ABSTRACT Editing of RNA changes the read-out of information from DNA by altering the nucleotide sequence of a transcript. One type of RNA editing found in all metazoans uses double-stranded RNA (dsRNA) as a substrate and results in the deamination of adenosine to give inosine, which is translated as guanosine. Editing thus allows variant proteins to be produced from a single pre-mRNA. A mechanism by which dsRNA substrates form is through pairing of intronic and exonic sequences before the removal of noncoding sequence(s) by splicing. Here we report that the RNA editing enzyme, human dsRNA adenosine deaminase (DRADA1, or ADAR1) contains a domain (Zα) that binds specifically to the left-handed Z-DNA conformation with high affinity (Kd = 4 nM). As formation of Z-DNA in vivo occurs 5' to, or behind, a moving RNA polymerase during transcription, recognition of Z-DNA by DRADA1 provides a plausible mechanism by which DRADA1 can be targeted to a nascent RNA so that editing occurs before splicing. Analysis of sequences related to Zα has allowed identification of motifs common to this class of nucleic acid binding domain.

A well characterized editing mechanism affecting double-stranded RNA (dsRNA) involves the deamination of adenosine to produce inosine, which is translated as guanosine (1). This activity has been reported to be widespread throughout metazoans. The first example showing the physiological relevance of dsRNA editing in mammals was the replacement of a glutamine (CAG) by arginine (CGG) in the ion channel of a glutamate-responsive neuroreceptor (GluR). This change decreased the Ca2+ permeability of the receptor (2–4). Subsequently, other examples of GluR RNA editing also have been identified (5, 6), as well as editing of the serotonin-2C receptor (7) and the 4f-rnp gene from Drosophila (8). So far, two types of enzymes have been reported that are capable of performing dsRNA editing in vitro, DRADA1 (the prototype of the ADAR1 family that includes dsRAD1 and dsRAD2) and RED1 (the prototype of the ADAR2 family that includes dsRAD2) (9–15). A third protein, RED2, which has strong sequence homology to RED1, has as yet no known in vitro or in vivo substrate (16). RED1 was cloned using low-stringency in vivo DRADA2) (9–15). A third protein, RED2, which has strong and RED1 (the prototype of the ADAR2 family that includes dsRAD1 and dsRAD2) uses double-stranded RNA (dsRNA) adenosine deaminase (DRADA1, or ADAR1) contains a domain (Zα) that binds specifically to the left-handed Z-DNA conformation with high affinity (Kd = 4 nM). As formation of Z-DNA in vivo occurs 5' to, or behind, a moving RNA polymerase during transcription, recognition of Z-DNA by DRADA1 provides a plausible mechanism by which DRADA1 can be targeted to a nascent RNA so that editing occurs before splicing. Analysis of sequences related to Zα has allowed identification of motifs common to this class of nucleic acid binding domain.

from RED1. The possibility therefore exists that this difference in structure determines how RED1 and DRADA1 are used within cells.

METHODS

Identification of the Zα Domain. The Z-DNA binding domain (Zα) initially was mapped to the N terminus of DRADA1 by testing baculovirus-expressed protein and showing that band shift activity required the presence of residues 1–296. This region of DRADA1 was then expressed as a C-terminal glutathione S-transferase fusion in Escherichia coli using a pGEX-5X1 cloning vector (Pharmacia) and shown to retain Z-DNA binding activity. The Zα binding domain was mapped further both by deletion and by PCR amplification of selected portions of DRADA1 cDNA. After expression in E. coli, glutathione S-transferase-fusion proteins were purified by affinity chromatography using glutathione-agarose (Sigma), and cleaved with factor Xa (New England Biolabs) to produce Zα peptide. The peptide was additionally purified by Mono S ion exchange chromatography (Pharmacia).

Z-DNA specific binding by the Zα domain of DRADA1 was demonstrated in a bandshift assay using nonadenaturing 6% polyacrylamide gel electrophoresis (19). All assays were performed in a final volume of 20 μl and contained purified Zα (5 ng), 1 μg of sonicated salmon sperm DNA, 10 mM MgCl2, 25 mM NaCl, 25 mM Tris-HCl (pH 7.4), and 100 pg of radiolabeled Z-DNA probe (19). Specificity of Zα-binding to the probe was tested by competition with unlabeled B-form poly(dC-dG) (Pharmacia) or unlabeled, chemically brominated Z-form poly(dC-dG) (20). In addition, unlabeled supercoiled plasmid pDHg16 (21) containing a (dc-dG)13 insert, which adopts the Z-DNA conformation under the conditions used, also was used as competitor. Competition also was performed using the parental plasmid pDPL6 (21) that has no Z-DNA forming insert under these conditions.

