA binding [Hagman et al. 1995], point into the groove. R163 binds the base and ribose of C8, whereas N172 forms a hydrogen bond with the ribose of C9 and, in addition, fixes the guanidino group of R163 in place by another hydrogen bond [Fig. 3D]. The Zn knuckle has several unusual features that, taken together, result in a unique structural motif [Supplemental Fig. S2]. With a core of just 14 residues, it is one of the shortest stretches of amino acids capable of forming an independent structural unit. In contrast to other zinc fingers [Schwabe and Klug 1994], most of its secondary structure does not consist of β sheets or loops, but very short α helices. Furthermore, DNA binding occurs in the minor groove, and is mediated by R163 in apical helix 3 and by residues in helix 4, which resembles a very short recognition helix.

**Functional validation of the Ebf1:DNA contacts**

The protein:base interactions schematically summarized in Figure 3E were validated in an electrophoretic mobility shift assay using a perfect palindromic Ebf1-binding site.
H235 (which structurally supports the C-terminal loop) and the DNA-contacting residues R63 and R163 are essential for DNA binding. Mutation of G203 to glutamate strongly reduces Ebf1:DNA complex formation by preventing binding of the GH loop in the minor groove. Mutation of other residues to alanine also reduces DNA binding (Fig. 4A). Using the natural Ebf1-binding site of the mb-1 (CD79a) promoter (Hagman et al. 1991), which consists of an imperfect palindrome, all mutations except N204A strongly reduce DNA binding (Fig. 4B). Mutation of both K146 and N147, which are involved in dimerization of the DBD (Supplemental Fig. S7), has only a modest effect on the binding of Ebf1 to the perfect palindrome, but severely affects the binding to the mb-1 site. This finding argues for a role of this contact in the cooperative binding of both DBDs to imperfect recognition sites (Hagman et al. 1995). In a model for such a cooperative mode of binding, the perfect half-site would first recruit one Ebf1–DBD, and the second DBD would then bind to a composite contact platform formed by the imperfect DNA half-site and the dimer surface of the first DBD. Natural Ebf1-binding motifs, including the single site found in the promoter of Cd79a (mb-1), frequently contain one perfect and one imperfect half-site (Hagman et al. 1991), suggesting that the dimerization of Ebf1–DBD is, at least in some promoters, functionally relevant.

In addition, we tested selected mutants of Ebf1 for their ability to activate the Ebf1 target gene Igll1 in the hematopoietic progenitor cell line BaF/3 (Fig. 4C). With the exception of K239A, all mutations tested markedly impaired the up-regulation of Igll1 by Ebf1, most likely reflecting a reduced binding to the regulatory elements in the Igll1 promoter. However, the effect of the K146A N147A double mutation is modest, which is probably due to the presence of multiple Ebf1-binding sites in this promoter. Taken together, the mutational and functional analysis confirms the role of specific residues in mediating protein–DNA interactions.

After completion of this study, the Structural Genomics Consortium (SGC) reported the structures of individual domains of Ebf1 and Ebf3 in the absence of DNA (Siponen et al. 2010). The domain structures presented by the SGC are essentially identical to the folds found in the DNA-bound dimers of our study. However, the SGC researchers did not recognize the novel topology of the DBD. In addition, they tried to computationally model the binding of Ebf1 to its recognition site. The resulting model shows two Ebf1–DBDs that form an incorrectly oriented dimer. The predicted binding to the DNA is highly asymmetric and involves contacts of the Zn knuckle to DNA bases in the major groove, none of which reflects the actual binding mechanism. Finally, Siponen et al. (2010) claim that the first two helices of the HLH domain form a new helical bundle-like fold, which is not supported by the obvious similarity with the dimerizing regions of other HLH factors (i.e., MyoD, RMSD 2.7 Å for 40 amino acids).
However, the duplicated HLH, which is not visible in our structure, is weakly defined in one monomer of the Ebf3 TIG/HLH structure, and is found to interact with the HLH of the other monomer, suggesting that it may contribute to dimerization.

