The Zipper Region of Epstein-Barr Virus bZIP Transcription Factor Zta Is Necessary but Not Sufficient To Direct DNA Binding

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Epstein-Barr virus (EBV) is associated with several pathologies, including Burkitt’s lymphoma, nasopharyngeal carcinoma, Hodgkin’s disease, and lymphoproliferative diseases in immunocompromised populations (6). EBV infects the host during early childhood and persists for the life of the individual with periodic bursts of replication and virus production (6).

The viral protein encoded by BZLF1, Zta (EB1, BZLF1, ZEBRA), is a key component of the induction of the lytic replicative cycle of EBV (8). Increased expression of Zta is one of the first events that can be detected following the induction of the lytic cycle in EBV-harboring B lymphocytes, and the enforced expression of Zta is sufficient to induce the lytic cycle in a cell containing latent EBV genomes (reviewed in references 21 and 24). Zta acts in part as a transcription factor; it activates its own expression and the expression of a subset of EBV genes through sequence-specific Zta response elements (ZREs) within their respective promoters (reviewed in references 30, 31, and 32). Zta also acts as a replication factor later in the lytic cycle; Zta interacts with the lytic origin of replication, again through specific ZREs (24, 28). Furthermore, Zta also reprograms the host cell cycle control machinery, since enforced expression of Zta induces cell cycle arrest in several cell lineages (2, 3, 11, 19, 20, 25, 26), although the effect is not universal (20). Interestingly the effect on cell cycle does not require a direct interaction with ZREs but occurs through the activation of a cellular transcription factor, C/EBPα (36). Zta also interacts with a number of other cellular factors (reviewed in reference 30) that may extend its ability to regulate the expression of genes that do not contain ZREs.

Zta is a member of the family of bZIP transcription factors (5, 7, 9, 14, 16, 23, 31, 34); it contains adjacent DNA contact (approximately amino acids 175 to 195) and multimerization domains (approximately amino acids 196 to 245) and can interact directly with specific DNA sequence elements, i.e., ZREs (5, 7, 10, 16, 23, 35) as a multimer (reviewed in references 24 and 31). By analogy with other members of the bZIP family, the multimerization interface of Zta has been predicted to fold through a coiled-coil structure (10, 14, 31). Biophysical evidence that this prediction holds true was recently provided (12). However, the thermal stability of the resulting structure is much lower than that of the coiled coil domains of canonical members of the bZIP family (12). This suggests that either the Zta dimerization interface is relatively weak or that elements outside the coiled coil act to stabilize dimer formation and thus the DNA-binding function of Zta.

We present an exploration of the dimerization region of Zta. By using a short synthetic peptide, homologous to the coiled coil region, we demonstrate that the coiled coil interface of Zta is relevant for the function of Zta as a protein, despite its low thermal stability. In addition, we explore the impact of the amino acid sequence variation found within the zipper region of Zta in natural isolates of EBV on DNA-binding dependent functions. These studies suggest that the zipper region of Zta is a suitable target for therapeutic agents designed to prevent viral lytic cycle reactivation. Furthermore, we unmask the function of a novel region (CT) of Zta, adjacent to the coiled coil, which is required for both DNA-binding and transactivation functions. The impact of this on the present model of Zta structure is discussed.

