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Coevolution of a Homing Endonuclease and Its Host Target Sequence

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Received 3 July 2007; received in revised form 20 July 2007; accepted 24 July 2007 Available online 2 August 2007 We have determined the specificity profile of the homing endonuclease I-AniI and compared it to the conservation of its host gene. Homing endonucleases are encoded within intervening sequences such as group I introns. They initiate the transfer of such elements by cleaving cognate alleles lacking the intron, leading to their transfer via homologous recombination. Each structural homing endonuclease family has arrived at an appropriate balance of specificity and fidelity that avoids toxicity while maximizing target recognition and invasiveness. I-AniI recognizes a strongly conserved target sequence in a host gene encoding apocytochrome B and has fine-tuned its specificity to correlate with wobble versus nonwobble positions across that sequence and to the amount of degeneracy inherent in individual codons. The physiological target site in the host gene is not the optimal substrate for recognition and cleavage: at least one target variant identified during a screen is bound more tightly and cleaved more rapidly. This is a result of the periodic cycle of intron homing, which at any time can present nonoptimal combinations of endonuclease specificity and insertion site sequences in a biological host.

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Homing is the transfer of an intervening genetic sequence (either an intron or an intein) to a homologous allele that lacks that same sequence, $^{1-3}$ leading to gene conversion and transmission of a mobile element. Homing endonucleases promote the mobility of these intervening sequences, which include their own reading frames, by generating doublestrand breaks in homologous alleles that lack the intron or the intein. Break repair leads to transfer of the element *via* homologous recombination, using the allele that contains a homing endonuclease gene as a template. In rare cases, homing endonucleases can also be encoded by free-standing genes.⁴ In either case, homing endonuclease genes are selfish DNA sequences that are inherited in a dominant, non-Mendelian manner.^{5–7}

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Abbreviations used: *A.n., Aspergillus nidulans; cob,* apocytochrome *B* oxidase; IC, information content; ITC, isothermal titration calorimetry; LHE, LAGLIDADG homing endonuclease.

Homing endonucleases and their mobile introns are found in virtually all microbial genomes, including phage, bacteria, archaea, protista, and organellar genomes in fungi and algae.^{1,2,8–10} A variety of analyses have indicated that mobile introns and homing endonucleases display a periodic life cycle in these host genomes:^{5–7} invasion of a DNA target is followed by subsequent vertical transmission, eventual loss of endonuclease activity in individual hosts, loss of the endonuclease reading frame and eventually the intron, and subsequent reinvasion by an active homologue. In the case of phage and prokaryotes, this cycle causes the persistence of introns in genomes that are otherwise subject to strong purifying selection and streamlining.¹¹

Homing endonucleases recognize long DNA sequences (typically 14–40 bp), which reduce their toxicity in the host organism. Homing endonucleases tolerate sequence variation at individual base pairs within their targets; this attenuated fidelity enables them to adapt to sequence drift in their target and increases their potential for ectopic transfer. There are at least five discrete homing endonuclease families, each of which has arrived at the optimal balance of specificity *versus* fidelity that is most suitable for evolutionary success in their host genomes. These families are classified and named according to their most conserved sequence and structural motifs, and generally are localized and constrained to distinct biological and genomic niches.¹²

Members of the largest family, termed 'LAGLI-DADG' homing endonucleases (LHEs), are most often encoded in archaea and in the mitochondrial and chloroplast genomes of single-cell eukaryotes.¹³ LHEs are segregated into two subfamilies, based on the presence of either one or two copies of the conserved LAGLIDADG motif per protein chain. Enzymes that contain a single copy of this motif act as homodimers and recognize consensus DNA target sites that are constrained to palindromic or near-palindromic symmetry. In contrast, LHEs that have two copies of the motif act as monomers, possess two structurally similar nuclease domains, and are free to evolve towards recognition of asymmetric DNA targets. Protein domains are connected by flexible linker peptides with lengths ranging from 3 residues to over 100 residues.¹⁴ Freestanding LAGLIDADG endonucleases (those that are not associated with additional protein domains) recognize DNA sites that range from 18 to 22 bp.¹⁵⁻¹⁸ They cleave both DNA strands, generating mutually cohesive four-base 3' overhangs.^{15,19,20} Like most nucleases, LHEs require divalent cations for activity.

The structures of several LHEs bound to their DNA targets have been determined. These include three homodimers (I-CreI,^{21–24} I-MsoI,²⁴ and I-CeuI²⁵), two pseudosymmetric monomers (I-AniI²⁶ and I-SceI²⁷), one artificially engineered chimeric protein (H-DreI²⁸), and one intein-associated endonuclease (PI-SceI²⁹). Structures have also been determined of several additional LHEs in the absence of DNA. Although the primary sequences, DNA target specificities, and oligomeric architectures of LHEs have diverged significantly, their overall topologies, tertiary folds, and DNA-binding mechanisms are still strongly conserved.

Many LHEs are encoded within introns that interrupt highly conserved rDNA genes, each of which can simultaneously harbor large numbers of separate group I introns with associated homing endonuclease reading frames (e.g., the 23S rDNA in *Chlorosarcina brevisisinosa*).^{30,31} The specificity of two of such endonucleases (I-CreI and I-CeuI, homodimers encoded in introns within the 23S rDNA of Chlamydomonas reinhardtii and Chlamydomonas eugametos, respectively) has been previously measured and found to be strongly correlated with the number and type of protein/DNA contacts found in a DNA-bound complex.^{24,25} Mobile introns can also inhabit protein-coding genes in many microbial species, including phage (e.g., the DNA polymerase gene in SPO1 and Bastille³²), in bacteria (e.g., the *nrdE* gene in *Bacillus anthracis*),³³ and in organellar genomes (such as the I-AniI endonuclease described in this work). However, these mobile introns are typically found at fewer

sites within individual host genes, and their specificities are generally less well characterized.