Circular Dichroism (CD). Experiments were performed using an Aviv model 62DS spectrometer. The DNA polymer stock, (Pharmacia, average length 2,900 base pairs) was dissolved in H2O and used without further purification. All measurements were performed at 30°C with 1 ml of 25 mM NaCl/50 mM Tris-HCl/0.1 mM Na2EDTA, pH 7.4, containing DNA at a final concentration of 46 μM. Samples were allowed to equilibrate for 15 min after addition of peptide before spectra were measured. The baseline was corrected against buffer.

Determination of Binding Constant. A BIAcore instrument (Pharmacia Biosensor) was used according to the manufacturer’s specification. Three hundred response units of biontinylated poly(dC-dG), stabilized in the Z-DNA conformation by chemical bromination (20), was immobilized on a streptavidin

Abbreviations: dsRNA, double-stranded RNA; CD, circular dichroism.
coated chip (SA5). All measurements were performed at 25°C in PBS buffer (1 mM KH$_2$PO$_4$/10 mM Na$_2$HPO$_4$/137 mM NaCl/1 mM KCl/1 mM Na$_2$EDTA/0.05% Tween 20, pH 7.4) with a continuous flow rate of 20 μl/min. The association rate ($k_a$) was determined by maintaining flow at each protein concentration for a 180-s period. The dissociation rate ($k_d$) was measured by changing the flow to buffer for an additional 200 s. These values were used to calculate the equilibrium constant $K_D$ for Zα and for the Fab of the Z-DNA specific Z22 mAb.

**Secondary Structure Analysis.** Sequences were analyzed by various sequence-structure programs located on the Internet at http://www.cse.ucsc.edu/research/compbio/sam.html (Hidden Markov Model, training, alignment and database searching/scoring); http://www.sdsc.edu/meme/meme/ website (motif analysis), http://blocks.fhcrc.org (BLOCKmaker and LAMA analysis); http://bonsai.tifr.res.in/bsm/dsc/dsc reads align.html (DSC structure prediction); and http://www.embl-heidelberg.de/predictprotein/ppDoPred.html (PHD structure prediction).

**RED1.** RED1 was obtained from J. Yang (Harvard University, Boston) who had purified it from a baculovirus expression system.

**RESULTS**

We have shown previously that the chicken homologue of DRADA1 binds Z-DNA in vitro with high affinity (23). Subsequently, we have expressed different regions of human DRADA1 in E. coli and mapped a tight Z-DNA binding site (Zα) to a domain encompassed by amino acids 121–197 (Fig. 1C). Binding of Zα to Z-DNA was demonstrated using a band-shift assay that uses an oligonucleotide (dC-dG)$_6$ modified by incorporation of 5-bromodeoxyuridine, which causes the probe to adopt a Z-DNA conformation under low salt conditions in the presence of Mg$^{2+}$ (19). As shown in Fig. 1, binding of Zα to the probe is competed by unlabeled poly(dC-dG) stabilized in the Z-form by chemical bromination (Fig. 1A, lanes 4–6), but not by the unmodified B-form of the polymer (Fig. 1A, lanes 1–3). Furthermore, competition is observed with an unlabeled plasmid, pDHg16, that contains an insert maintained in the Z-conformation by negative supercoiling (Fig. 1B, lanes 4–6), but not with the parental plasmid pDPl6 that lacks this insert (Fig. 1B, lanes 1–3). These results indicate that Zα has specificity for Z-DNA, and not some other feature of the probe, such as a bromine atom. RED1, which does not have a domain equivalent to Zα, fails to bind to the probe used in this assay (data not shown). Another Z-DNA binding region, Zβ, has been identified in DRADA1 (Fig. 1C) and will be discussed elsewhere.