In order to probe the relevance of the coiled coil region of Zta, we attempted to disrupt the formation and/or stability of the coiled coil within the context of the full-length Zta protein. The principle behind this has been exploited previously to generate reagents that block the function of the coiled coil containing proteins APC and human immunodeficiency virus gp41 (4, 15, 29, 37) and is illustrated in Fig. 1A. To evaluate whether this approach is viable for Zta, a synthetic peptide (ZEDpep) corresponding to residues spanning the coiled coil region of Zta (amino acids 196 to 227 of the B95-8 sequence) was used. Initial experiments revealed that the addition of ZEDpep reduced DNA complex formation by 76% without altering the stability of Zta under the reaction conditions (Fig. 1B).
We previously demonstrated that the three natural sequence variants within the dimerization domain of Zta fold as dimeric coiled coils but that these structures have different stabilities (the $T_m$ ranges from 17 to 25°C) (12). This was surprising, since the variant amino acids fall at the b and c positions of the heptad repeat, distant from the critical hydrophobic core formed by the a and d residues (Fig. 2A). It is therefore relevant to ask whether the different stabilities of the coiled coil motifs have an impact on dimerization-dependent functions of Zta, namely, the ability to interact with DNA in vitro and to transactivate ZRE-dependent promoters in vivo. In order to establish whether the three natural variants of the coiled coil sequence confer different properties to Zta, we reconstructed each variant within the context of the B95-8 Zta sequence, which generated vectors capable of expressing three forms of Zta termed as follows: B95-8, A205S, and A206S (Fig. 2B). All three forms of Zta interacted specifically with two different ZREs (AP1 and M) (Fig. 2C). Further analysis of the strength of the interaction revealed similar $K_d$ values for each Zta variant (Fig. 2D). The ability of each of the three variants to transactivate through ZREs in vivo was also compared. Using a reporter vector bearing seven ZREs upstream from a minimal promoter (1), together with a TK promoter as an internal control (Fig. 3A), we observed that all three Zta proteins transactivate well through the ZREs with A206S, effecting a marginally higher activation (Fig. 3B). Given that both A205S and A206S were expressed at lower levels than was B95-8 Zta in this experiment, it is clear that neither is compromised in function. Furthermore, we assessed the ability of these variants to up regulate the natural BSLF2-BMLF1 EBV promoter (27) in B cells (Fig. 3C). All three natural variants directed the up regulation of this promoter to a similar degree.

This demonstrates that, despite the differences in the stability of the coiled coil structure within the zipper region of these variants, all have similar properties with respect to the ability to bind to ZREs and to transactivate gene expression through them.

A detailed evaluation of the ability of ZEDpep to interfere with the ability of the full-length Zta proteins B95-8, A205S, and A206S was then undertaken (Fig. 4). The addition of between 4 and 10 nmol of ZEDpep compromised the ability of all three forms of Zta to interact with DNA in a dose-depen-
FIG. 3. All three forms of Zta transactivate a synthetic ZRE containing promoter equivalently in vivo. HeLa cells (obtained through ECACC) were transfected by using Effectene (Qiagen) with a total of 2 μg of DNA containing the reporter vectors, ZRE-CAT (0.5 μg) and HSVTK luciferase (0.5 μg) (Promega) (A), and either a Zta expression vector containing the full-length Zta sequences (1.0 μg) or the respective “empty” vector pBabe (22) (1.0 μg) as a control. DG75 cells were transfected by using electroporation with a total of 20 μg of DNA containing the reporter vectors, SCAT (5.0 μg) (27) and HSVTK luciferase (5.0 μg) (Promega) (C), and either a Zta expression vector containing the full-length Zta sequences (10.0 μg) or the respective “empty” vector pBabe (22) (10.0 μg) as a control. Forty-eight hours later the chloramphenicol acetyltransferase and luciferase activities were determined (Promega). The relative activation (n-fold) after correction for different transfection efficiencies is shown above the chloramphenicol acetyltransferase activity data in panels B and C. Western blot analysis of the expression of Zta given by using the monoclonal antibody BZ1 (38) is shown in panel D.