The I-AniI protein is a monomeric 250-residue LAGLIDADG endonuclease encoded within a group I intron that interrupts the mitochondrial apocytochrome *B* oxidase (*cob*) gene of *Aspergillus nidulans* (A.n.).³⁴ I-AniI recognizes and cleaves a 19-bp asymmetric DNA target site. The enzyme binds its DNA substrate with approximately nanomolar affinity and cleaves the noncoding upper strand more rapidly than the coding lower strand.35,36 Homologous introns and homing endonucleases are found in the same genetic location in related host organisms, including Saccharomyces cerevisiae³⁷ and Venturia inaequalis.³⁸ These two proteins display 49% and 65% sequence identity, respectively, to I-AniI. Expanded analyses of recently deposited cob sequences from microbial genomics projects indicate that at least one additional position in the gene has been invaded by a mobile intron³⁹; the specificity of one of these (I-CsmI, from Chlamydomonas smithii) has also been studied in detail, with many of those results agreeing with the analysis described here.⁴⁴

I-AniI and its host gene offer an excellent system for a detailed analysis of the coevolution of a homing endonuclease and a corresponding protein-encoding host gene and target site. In this study, we describe the specificity profile and overall fidelity of the I-AniI endonuclease at each base pair of its target site using a high-throughput cleavage screen of a randomized substrate library. We identify and characterize one target variant that is a significantly superior substrate as compared to the physiological DNA target sequence in *Aspergillus*. The specificity profile of the endonuclease is significantly correlated to the reading frame of the host gene, down to the position of individual base pairs at wobble versus nonwobble positions in individual codons. This analysis illustrates the degree to which homing endonuclease specificity is 'fine-tuned' to its natural target site and also demonstrates the extent to which a viable intron invasion site can be diverged from an optimal DNA target sequence for a homing endonuclease.

Results

Recovery of cleavable DNA substrates

The site specificity of the I-AniI homing endonuclease was assayed across its entire 19-bp DNA recognition target by the selection of cleavable DNA site variants from a randomized site library under optimal cleavage conditions as described above. Information content (IC; specificity) was calculated at each base pair using the published algorithm of Schneider *et al.*⁴¹

Seventy-five unique cleavable target sequences were identified in the screen (Figs. 1 and S2). Enriched from an initial library containing an average of 6.7 mutations per target site, two thirds of cleavable substrates (50 of 75) contain a single

| Wild-type | TGAGGAGGTTTCTCTGTAA | | |
|---------------------------|---|--------------|--------------|
| Recovered (out of 428) | Sequence | # of parents | # Mismatches |
| 114 | TGAGGAGGTT <mark>G</mark> CTCTGT <mark>T</mark> A | 7 | 2 |
| 50 | TGAGGAGGTTGCTCTGTAA | 9 | 1 |
| 49 | TGAGG <mark>T</mark> GGTT <mark>G</mark> CTCTGT <mark>C</mark> A | 2 | 3 |
| 30 | TGAGG <mark>T</mark> GG <mark>G</mark> TCCTCTGTTA | 1 | 4 |
| 23 'Lib4' | TGAGGAGGTTACTCTGTTA | 2 | 2 |
| 18 | TGAGGAGG <mark>G</mark> TTCTCTGTAA | 11 | 1 |
| 10 | TGAGG <mark>T</mark> GGTT <mark>G</mark> CTCTGTAA | 3 | 1 |
| 7 | TGAGGAGG <mark>G</mark> TTCTCTGT <mark>T</mark> A | 2 | 2 |
| 6 | TGAGGAGG <mark>C</mark> TTCTCTGT <mark>T</mark> A | 3 | 2 |
| 5 | TGAGG <mark>G</mark> GG <mark>G</mark> T <mark>C</mark> CTCTGT <mark>T</mark> A | 1 | 4 |
| 5 | TGAGGAGGTT <mark>C</mark> CTCTGTAA | 5 | 1 |
| 4 | <mark>G</mark> GAGGAGGTT <mark>G</mark> CTCTGTAA | 1 | 2 |
| 4 | TGAGGAGGTT <mark>G</mark> CTCTGT <mark>C</mark> A | 2 | 2 |
| 3 | TGAGGAGGTT <mark>A</mark> CTCT <mark>C</mark> T <mark>A</mark> | 1 | 3 |
| 3 | TGAGGAGG <mark>G</mark> TTCTC <mark>C</mark> GT <mark>T</mark> A | 1 | 3 |
| 3 | TGAGGAGGT <mark>CA</mark> CTCTGTAA | 1 | 2 |
| 3 | C GAGGAGGTT <mark>A</mark> CTCTGTAA | 2 | 2 |
| 3 | TGAGGAGG <mark>G</mark> T <mark>A</mark> CTCTGT <mark>C</mark> A | 1 | 4 |
| 3 | TGAGGAGGT <mark>C</mark> TCTCTGT <mark>T</mark> A | 1 | 2 |
| 2 wt → | TGAGGAGGTTTCTCTGTAA | 2 | 0 |
| 2 | TGAGG <mark>G</mark> GGTTTCTCTGT <mark>C</mark> A | 2 | 2 |
| 2 | TG <mark>T</mark> GG <mark>T</mark> GGTT <mark>C</mark> CTCTGT <mark>T</mark> A | 1 | 4 |
| 2 | TGAGGAGG <mark>C</mark> T <mark>C</mark> CTCTGTAA | 1 | 2 |
| 2 | TGAGGAGG <mark>A</mark> T <mark>GT</mark> TCTGTAA | 1 | 3 |
| 2 | TGAGG <mark>G</mark> GGTTTCTCTGT <mark>T</mark> A | 1 | 2 |
| 2 | TGAGGAGGTT <mark>A</mark> CTCTGT <mark>TG</mark> | 1 | 3 |
| 2 | TGAGGAGGTT <mark>A</mark> CTCTGTAA | 1 | 1 |
| 2 | TGAGGAGG <mark>C</mark> TTCTCTGT <mark>C</mark> A | 1 | 2 |
| 2 | TGAGGAGG <mark>A</mark> T <mark>A</mark> CTCTGTAA | 1 | 2 |
| 2 | TGAGGAGGT <mark>C</mark> TTCTGTAA | 1 | 2 |

(+ 45 additional sequences that appeared once each: see Figure S2 in supplementary information)

Fig. 1. Target sites recovered in a specificity measurement screen of the I-AniI endonuclease. In this experiment, 428 substrate targets were sequenced. The wild-type site was isolated twice in this screen and is indicated; individual mismatches in other sites, relative to the wild-type target, are highlighted in yellow. The center of the target site (dividing the left and right half-sites) is indicated with the vertical line. In addition to individual sequences of cleavable substrates, the number of individual parental clones for each target can be identified, based on unique sequences of randomized, addressable flanking bases. Out of 76 unique sequences, 31 appeared more than once, out of which half were generated from multiple parental site clones. The frequency at which individual sites were recovered did not correlate strongly with their performance in kinetic cleavage assays; however, at least one site ('Lib4') is a much better substrate than the physiological, wild-type target sequence (see Fig. 4).

base-pair substitution, while the majority of the remainder contain two (15 of 75) or three (5 of 75) mismatches.