The structural specificity of Zα was examined further by using CD to follow the conversion of B-form poly(m$^5$dC-dG) to the Z-DNA conformation induced by Zα. Poly(m$^5$dC-dG) has been shown previously to form Z-DNA readily under physiological conditions in the presence of metal ions, polyanines, and synthetic peptides (24, 25). Formation of Z-DNA is indicated by the loss of a negative peak at 258 nm and the appearance of a new one at 295 nm. Such a change in the poly(m$^5$dC-dG) CD spectrum is indeed caused by Zα (Fig. 2), indicating that Zα stabilizes the Z-DNA conformation of DNA. This effect was maximal under the conditions used at a molar ratio of one Zα peptide per 5 base pairs of DNA. The isosbestic point of the Zα-induced transition differs from that obtained with MgCl$_2$, which also is shown in Fig. 2. The trough at 295 nm is also less negative. These results suggest that Zα may stabilize a left-handed conformation with a CD spectrum that differs slightly from the Mg$^{2+}$-stabilized Z-DNA, and indicates that the transition from B-form poly(m$^5$dC-dG) occurs in a single step. Such findings are similar to those found with an anti-Z-DNA antibody in which the spectrum of the protein-DNA complex differs slightly from that of metal-ion stabilized Z-DNA (26). In both cases, the protein traps the DNA polymer in a left-handed conformation.

The affinity of Zα for Z-DNA was further assessed by using a BLAcore instrument to measure, under physiological conditions, the direct interaction between peptide and poly(dC-dG) stabilized in the Z-DNA conformation by chemical bromination. For comparison, binding to the same polymer of an Fab prepared from the anti-Z-DNA mAb Z22 was measured (Fig. 3) (22). The equilibrium dissociation constant between Zα and Z-DNA was 4 nM and that of the Fab was 9 nM. The latter result is in agreement with previous measurements (27). However, the rates of association and dissociation of the two molecules were markedly different. Whereas the Fab demonstrated slow on and slow off kinetics, Zα showed the opposite
Quantitative association of Z-DNA with Za peptide. The Za peptide alone had no observable effect on spectra in the region of 250 to 320 nm, so spectra with DNA were not corrected for the presence of peptide. The spectra measured using Za therefore contain a large negative peak present below 245 nm that is attributable to the peptide. Molar ellipticities were calculated using base pairs of DNA. Numbers next to each spectra (solid lines) reflect the molar ratio of base pairs to peptide. The spectra obtained with DNA alone (i.e., no added MgCl₂) (B form) and DNA in the presence of 10 mM MgCl₂ (Z form) are also shown for comparison.