FIG. 4. The synthetic peptide is specific for Zta and can disrupt all three natural variants. The ability of ZEDpep or of an unrelated coiled coil control peptide, SKIP1 (IAALERKNALEQKAIASEYKIA LEKK [13]), to disrupt the ability of each of the three Zta full-length proteins to interact with AP1 sites was evaluated with EMSA. The DNA binding relative to the no-peptide control value is indicated. The error bars represent the standard deviation from two independent experiments.

evaluated by electrophoretic mobility shift assays (EMSA) (Fig. 5C). Data obtained from one ZRE (AP1) are shown; equivalent results were obtained for a second ZRE (M) (data not shown). Initial results from DNA-binding assays undertaken at 20°C did not fully correlate with the transactivation activity that we observed in vivo, which prompted us to further analyze DNA binding at the physiological temperature of 37°C (Fig. 5C). Truncation at 199H, carboxy terminal to the DNA-binding motif, resulted in negligible DNA binding at either temperature, emphasizing the requirement for the zipper region. Interestingly, truncation at K219 abolished DNA binding at both temperatures, but truncation at M221 retained some DNA binding at 20°C. This suggests that the zipper facilitates DNA binding but that this is insufficient at 37°C. In vivo transactivation assays on the reporter construct containing seven ZREs (Fig. 6A) and the natural BSLF2-BMLF1 promoter (Fig. 6B) revealed that only full-length Zta is capable of transactivating. It can be concluded that, although the coiled coil region is sufficient to direct DNA complex formation at 20°C, sequences C terminal to L225 (the CT region) are required for both DNA complex formation and transactivation at 37°C.

Small differences in the stability of the coiled coil structure of the natural variants of Zta that suggested that they might differ in their ability to direct stable dimer formation and thus DNA-binding-dependent functions were previously identified (12). In addition, in a recent study a 48% decrease in dimer formation was observed for A206S compared to the result for B95-8 (18). Here we assessed the potential impact of these coiled coil variants within the context of full-length Zta protein. No significant differences were found in their ability to interact with two ZREs or to direct transcription through two further ZREs. This suggests that the small differences in stability of these coiled coil structures observed by Hicks et al. and by Martel-Renoir et al. (12, 18) do not reflect differences in the function of the coiled coil when it is within the context of full-length protein. Interestingly, both of the previous studies measured dimerization in the absence of DNA and so neither would reflect any potential effects of increased stability of the
Based on our previous structural analysis of the coiled coil region of Zta, we designed a short synthetic peptide that was predicted to dimerize with Zta. This successfully disrupted the ability of the full-length Zta protein to form DNA complexes, which emphasizes the relevance of the coiled coil region for the function of Zta. In addition, it establishes the principle that the coiled coil region of Zta is a suitable target for drug design. The 50% inhibitory concentration for the Zta peptide was equivalent for all three of the naturally occurring coiled coil sequence variants of Zta, which suggests that this approach may be effective for all isolates of EBV; however, since they are all in the high-micromolar range, the synthetic peptide is unlikely to be a drug candidate itself.

The contribution of residues within the CT region (amino acids 222 to 245) of Zta to its function has been addressed previously, but the various studies reached different conclusions. Deletion of the carboxy-terminal 18 amino acids did not prevent either DNA binding in vitro or transactivation of a ZRE-dependent promoter construct in vivo (23). In contrast to this, in a recently published study of a hybrid protein containing part of Zta, a strong drop in transactivator function was observed when the carboxy-terminal 5 amino acids were deleted (pZ95dletaC5) (18). Our demonstration of temperature dependence by some of the C-terminal mutants of Zta (described in Fig. 6) may explain why the contribution from the CT region had been overlooked previously. DNA complex formation is normally assayed at temperatures between 4 and 20°C, where M221ter and L225ter retain some ability to form DNA complexes. The data presented here clearly demonstrate...
that, under the stringent assay conditions (DNA-binding assays at 37°C), the CT region is required for the DNA-binding-dependent functions of Zta. Together this demonstrates a clear role for the CT region for dependent functions of Zta both in vitro and in vivo.

It is important that some functions of Zta are independent of complex formation with ZREs (reviewed in reference 30) and that the contribution of the CT region to those functions remains unknown. The CT region has no homology to other members of the bZIP family or other proteins in available species-wide databases (as of January 2003), so further investigation is required to ask whether the CT region may act to stabilize the coiled coil of Zta or to enhance DNA-binding function through a different mechanism.

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