Seven unique sites account for 294 of 428 recovered DNA targets, corresponding to over 65% of the isolated endonuclease substrates in this experiment (Fig. 1). The remainder of cleavable DNA sequences were recovered at much lower frequencies. The enhanced recovery of the most frequently isolated target sequences was not due to either the initial bias of the initial DNA template synthesis (as demonstrated by the sequencing and analysis of clones in the naive library) or a significant bottleneck in early cycles of selection (all but one of these frequently recovered substrates arose from multiple parental clones).

As described below, the more frequently recovered DNA target sequences are usually found to be cleaved slightly more rapidly than are those that were recovered only once. One of the most frequently recovered substrates (labeled 'Lib4' in Fig. 1), containing 2-bp substitutions, was found to be a superior substrate as compared to the wild-type sequence from *Aspergillus* and to other sequences recovered from the library that were tested in detail.

In addition to the cleavable targets analyzed in this study, one clone containing a site that harbored an unusually high number (8) of mismatches relative to the wild-type I-AniI target was recovered once in the final output of the selection. Subsequent *in vitro* analyses of this site indicated that I-AniI exhibits no cleavage activity on this site under the conditions described above and likely represents a spurious target site sequence that slipped through the target site screen. The remaining target sites that were retested individually for cleavage were viable substrates for the enzyme.

Site specificity and fidelity

While individual base-pair substitutions can apparently be tolerated by the endonuclease at

many positions under the reaction conditions used in this experiment, on average, cleavable substrate sequences only deviated by 1.4 mismatches from the cognate physiological target site (relative to the average of seven mutations per site that were incorporated into the naive library). The specificity of DNA recognition at individual base pairs is highly variable across the target site (Fig. 2a). Nine positions (-9, -7, -6, -4, -3, +3, +4 +6, and +7) display nearly invariant base preference, four positions (-5, -2, +1, and +8) display little or no sequence preference, and the remaining six sites (-10, -8, -1, +2, +5, and +9) display attenuated specificity. In addition, two positions (-1 and +1)display mild base covariation, with an $A \rightarrow C$ substitution at position +1 being dependent on the conservation of the wild-type A:T base pair at the neighboring -1 position. These positions correspond to the direct center of the DNA target site and are subject to significant base unstacking upon binding of the enzyme. Such positions in DNA target sequences often display sequence-specific conformational preferences that contribute to overall protein-binding specificity.

Across the central-four base pairs of the target site (which are not in direct contact with protein atoms and are subject to binding specificity primarily as a result of DNA bending), three individual base-pair substitutions ($-1 \text{ T} \rightarrow G$, $+2 \text{ C} \rightarrow G$, and $+2 \text{ C} \rightarrow A$) completely inhibit target site cleavage, while two substitutions ($+1 \text{ T} \rightarrow C$ and $+1 \text{ T} \rightarrow A$) yield substrate cleavage efficiencies equal or superior to the wild-type target (Fig. 2b). The remaining seven individual substitutions across the central-four bases yield relatively minor changes in cleavage efficiency. The T $\rightarrow A$ substitution at position +1 is also observed in the hypercleavable 'Lib4' target site (Fig. 3) and was isolated as an individual substitution from the randomized site library screen.

The overall specificity of I-AniI is extremely high, even under optimal *in vitro* cleavage conditions as described here. The total number of potential DNA target sequences for this endonuclease is 4^{19} or 2.7×10^{11} sites for the 19-bp target site. Based solely on the IC at each base pair described above, approximately 16,384 of these sequences are predicted to be cleavable (corresponding to nine positions that are invariant or nearly so, four positions



Fig. 2. Specificity of DNA recognition by I-AniI. (a) IC at individual positions across the I-AniI target site, calculated using recovered sequences from a screen of site variants. The corresponding translated protein sequence of the host gene is provided below the DNA target sequence (note that in fungal mitochondrial genomes, 'TGA' encodes for Trp rather than for 'stop'). IC (specificity of recognition) was calculated at each base pair according to the method of Schneider *et al.* ("2" = invariant base preference). Arrows indicate wobble positions in host-coding frame (average IC=1.0, compared to 1.7 for nonwobble positions). (b) Cleavage of individual single base-pair substitutions across the 'centralfour' positions in the target site. Three substitutions (-1 G, +2 G,and +2 A) are strongly disfavored, two (+1 G and A) are equivalent or superior to wild-type, and the remainder are slightly reduced but within 2-fold of wild-type cleavage efficiency.



Fig. 3. Schematic of the methodology for *in vitro* screen for cleavage target site variants. See main text and Supplementary Data for a full description of experimental details. (a) Design of the cyclical screen and enrichment for cleavable target sites. (b) Progression of the screen from the naive library to the final convergence in round 4.