Sequences that show strong similarity to Za were identified by a BLAST search (28) of GenBank. In total, eight sequences from DRAD1-related proteins, two E3L proteins from vaccinia, and variola virus and a murine expressed sequence tag were found. A further search with an Hidden Markov Model (29) trained using these sequences failed to identify any more related domains (240,346 sequences searched, 0.01 significance level). Fig. 4D shows an HMM-generated multiple alignment of the Z-DNA binding domains (29). In addition to the Za domain, the human, rat, bovine, and Xenopus genes all have another Z-DNA binding domain, Zβ, that differs from Za in the amino terminus. E3L is of interest because it contains a dsRNA binding site with similarity to the dsRNA binding motifs of DRAD1, but it has no deaminase domain and is not an editing enzyme. The sequences in Fig. 4E were analyzed as a group, using position-specific scoring matrices, to identify conserved sequence and structural elements. Multiple alignment using MEME (30), a program that creates letter probability matrices for each sequence position, identified three conserved consensus motifs, as indicated in Fig. 4E. In addition, secondary structure elements were analyzed using the DSC program (31), which uses sequence variation at each position to refine structural predictions, and the PHD program, which uses a neural net strategy to achieve the same outcome (32). Three regions of strong helix formation are predicted by both programs, and these are labeled helix A, B, and C in Fig. 4C. Two of these helical regions incorporate motif I and motif II identified by the MEME program. The carboxy terminal motif III identified by MEME is predicted to be unstructured by DSC and to contain two strands of a β-sheet by the PHD program. Another conceptually different method also was used to examine the potential of these sequences to adopt particular structures. This approach uses two programs, BLOCKmaker (33) and LAMA (34), that attempt to identify structural and functional properties of proteins through sensitive detection of sequence similarities. BLOCKmaker searches protein databases for stretches of highly conserved sequence, called blocks, and records these in the BLOCKS database that can be assessed through the Internet address given in Methods. LAMA scores the relationship between blocks, allowing comparison of newly found blocks with those already known. The output of LAMA is a Z-score that is derived by Pearson correlation analysis of amino acid usage within blocks. A Z-score of 3.83 or greater almost certainly establishes that blocks are related, as all scores generated with blocks created by randomizing the BLOCKS database fall below this value. Scores of 3.83 or greater allow predictions based on the properties of a related block to be made with great confidence. Scores between 5.6 and 8.3 are suggestive of a relationship between blocks. Indeed, scores in this range are found when blocks belonging to different protein families with a common structural fold are compared (34). The predictive value of scores between 5.6 and 8.3 is increased when a particular block shows relationships to numerous other blocks with the same structural fold. BLOCKmaker was used to create a block (ZA_Block, Fig. 4D) from the aligned sequences shown in Fig. 4A. To maintain the predicted alignment of residues and because BLOCKmaker does not compensate for insertion of residues, the analysis was performed using Zβ sequences with the insertion between motif I and motif II removed. The block was tested against the BLOCKS database (33) with LAMA. Significant similarity was found with BL00043 (the helix–turn–helix ICLR family, alignment score 39, Z-score 8.1 in the top 6e⁻¹⁴ percentile of scores), BL00043 (the helix–turn–helix ICLR family, alignment score 29, Z-score 7.9 in the top 0.0e⁻¹⁰ percentile of scores), BL00042B (the helix–turn–helix CRP family, alignment score 25, Z-score 7.3 in the top 7.5e⁻⁰³ percentile of scores), BL000894A (the helix–turn–helix DEOR family, alignment score 23, Z-score 6.9 in the top 2.5e⁻⁰² percentile of scores) and with BL000356 (the helix–turn–helix LACI family, alignment score 21, Z-
score 6.7 in the top 3.5e 2 percentile of scores). When the analysis was performed with a block made without using Zb sequences, the Z-score was 7.6 with BL01051 and 7.2 with BL00043. The alignment between the COBBLER, or composite sequence, of each of the above blocks is shown in Fig. 4D. The regions of similarity map to helix B and helix C predicted by the DSC and PHD programs, and also to the helix–turn–helix region of BL00043, the CRP helix–turn–helix family (HTH CRP, BL0042B), the DEOR helix–turn–helix family (HTH DEOR, BL00894A), the LACI helix–turn–helix family (HTH LACI), and the Za group of related sequences (ZA BLOCK). The ZA BLOCK was generated after removal of the inserted R and P residues present in helix C of Zb-related sequences, a move necessary to maintain alignment of other residues during analysis. The Z-scores indicate a high probability that the relationships revealed by LAMA (34) are true positive results. For the HTH families, sequences corresponding to the DNA binding region are capitalized.

Fig. 4. Sequence and structural analysis of the Za and related domains. Human Za and Zb (hza, hzb, and HSU10439A), rat Za and Zb (rza, rbz, and RNU18942), bovine Za and Zb (bza, bzb, and this paper), two Za-related sequences (xal and xaz2) and two Zb-related sequences (xb1 and xb2) present in xenopus dsRAD1 and dsRAD2 (XLU88065 and XLU88066, respectively), the vaccinia E3L protein (S64006), and the variola equivalent (var, VVCGAA), as well as a mouse expressed sequence tag (AA204007) with relationship to Za are shown. An HMM-generated alignment of sequences (29) is presented in A, with dots indicating gaps and blanks inserted to aid in viewing the data. Residues conserved in all sequences are shown in bold. (B) Common sequence motifs present in this group of sequences. These motifs were extracted using the MEME program (30) and are presented as a multilevel consensus sequence. (C) Structural prediction for Za made using the DSC program (31), which incorporate sequence variation at each position to improve accuracy. H is used to indicate predicted regions of helix, E for b-sheet residues, and C for coiled or loop elements. An alternative prediction made using the PHD program also is shown (32). (D) Alignment between the COBBLER (or composite) sequence of the blocks (33) that characterize the ICLR helix–turn–helix family (HTH ICLR, BL01051A), the GNTR helix–turn–helix family (HTH GNTR, BL00043), the CRP helix–turn–helix family (HTH CRP, BL0042B), the DEOR helix–turn–helix family (HTH DEOR, BL00894A), the LACI helix–turn–helix family (HTH LACI), and the Za group of related sequences (ZA BLOCK). The ZA BLOCK was generated after removal of the inserted R and P residues present in helix C of Zb-related sequences, a move necessary to maintain alignment of other residues during analysis. The Z-scores indicate a high probability that the relationships revealed by LAMA (34) are true positive results. For the HTH families, sequences corresponding to the DNA binding region are capitalized.