Implications for the function of Ebf1 as a ‘pioneer’ factor

Ebf1 has been proposed to act as a “pioneer” transcription factor in the early steps of activation of developmentally regulated genes [Maier et al. 2004, Treiber et al. 2010]. The overall contact surface of the Ebf1 DBD dimer with the DNA duplex is nearly 3000 Å², and thus represents one of the largest interfaces reported for transcription factors. The dimerization contacts of the TIG and HLH domains on one side of the DNA and the interface between the DBDs on the opposite side lead to a complete encirclement of the DNA by Ebf1, similar to other members of the NFAT and NF-kB families [Ghosh et al. 1995; Muller et al. 1995; Stroud et al. 2002]. However, unlike the topologically constrained TIG domain of NF-kB, which mediates dimerization and DNA contacts, the Ebf1 TIG domain, which is attached via a flexible linker, contributes only to dimerization and may necessitate compensatory DNA contacts via the Zn knuckle in the Ebf1 DBD. The encirclement of DNA, which allows c-Rel homodimers to bind a consensus NF-kB site with an affinity of 0.5 nM [Sanjabi et al. 2005], most likely does not allow Ebf1 to bind its recognition motif in the context of nucleosome-wrapped DNA. In the presence of nucleosome-remodeling activity, however, the extensive DNA contacts may allow EBF proteins to stably occupy their binding sites and form barriers for nucleosome sliding. Consistent with this view, our recent ChIP-seq and ChIP analysis showed that Ebf1 needs a preconditioned or permissive chromatin context for binding to tissue-specific genes [Treiber et al. 2010]. Thus, the multiple DNA recognition and dimerization modules of the Ebf1–DNA complex may stabilize the complex in chromatin and allow for the “pioneer” function of Ebf1 in lineage-specific gene regulation.

Materials and methods

Crystallization and structure solution

Ebf126–240 (DBD) and Ebf126–422 were expressed as His, fusion proteins in Escherichia coli, and were purified using Ni²⁺ and heparin affinity chromatography and gel filtration. Detailed purification procedures are given in the Supplemental Material. Both proteins were complexed with a duplex of the palindromic 22-nt oligonucleotide 5’-CTTTATTCCTCCCTGGGAATTAAG-3’, and were crystallized using the hanging drop vapor diffusion method. Crystals of the Ebf126–240 complex were obtained at 20°C in 6% [w/v] PEG 4000, 100 mM MES (pH 5.7), and 200 mM KCl (crystal form I), 10% [w/v] PEG 4000, 100 mM NaCitrate (pH 5.4), and 10% [w/v] isopropanol. X-ray diffraction data of the three crystal forms were obtained at the Swiss Light Source (SLS) and the BESSY, including three-wavelength MAD data of Ebf126–422 complex grown at 37°C in 10% [w/v] PEG 4000, 100 mM NaCitrate (pH 5.4), whereas crystals of the Ebf126–422 complex grew at 20°C in 6% [w/v] PEG 4000, 100 mM MES (pH 5.7), and 200 mM KCl (crystal form II), whereas crystals of the Ebf126–422 complex grew at 20°C in 6% [w/v] PEG 4000, 100 mM NaCitrate (pH 5.4), and 10% [w/v] isopropanol. X-ray diffraction data of the three crystal forms were obtained at the Swiss Light Source (SLS) and the BESSY, including three-wavelength MAD data of Ebf126–422 crystals of the Ebf126–422 complex I soaked with Ta₆Br₁₄ clusters. Statistics of the acquired data sets are given in Supplemental Table S1. Phases were derived from the MAD data using SHARP [de la Fortelle and Bricogne 1997]. The other data sets were phased by molecular replacement using CCP4 Phaser [McCoy et al. 2007] and the structure model of the Ebf1 DBD-DNA complex derived from crystal form I.

Refinement

Model building was performed in COOT [Emsley and Cowtan 2004], and refinement was carried out using CCP4 Refmac5 [Murshudov et al. 1997]. The individual protein domains were defined as TLS groups in the refinement process. For structure validation, Molprobity [Davis et al. 2007] was employed. Refinement statistics are given in Supplemental Table S2. Superpositions of structures were done with LSQMAN [Kleywegt 1996], and interfaces, given as one-half BASA, were analyzed using PISA [Krissinel and Henrick 2007]. Figures of the structure models were created using MOLSCRIPT [Kraulis 1991] and rendered with POVRAY [http://www.povray.org]. Additional methods can be found in Supplemental Material.

Database entries

The coordinates of the presented structures were deposited in the Protein Data Bank [http://www.pdb.org] under accession codes 3MLN, 3MLO, and 3MLP.

Acknowledgments

We thank the staff at the SLS and BESSY synchrotrons for assistance during data acquisition. We thank P. Cramer for advice in the design of oligonucleotides, and K.P. Hopfner for discussions and comments on the manuscript. The initial crystallization and X-ray analysis of Ebf126–422 were carried out in the laboratory of G.E. Schulz at the University of Freiburg, Germany. This work was supported by funds of the Max-Planck Society and German Research Foundation.
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Genes Dev. 2010, 24: originally published online September 28, 2010
Access the most recent version at doi:10.1101/gad.1976610

Supplemental Material
http://genesdev.cshlp.org/content/suppl/2010/09/20/gad.1976610.DC1

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