that can tolerate any of the possible base pairs, and six positions that can approximately accommodate either of two base pairs: $1^9 \times 2^6 \times 4^4 = 16,384$). However, this number represents a significant underestimate of the enzyme's substrate specificity, as it assumes that substitutions at all the 10 variable positions listed above can be accommodated by the enzyme simultaneously (taking into account neither covariation between adjacent base pairs nor the observation that cleavable substrates isolated in this screen harbor an average of only 1.4 mutations per site). Thus, the number of theoretically cleavable targets should be reduced by at least 7-fold (the ratio of 1.4 mutations per cleavable site, on average, to 10 base-pair positions that can potentially be altered in any given substrate). This results in a prediction of

approximately 2340 cleavable sites out of over 2×10¹¹ potential substrate sequences (or one cleavable site per 10⁸ random sequences). Extrapolated to a mammalian genome ($\sim 10^9$ bp), one would predict that I-Anil will be capable of cleaving approximately 10 sites per genome under optimal in vitro cleavage conditions. However, under less optimal reaction conditions (e.g., reduced enzyme concentration, or in the presence of high concentrations of competing DNA-binding proteins and highly ordered chromatin structure as found in vivo), as many 'marginal' substrates would likely be poorly bound and cleaved. These observation and interpretation are in agreement with many reported studies on the transfection of mammalian and/or human cells with nuclearly localized homing endonucleases, including I-SceI^{42,43} and I-PpoI,⁴⁴ that demonstrate levels of double-strand-break foci and cell toxicity that are not significantly above background measurements for untransfected cells. Similarly, the transformation of human cells with the I-AniI reading frame, and the subsequent expression and nuclear localization of high levels of active endonuclease do not appear to cause measurable cell toxicity or elevated genomic double-strand breaks, even though at least four potentially cleavable targets are known to exist in the human genome (M.S.K. and B.L.S., unpublished observations).

Correlation of I-Anil specificity with constraints of its host target gene sequence

The mitochondrial *cob* gene in fungi has been the target of at least two historical invasion events by an LHE, one of which is represented by I-AniI and its close homologues in *S. cerevisiae* and *V. inaequalis*.^{37–39} This protein-encoding host gene shows a typical pattern of conserved and nonconserved residue positions, with nonconserved amino acids observed at approximately 25% of the positions in its reading frame (Fig. 4). Both group I introns are located at positions in the cob gene that encode exceptionally well-conserved amino acid sequences, each extending over approximately six codons. The translated sequence for the I-AniI target site (using the fungal mitochondrial genetic code, where TGA='Trp,' rather than 'stop') is 'WGGFSV,' a peptide sequence that lies within a surface-exposed tight turn that connects two helices near the enzyme active site (Fig. S3). The two glycine residues in this sequence appear to be essential for the structure of this region of the enzyme subunit, as both exhibit backbone dihedral conformations that are disallowed for all other protein residues. Two of the three large hydrophobic residues (W and V) lie in the core of the protein, while phenylalanine (F) is completely solvent-exposed. Apparently, this particular protein sequence is tightly constrained to maintain the function of the enzyme.

A comparison of the specificity of I-AniI at each base pair to its cognate target sequence with the reading frame of the host gene demonstrates a very strong correlation between specificity at individual DNA bases and the constraints on each of those bases imposed by the degeneracy of the amino acid code (Fig. 2a). The average IC at nonwobble positions in the target sequence is approximately 1.7, whereas the same measurement of specificity at wobble positions indicates a much lower value of 1.0 (i.e., the endonuclease is more tolerant of target site variation at those positions). Furthermore, the reduced specificity at the wobble positions of the host gene is most pronounced for three of the more degenerate codons in the target site that encode two glycines and one valine. In contrast, two of the wobble positions that display only slightly reduced, 'intermediate' specificities correspond to host codons for Trp and Phe (each of which allows only two possible bases at their wobble position). Therefore, the correlation of the specificity of the homing endonuclease, relative to sequence constraints on the host-gene-coding sequence, is extraordinarily 'fine-tuned' to the most possible forms of genetic variation that can occur in a highly constrained target position. Once incorporated into a host target site, it seems likely that subsequent vertical transmission of a homing endonuclease gene is maximized by continuing to evolve to be particularly tolerant of those DNA base substitutions that are silent during protein translation and thus are likely to experience accelerated genetic drift.

The one notable 'outlier' in the specificity profile and trend described above is the lack of obvious specificity at position +1. While there is no obvious explanation for this result, it is worth noting that this



Fig. 4. Conservation of the *cob* gene from sequenced microbial genomes, and the position of known inserted group I introns. For the *A.n.* cob protein sequence, a set of homologous sequences was collected using an iterative PSI-BLAST search of the nonredundant protein sequence database. A multiple-sequence alignment was made from the PSI-BLAST output, allowing for the calculation of IC at each position. IC values were linearly normalized from 0 to 1, with 0 corresponding to the least conserved position and 1 corresponding to the most conserved positions. The arrows point to conserved regions of the host gene targeted by I-CsmI and I-AniI.

position happens to be one of two positions that are decidedly 'suboptimal' for recognition (being one of two base pairs that are mutated in a highly cleavable 'Lib4' site, as described below), which may be reflected in the low measured IC at that position.

Comparison of wild-type and variant target sites as endonuclease substrates

In order to determine whether the relative recovery of various DNA target sequences was strongly correlated with their behavior as endonuclease substrates, a panel of sites from the screen was characterized in more detail using *in vitro* cleavage assays (Fig. 5). Five of the most frequently isolated targets, five of the targets each isolated once, and the wild-type target from *Aspergillus* were subjected to individual kinetic cleavage analyses. These experiments demonstrate that the more 'successful' targets from the screen are cleaved, on average, 50% more rapidly than those that were each isolated only once.

One of those target sites ('Lib4') displayed much more efficient cleavage than the wild-type target (Fig. 5). Binding analyses of the relative affinity of



Fig. 5. Relative behavior of isolated target sites as cleavage substrates for I-AniI. (a) Kinetics of cleavage under reaction conditions as described in Materials and Methods. Extent of cleavage quantitated at 30, 60, and 120 min. The numbers above the individual substrate bars correspond to the number of times each sequence was isolated in the screen, as shown in the first column of Fig. 3. Note the complete rapid digestion of 'Lib4' relative to the wild-type target sequence. (b) Relative binding of wild-type and 'Lib4' substrates by the I-AniI endonuclease, as determined by ITC. K_d values are averages of multiple independent experiments conducted on separate days, which reproducibly indicate that the 'Lib4' sequence is bound approximately 10-fold more tightly than the wild-type sequence. While the estimated molar ratios of binding for the experiments shown in these panels are slightly above and below 1:1, respectively, the average of multiple runs in each case indicates a 1:1 binding stoichiometry of protein to DNA duplex. The variation in this value illustrated in this figure is typical of ITC runs with homing endonuclease/DNA systems.

this substrate compared to the wild-type target indicated that this target site is bound approximately 10-fold more tightly (with a reduction of -1.5 kcal/ mol in $\Delta G_{\text{binding}}$) than the wild-type sequence. This improved affinity is largely driven by a significant improvement (-4.5 kcal/mol) in enthalpy change (ΔH) upon binding, which implies that the new target site provides new contact points (presumably additional hydrogen bonds) to the protein.