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Intercalation of tryptophan between a dG-dC step has been reported for the human ETS1-DNA complex (38) (discussed in ref. 43), whereas intercalation of proline is found in an IHF-DNA complex (46). Alternatively, the conserved proline and tryptophan may be required for correct protein folding. The variation in the amino terminus between Za, Zb, and E3L may allow specific interaction with other proteins, and thus affect binding specificity or function of these domains.

**DISCUSSION**

The potential for involvement of Z-DNA in biological processes has been much discussed since the x-ray crystallographic description of this structure (47). The demonstration that Z-DNA could be stabilized by negative supercoiling brought this conformation into the realm of the biologist (48, 49). Subsequently, it was shown that formation of Z-DNA occurs *in vivo* and in agarose-embedded, permeabilized, metabolically active nuclei as a result of transcription-induced supercoiling in the underwound region 5′ or behind a moving RNA polymerase (50–52). The level of unrestrained supercoiling present *in vivo* nevertheless is limited by the relaxing action of topoisomerases and the accommodation of negative supercoils into nucleoprotein structures. Due to the transient nature of Z-DNA in *vivo*, direct experimental demonstration of the involvement of Z-DNA in biological processes has been difficult. The indirect approach of finding Z-DNA binding proteins also has been beset by methodological problems (53, 54).

The data presented here shows that a natural protein exists that is specific for Z-DNA. The nuclear location of this protein and the high affinity for Z-DNA make it unlikely that this finding is adventitious, underscoring the possibility that this non-B-DNA structure is exploited by nature in regulation of biological processes.

The nature of the interaction of Za with Z-DNA will await structural studies. The CD experiments show a difference between salt-induced and Za-stabilized Z-DNA. This outcome may reflect the sensitivity of CD to changes in the close environment of Z-DNA, possibly due to binding of Za to the convex outer surface of Z-DNA. Alternatively, Za may induce changes to the helical parameters of Z-DNA. For example, Za may interact with the minor groove of Z-DNA in a manner analogous to that seen with some transcriptional regulators that bend the DNA helix (38, 55, 56). Alternatively the B-Z junction may be recognized by Za. At the B-Z junction, the change in helical direction is associated with an inversion of base pairs. The major groove of B-DNA comes to overlie the minor groove of Z-DNA whereas the minor groove of B-DNA merges with the major convexity of Z-DNA. This area of transition has the potential to create structurally unique shapes to which Za might bind (see for example ref. 57). The junction area also may be bent (58), providing sites where the intercalation of the carboxy tryptophan (or other amino acids) could occur. However, binding involving intercalation might be expected to involve slower kinetics than those shown in Fig. 3. It is of interest that extremely slow association and dissociation phases, suggestive of a second binding mode, are found when Za dimers rather than monomers are studied (data not shown). Potentially Za could bind solely to the major groove of B-DNA adjacent to a B-Z junction. This event is unlikely as the B-DNA form of dC-dG (Fig. 1) does not affect binding of Za to Z-DNA polymers.

An unresolved question in these studies is the role of Za in the biology of the editing enzyme DRADA1. Use of introns to form dsRNA, as has been demonstrated for editing of glutamate-responsive neureceptor subunit RNAs (5, 6, 59–61), means that DRADA1 must act soon after transcription and before splicing of the nascent pre-mRNA. Targeting of DRADA1 to a transcription-dependent structure, such as Z-DNA would facilitate its localization to nascent transcripts, so that editing occurred before splicing. It may turn out, however, that Z-DNA binding domains of DRADA1 regulate transcriptional events other than editing.

Demonstration of an interaction between Za and DNA within living cells may be possible using rapid UV-laser-induced crosslinking techniques (62). Potential Z-DNA-forming sequence motifs are present in many human genes (63). Binding of Za to such segments could indicate that under appropriate levels of negative supercoiling, pre-mRNA from that gene is edited. Such studies might provide further insight into the possible regulation of DRADA1 by Z-DNA and provide insight into the evolution of this editing mechanism (64).

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