This target site contains two base-pair substitutions, both on the right DNA half-site, at base pairs +1 and +8. The endonuclease does not display obvious direct contacts to either of these base pairs in the wild-type complex, also implying that these two substitutions might introduce novel interactions in the protein/DNA interface that enhance affinity and improve cleavage efficiency. Subsequent cleavage analyses of two separate sites that each harbor a single individual substitution present in the 'Lib4' site demonstrate that both contribute to tighter binding and more rapid cleavage, with the majority of improved recognition derived by the $A \rightarrow T$ substitution at position +8 (data not shown). Neither the 'Lib4' sequence nor a sequence containing the single mismatch at position +1 (both of which would encode a Phe→Leu point mutation in the translated host protein) was found in a search of all currently sequenced *cob* genes. In contrast, approximately 30 known cob sequences contain the 'Lib4' mismatch at base pair +8, which corresponds to a silent mutation at the wobble position within the valine codon.

Given the evolutionary dynamics of mobile introns and homing endonucleases (and most, if not all, other invasive genetic elements), it would not be

surprising if many naturally isolated combinations of endonuclease and homing site sequences represented nonoptimal pairing of enzyme specificity and substrate sequence, as described above for I-AniI. A variety of analyses have indicated that mobile introns and homing endonucleases display a periodic life cycle in their host genomes.^{5–7} Horizontal transfer into a novel DNA target is followed by subsequent vertical transmission and expansion across closely related homologous genes, eventual loss of endonuclease activity in individual hosts, loss of the endonuclease reading frame and eventually of the intron, and subsequent reinvasion by an active homologue. Estimates of the periodicity of intron invasion, loss, and reinvasion are typically several million years.

The structure of the DNA-bound endonuclease complex: details of recognition and cleavage

The I-AniI/DNA complex structure²⁶ was extended to 2.4 Å resolution, giving a more detailed view of both the protein/DNA interface and the I-AniI active site when compared to the previously determined structure. The structure consists of a full-length enzyme monomer and a 28-bp DNA duplex containing the 19-bp native homing site from the *Aspergillus* mitochondrial cob intron. The refined structure additionally includes 47 water molecules, 10 of which mediate contacts in the protein/DNA interface. Three metal ions were modeled into overlapping active sites, and the DNA was observed to be uncleaved. Details of active-site metal coordination are provided in Supplementary Data and Fig. S4.



Fig. 6. I-AniI/DNA interface and contacts, modeled upon the refinement of the structure to a resolution of 2.4 Å. The top strand corresponds to the noncoding strand of the *cob* host gene. The acidic residues from the LAGLIDADG motif that are involved in metal binding are D16 and E148.

The variability of specific recognition and fidelity across the DNA target site is a function of the contacts made between the endonuclease and the individual base pairs of the substrate. Side chains from the β -sheet DNA recognition surface of each individual I-AniI domain contact nucleotide bases in each of the two corresponding DNA target half-sites (Fig. 6). All contacts to nucleotide bases are made in the major groove. Eighteen of the 38 phosphate groups (47%) are in direct contact with the protein. A total of 18 direct contacts, 10 of which are watermediated, are made between hydrogen-bond partners on the protein and corresponding hydrogenbond donors and acceptors on DNA bases in the target site. This corresponds to roughly 30% of potential hydrogen-bond sites across the DNA target's major groove.

I-AniI interacts with its DNA target through a strikingly asymmetric distribution of contacts, with 12 contacts made to bases in the left half-site and with five contacts made to bases in the right half-site. In addition, the majority of direct nucleotide base contacts (15 of 18; 83%) are made on the top noncoding strand of the target DNA. Arginine contacts to a run of purine bases on the top strand of the left half-site account for eight of the direct contacts. These interactions correspond to the DNA target sequence 5'-GGAGG-3' from base pairs -7 to -3, and protein residues Arg72, Arg70, Arg59, and Arg61. Two of these residues, Arg59 and Arg72, contact two neighboring purine residues simultaneously.

Unlike the previously studied I-CreI LHE,24 I-AniI exhibits a relatively loose correlation between measured IC at individual base pairs and the number of direct contacts at each position within the target site. Positions +7 and +9, for example, exhibit high IC (1.9 and 1.6, respectively), yet I-AniI makes no direct contacts to nucleotides at these positions. One partial explanation for this difference in behavior might be that whereas I-CreI appears to be highly optimized for the recognition of its cognate target site in Chlamydomonas (displaying a cognate K_d value of ~1 nM), I-AniI is clearly less well optimized towards its target site in Aspergillus, as described above: at least two positions in the target site (not visualized in the 'wildtype' complex) yield tighter binding when mutated and may provide additional contacts that improve the correlation between structure and binding specificity.

In addition, the structure of I-CreI was determined at 1.9 Å resolution, allowing us to model a total of 28 water molecules that mediate contacts in the protein/DNA interface, accounting for about half of the protein/DNA contacts. In contrast, only 10 water molecules can be confidently modeled into the I-AniI protein/DNA interface in the 2.4-Å resolution structure. Therefore, the structure may be missing several important contactmediating water molecules from the protein/DNA interface that can only be visualized at higher resolution.

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Discussion

The success or failure of a homing endonuclease lineage is dictated by the balance of specificity and fidelity across its target site, relative to host constraints of the same sequence

Consideration of the I-AniI homing endonuclease and its most well-studied cousins from the LAGLI-DADG family, particularly similar studies on I-CsmI from the same host gene,⁴⁰ illustrates that the most successful homing endonucleases display several evolutionary mechanisms for optimizing their recognition of and persistence in genomic host sites:

(1) Such elements have arrived, by random sampling, at substrate specificities that correspond to highly conserved, and therefore static, sequences in potential host genes. This observation has been made previously for mobile introns specific both for structural RNA genes (e.g., Refs. 30,45) and for protein-encoding genes (e.g., Ref. 46). In this latter study, the homing endonuclease I-TevI (a member of the much less specific GIY-YIG endonuclease family, which is generally found in phage genomes) was shown to bind a DNA target site in its phage thymidylate synthase host gene that corresponds to functionally critical active-site residues in the host enzyme, including one critical base common to all known TS gene sequences.

If a host gene is essential or valuable to the host organism, then purifying selection pressure across the gene is likely to be higher and therefore more likely to produce 'static' sequence targets for invasion. Furthermore, once an endonuclease and its element (intron or intein) have become established, only a precise deletion of the element from the host gene would reestablish an intronless or an inteinless allele for reinvasion. Any 'sloppy' deletion event would likely create a mutation within a critical region, rendering the gene product nonfunctional. Thus, endonucleases select against deletion by targeting essential genes, especially if the target site corresponds to an active site or to other functionally critical regions of the protein.

(2) Homing endonucleases optimize an appropriate balance of relative specificities at individual base pairs of the target site that corresponds to the underlying constraints imposed by the host gene sequence (for additional experimentally validated examples, including a similar study of the I-CsmI homing endonuclease from the mitochondrial cob gene in *C. smithii*,⁴⁰ see Refs. 16,18,47–49). In the case of I-CsmI, the endonuclease is shown to display a "slight but significant tendency...to cleave substrates containing a silent or tolerated amino acid change more efficiently than nonsilent or nontolerated ones."⁴⁰

In the case of host genes encoding structural tRNA and ribosomal RNA molecules, constraints imposed by the gene product itself would include conservation of base pairs and longer-range tertiary interactions required for function in folded RNA

molecules. These constraints can be quite strong, as only four unique bases are encoded within these nucleic acids. Therefore, many highly structured RNA molecules tend to be extremely well conserved across biological kingdoms and therefore somewhat static targets for invasion by homing endonucleaseintron systems. Additionally, the preponderance of base-paired structural elements in RNA structures increases the amount of palindromic symmetry in many potential target sites, encouraging the success and persistence of homodimeric homing endonucleases. These host genes can thus serve as a reservoir for such proteins, which can then undergo gene duplication and fusion into monomeric endonucleases, greatly expanding their available target site repertoire.

In contrast, host genes that encode proteins appear to be more challenging targets for invasion, for at least three reasons: (i) the combination and interactions of the 20 possible amino acids within unique protein fold families are more readily and rapidly diverged during evolution than are the interactions of nucleic acids; (ii) the genetic code for amino acid sequences is partially degenerate, allowing synonymous (silent) mutations that can inhibit homing endonuclease binding while maintaining the structure and function of the host protein; and (iii) protein-coding sequences are inherently less symmetric than are those that code for ribosomal RNA and tRNA.

(3) Many persistent homing endonucleases also acquire secondary activities that provide an advantage to the host, such as assisting in intron splicing, thus generating selective pressure to help maintain endonuclease fold and stability.^{37,50,51} In the case of I-AniI, the homing endonuclease has also evolved to become an essential cofactor for the splicing of its cognate group I intron. This 'maturase' activity is required for successful expression of the cytochrome b oxidase gene.^{52,53} In such a case, one would expect that the loss of endonuclease activity due to random mutations would be reduced because additional purifying selection pressure will act at many positions on the endonuclease reading frame to preserve its folding and stability, in order to maintain the function of the host gene. While a small number of mutations in I-AniI can upcouple its homing endonuclease and maturase activities,^{26,35,54} the number of random mutations in the LHE gene that can abrogate homing activity without reducing the fitness of the host (by decreasing the expression of the *cob* gene) is far less than that for a comparable homing endonuclease with no additional function. The moonlighting relationship of the two activities can thereby 'stabilize' the endonuclease reading frame in the host genome.

The cost to the endonuclease of developing this secondary 'moonlighting' activity as a mechanism to help maintain its primary function as a DNA endonuclease is that each activity imposes different, partially overlapping constraints on the identity of individual amino acids in the LHE scaffold. While some of those sequence constraints are primarily structural and thus help to preserve both activities, others are distinct to the two separate functions of the protein and are potentially noncomplementary. Thus, the protein may have difficulty in optimizing either or both activities during evolution, leading to a mobile element that is relatively stable in its particular biological host, but at the cost of reduced efficiency for horizontal transfer.

Concluding remarks: several properties of wild-type homing endonucleases should be optimized for gene targeting

LHEs are under intense study by several groups as possible reagents for gene-targeting applications.^{55,56} The experiments reported here, as well as a variety of published studies from several groups,⁵⁵ clearly indicate that (i) the optimal DNA substrate of an LHE does not necessarily coincide with its physiological target in its cognate biological host and (ii) most, if not all, homing endonucleases, while inherently very specific due to their long target sites, are capable of even greater specificity because they have evolved to only being specific enough to avoid host toxicity.

This study demonstrates that the I-AniI endonuclease and presumably its cousins throughout the LAGLIDADG family display more than sufficient specificity to be potential reagents for gene-specific applications. However, wild-type homing endonucleases often display a variety of biophysical characteristics that are suboptimal for use in biotechnological or therapeutic applications, as a result of the lack of host pressure to maintain their homing function. This can include instability and/or insolubility upon expression and nonoptimal DNA affinity for various purposes. As well, the presence of additional biochemical activities on the same protein scaffold (such as protein splicing for inteinassociated homing endonucleases, or protein and RNA binding for various maturase-homing endonuclease proteins) can impose constraints on their structure that interfere with optimal DNA cleavage activity. In the case of I-AniI, maturase activity may be responsible for the conservation of surface residues on the C-terminal domain that reduce the protein's solubility and also impart RNA-binding function that may interfere with its DNA endonuclease activity. Even with these caveats, wild-type homing endonucleases have been used in many studies to drive gene conversion events, with little or no attendant issues of cell toxicity or elevated nonspecific DNA cleavage,^{44,55} and appear promising enough as reagents for targeted genetic applications to be well worth the effort required to optimize their stability, specificity, and biochemical behavior. As a case in point, during the experiments described in this study and in an earlier study on I-AniI, we were able to easily (i) eliminate RNA intron association as competing binding activity, via a single-point mutation²⁶; (ii) improve the solubility and solution behavior of the purified protein, via two mutations on its surface distant from the DNA-

binding surface; and (iii) identify a substrate variant of the endonuclease target site that is bound and cleaved significantly more efficiently than wild-type targets.

Materials and Methods

Materials

The I-AniI homing endonuclease was expressed and purified from *Escherichia coli* strain BL21 (DE3)lysS as previously described, ²⁶ with the exception that expression was induced at 37 °C for 2–3 h after the culture had achieved early log growth phase (OD₆₀₀ \sim 0.6). Wild-type I-AniI was used for substrate-selection experiments. For binding assays, two point mutations (F80K and L232K) that improve the solubility of the enzyme and facilitate measurements using isothermal titration calorimetry (ITC) were introduced. Both mutations are far removed from the DNA-binding surface of the enzyme and are on exposed surface loops (Fig. S1).

DNA oligonucleotides for the generation of the randomized substrate library and for subcloning and amplification steps were purchased from Operon, Inc. (50-nmol scale, salt-free). DNA oligonucleotides for cocrystallization studies and for ITC were ordered from IDT, Inc. (1-µmol scale; HPLC and salt-free purifications, respectively).

In vitro site specificity screen

A plasmid library containing randomized target site variants for the I-AniI endonuclease was constructed using a straightforward protocol described here and shown schematically in Fig. 3a. Additional details of the protocol for library construction are provided in Supplementary Data. A single-'template' oligonucleotide that contains, in order, a 5' EcoRI restriction site for subcloning, the randomized target site (mixed at each position in the target site at a ratio of 60% wild-type base to 13.3% of the three possible substitutions), a 3' XhoI restriction site for subcloning, and a nondegenerate primer binding site for generation of the corresponding randomized doublestranded DNA was designed and synthesized. After annealing a universal primer to the 3' end of the template, a double-stranded duplex was generated by polymerase extension using Klenow fragment. The site library was digested to yield cohesive ends for subcloning, gelpurified, and ligated into linearized BlueScript plasmid. To measure the complexity and quality of the library (and, subsequently, the progression and convergence of the selection method), randomly selected members of the library were sequenced, and the complexity and average mutational load of the naive library were calculated at each base (Fig. 3b). A complexity of 10⁶ unique sites, containing an average of 6.7 mutations in each unique target site variant, was generated.

Base-pair positions that immediately flank the I-AniI target site were also randomized and thus served as unique sequence 'labels.' This allowed us to distinguish, for any sites that were recovered more than once, those that arose independently from separate parental clones *versus* those derived from single parental clones as a result of a 'bottleneck' in the cloning or screening process.

The naive library was digested for 2 h at 37 $^{\circ}$ C with a purified enzyme under optimal conditions: 10 nM DNA substrates and 10 nM enzyme (a 1:1 ratio is required

because LAGLIDADG endonucleases do not readily release free product ends), 50 mM Tris-Cl (pH 7.5), 50 mM NaCl, 10 mM MgCl₂, and 1 mM DTT. This protocol provides for optimal cleavage of DNA targets by the endonuclease and therefore yields the most conservative estimate of its specificity. Typically, approximately 1% of the naive library is prone to cleavage in the initial round of digests. Linearized products were purified, religated, and passed on to the next round of selection, and cleavable sites were sequenced at each cycle to monitor progression of the selection. Cycles of site digests and recovery were continued until convergence (which was observed when the number of mutations per site and the percentage of unique sites had both reached a minimum). Progression of the screen and of the enrichment of cleavable DNA sequences is illustrated in Fig. 3b.

From the final pool of recovered DNA substrates, 428 cleavable target sites were sequenced in order to align sufficient unique target sequences to allow accurate calculation of IC and base-pair covariation within the site. All sequences were determined in a high-throughput 96-well-plate sequencing format using the HLA genotyping facility at the Hutchinson Center.

Using an alignment of all cleavable target sites identified in this screen, the specificity of DNA recognition by I-AniI was calculated at each base-pair position of the target site using the method of Schneider et al.41 This method generates a descriptor of specificity at each DNA base pair (termed 'information content,' quantitated as 'bits' of information) by summing the probability of each possible base found at each position across the site, relative to the background base content expected at each position based on library design. This form of analysis can be directed against collections of either naturally occurring sites that are recognized by a single protein or sites that are recovered from a library screen as described in this study. IC at individual base pairs can range from 0 bits, corresponding to complete degeneracy (25% probability of any base at a position), to 2 bits, corresponding to no degeneracy (100% chance of a unique base at that position).

Discrete cleavage assays of base-pair variants at the 'central-four' base-pair positions

In addition to the screen from a randomized site library described above, all individual single base-pair mismatches were generated across the central-four positions of the I-AniI target site and tested individually for relative cleavage activity. This was done because these four positions, which are located in between two scissile phosphates, are within a region of significant proteininduced DNA bending and are not in direct contact with the enzyme. Therefore, recognition of these bases is largely driven by indirect readout of sequence-specific DNA conformational preferences and is prone to particularly strong patterns of high or low specificity and fidelity that are independent of protein contacts.

The target site variants containing individual base-pair mismatches across these four positions were generated using the same templated polymerase extension protocol described above, except that four separate mutant oligonucleotide templates were used (each randomized at only one of the central-four positions of the target site). These templates were combined in a single reaction mixture, annealed with a common universal primer, converted to individual double-stranded duplexes, and cloned in a single ligation-and-transformation step into BlueScript. In this case, each product construct contains a single-point substitution within the central-four bases of the target site. Individual constructs were isolated, prepped, and sequenced, and a complete matrix of all 12-point substitutions across the central-four bases was generated. All substrate concentrations were normalized and aliquotted for individual digests, and each target site variant was individually tested in parallel cleavage assays.

Sequence alignment and IC of the cob host gene

For the *A.n.* cob protein sequence, a set of homologous sequences was collected using an iterative PSI-BLAST (Position-specific iterative BLAST) search of the nonredundant protein sequence database. Using the output, a multiple-sequence alignment consisting of the 500 most homologous *cob* sequences was created, corresponding to protein sequence identities ranging from 55% to 95% and to *E*-values ranging from 10^{-105} to 10^{-164} . The degree of conservation of each protein residue in the host gene was observed by calculating IC, as described previously.⁵⁷ For simplicity, IC values were linearly normalized from 0 to 1, with 1 corresponding to the most conserved positions.

Cleavage assays

Cleavage assay reactions were carried out using target sites in supercoiled plasmid substrates. The I-AniI cleavage buffer and digest condition were both identical with that used for the site library screen: 2 h at 37 °C with 10 nM DNA substrates and 100 nM enzyme in 50 mM Tris–Cl (pH 7.5), 50 mM NaCl, 10 mM MgCl₂, and 1 mM DTT. Reactions were terminated by the addition of stop buffer (2% SDS, 100 mM ethylenediaminetetraacetic acid (EDTA), 20% glycerol, and 0.2% bromophenol blue). Samples were run on a 0.8% agarose gel, stained with ethidium bromide, and imaged with Eagle Eye II and EagleSight, version 3.22 (both from Stratagene, La Jolla, CA). Quantification was performed with the program ImageJ 1.32j.

Isothermal titration calorimetry

Two sets of DNA oligonucleotides containing either the wild-type I-Ani target site (5'-CCTTCCCTGAGGAGGT-TTCTCTGTAAACCCTTCC-3' and its complement; the target site is in bold) or the 'hypercleavable' Lib4 target identified during target site screening experiments (5'-CCTTCCCTGAGGAGGTTACTCTGTTAACCCTTCC-3' and its complement; base-pair substitutions relative to wild-type are underlined) were purchased from Operon Biotechnologies. After suspending in 1 mM STE buffer (10mM Tris-Cl pH7.5, 10mM NaCl, 1mM EDTA), complementary strands were combined in a 1:1 ratio and annealed by incubating at 95 °C for 10 min and by slowly cooling to room temperature to form DNA target sites for ITC-binding analysis. G-C anchors at each oligonucleotide end promote stable heterodimer formation and prevent hairpin formation of the pseudopalindromic site.

Protein and DNA samples were dialyzed overnight into Ani ITC buffer (50 mM Tris (pH 7.5), 50–100 mM NaCl, and 10 mM CaCl₂) at 4 °C. Dialyzed samples were quantified by UV spectroscopy, using extinction coefficients to calculate concentrations. Protein concentrations were further confirmed by Bradford assay, as well as by estimation on SDS-PAGE gels against commercial protein standards. Sample concentrations ranged from 4 to $14 \,\mu M$ I-AniI and from 22 to 50 μM double-stranded DNA target.

A VP-ITC MicroCalorimeter was used for all experiments, with the protein sample in the cell and with the DNA target in the autopipette. Individual runs consisted of 30–40 injections of 5–10 μ L each, depending on sample concentrations, and were conducted at 30 °C with a 351-rpm stirring speed. A full experiment consisted of three experimental runs injecting DNA into the protein-containing cell, as well as a control run injecting DNA into the buffer. Data were fit using Origin 7 SR2 software; both sets of data were well modeled by the one-site fitting algorithm in the Origin 7 SR2 software, and K_d values were calculated from the resulting fit parameters.

X-ray crystallography

The resolution of the I-AniI/DNA cocrystal structure was extended to 2.4 Å after screening crystals for improved diffraction behavior. Complementary DNA strands were annealed by incubating for 5 min at 90 °C and then allowed to cool to room temperature. The protein was diluted to a concentration of 5 mg/mL, mixed with a 1.2-M excess of DNA duplex at room temperature, and used directly for cocrystallization. Crystals containing wild-type DNA target site were grown at 22 °C by vapor diffusion. A 500-µL reservoir containing 100 mM KCl, 20 mM MgCl₂, 50 mM sodium citrate (pH 5.0), and 16-22% PEG 3350 was equilibrated against a 2-µL drop containing a 1:1 mixture of the protein/DNA complex and the reservoir solution. Crystals grew as plates within 2 weeks, were flash-frozen in the mother liquor described above, and were augmented with 30% PEG 3350 and 20% glycerol v/v. The crystals are space group $P2_1$ and diffract to a resolution of approximately 2.4 Å.

The structure of the I-AniI/DNA complex was solved by molecular replacement using CNS (Crystallography and NMR System),⁵⁸ with the lower-resolution I-AniI/ DNA complex structure as an initial search model. The

 Table 1. Crystallographic data and refinement statistics for wild-type I-AniI/DNA

| Data collection | | |
|---|---------------------|--|
| Space group | $P2_1$ | |
| Resolution (highest shell) | 50-2.4 (2.47-2.4) | |
| Unit cell dimensions (Å) | a = 60.4, b = 72.7, | |
| | c=61.1 | |
| | $\beta = 103.9$ | |
| Wavelength | 1.000 | |
| Total reflections | 48,322 | |
| Unique reflections | 14,643 | |
| Completeness (%) | 92.1 (76.75) | |
| Redundancy | 3.3 (2.1) | |
| R _{merge} | 0.035 (0.140) | |
| Average I/σI | 18.5 (5.7) | |
| Refinement | | |
| R _{work} | 0.219 (0.229) | |
| R _{free} | 0.276 (0.320) | |
| R.m.s.d. bond length (Å) | 0.001 | |
| R.m.s.d. bond angles (°) | 1.25 | |
| Protein residues | 275 | |
| Nucleotides | 60 | |
| Metal ions | 3 | |
| Water molecules | 47 | |
| Ramachandran distribution | 92.3, 5.4, 1.6, 0.7 | |
| (% core, allowed, generous, disallowed) | | |
| Average <i>B</i> -factors (\dot{A}^2) | 29.9 | |
| | | |

structure was modeled in COOT (Crystallography Object-Oriented Toolkit)⁵⁹ and refined to 2.4 Å using REFMAC, with 5% of the data set aside for cross-validation. The final refinement statistics are $R_{\rm work}/R_{\rm free}$ =0.216/0.276. Geometric analysis of the structures using PROCHECK⁶⁰ indicates that over 90% of the residues are in the most favored region. Refinement statistics are provided in Table 1.

Protein Data Bank accession number

The coordinates of the I-AniI/WT DNA complex have been deposited in the RCSB Protein Data Bank with accession number 2QOJ.

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Supplementary Data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j. jmb.2007.07.052

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