Structure of *Aquifex aeolicus* Argonaute Highlights Conformational Flexibility of the PAZ Domain as a Potential Regulator of RNA-induced Silencing Complex Function*†§*

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Gene silencing mediated by RNA interference requires the sequence-specific recognition of target mRNA by the endonuclease Argonaute, the primary enzymatic component of the RNA-induced silencing complex. We report the crystal structure of *Aquifex aeolicus* Argonaute, refined at 3.2 Å resolution. Relative to recent Argonaute structures, a 24° reorientation of the PAZ domain in our structure opens a basic cleft between the N-terminal and PAZ domains, exposing the guide strand binding pocket of PAZ. This rearrangement leads to a branched, Y-shaped system of grooves that extends through the molecule and merges in a central channel containing the catalytic residues. A 5.5-ns molecular dynamics simulation of Argonaute monitors DNA binding, conformational rearrangement of the PAZ domain in our structure opens a basic cleft between the N-terminal and PAZ domains, exposing the guide strand binding pocket of PAZ. This rearrangement leads to a branched, Y-shaped system of grooves that extends through the molecule and merges in a central channel containing the catalytic residues. A 5.5-ns molecular dynamics simulation of Argonaute shows a strong tendency of the PAZ and N-terminal domains to be mobile. Binding of single-stranded DNA to Argonaute monitored by total internal reflection fluorescence spectroscopy shows biphasic kinetics, also indicative of domain rearrangement upon DNA binding. Conformational rearrangement of the PAZ domain may therefore be critical for the catalytic cycle of Argonaute and the RNA-induced silencing complex.

RNA interference, an ancient mechanism of gene silencing mediated by small RNAs, is widely distributed throughout most eukaryotes (1, 2). Potent silencing of genes is achieved through the processing of double-stranded RNA (dsRNA) into small interfering RNAs (siRNA) by the enzyme Dicer. The ~22-nt-long siRNAs contain a signature 5′-phosphate group and a 2-nt-long 3′-overhang (3). The siRNA is then loaded into the RNA-induced silencing complex (RISC), of which Argonaute is the primary catalytic component (4, 5). Energetic asymmetry of the siRNA ends allows for its directional loading into RISC (6–8). Argonaute cleaves the passenger strand of the siRNA, leaving the guide strand of the siRNA bound to RISC (9–11). This single-stranded guide strand siRNA bound to Argonaute is required for the sequence-dependent recognition of mRNA by RISC. Recombinantly expressed human Argonaute 2 in complex with single-stranded siRNA forms a minimal RISC, capable of mRNA recognition and cleavage (12). Recent crystal structures of Argonaute apoenzymes show the architecture of Argonaute to be a multidomain protein composed of N-terminal, PAZ, MID, and PIWI domains (13, 14).

The catalytic cycle of RISC requires the binding of a number of different nucleotide structures to Argonaute, and we expect Argonaute to undergo a number of conformational changes during the cycle of mRNA recognition by RISC (15, 16). Nevertheless, it remains unclear how the multidomain arrangement of Argonaute recognizes and distinguishes between single-stranded and double-stranded oligonucleotides that correspond to the Dicer-processed siRNA product, guide strand siRNA, and the guide strand/mRNA duplex.

We report the crystal structure of *Aquifex aeolicus* Argonaute (AaAgo), crystallized in the presence of a single-stranded siRNA and refined using data to 3.2 Å resolution. We propose a model for Argonaute activity in the context of the RISC complex. We also report a kinetic and 5.5-ns molecular dynamics characterization of Argonaute, which highlights biphasic binding kinetics with ssDNA, and a tendency for pronounced motion of the PAZ and N-terminal domains. Together, these studies highlight a functional role for domain flexibility, especially of the PAZ, in Argonaute activity that may be crucial for the function of the RISC.

**EXPERIMENTAL PROCEDURES**

Protein Expression and Purification—The full-length Argonaute gene from *Aquifex aeolicus* was amplified by PCR from genomic DNA and cloned into a modified pET16b vector (Novagen) containing an N-terminal His tag. AaAgo was overexpressed in *Escherichia coli* host cell strain BL21 (DE3) pLysS (Strategene) and induced by 1 mM isopropyl-1-thio-β-D-galactopyranoside (A₄₅₀ ~ 0.8) at 37 °C for 3 h. Cells were resuspended and sonicated in 10 mM Hepes, 300 mM NaCl, pH 7.4. Non-thermostable proteins were removed by a heating step of 72 °C for 30 min. Precipitated proteins were removed by centrifugation at 20,000 rpm, and the supernatant was applied to an immobilized metal affinity column (nickel-nitrilotriacetic acid-agarose; Qiagen), with the partially purified target protein eluting at 300 mM imidazole. The N-terminal His tag was

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The atomic coordinates and structure factors (code 2NUB) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http://www.rcsb.org/).

†The on-line version of this article (available at http://www.jbc.org/) contains supplemental Figs. S1 and S2 and supplemental data.

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2The abbreviations used are: dsRNA, double-stranded RNA; ssRNA, single-stranded RNA; RISC, RNA-induced silencing complex; AaAgo, *Aquifex aeolicus* Argonaute; ssDNA, single-stranded DNA.
removed by addition of TEV protease overnight at 4 °C. The target was further purified by gel filtration on a Superdex 200 column (Amersham Biosciences), equilibrated with 10 mM Hepes, 300 mM NaCl, pH 7.4. For binding experiments, the protein was purified by affinity chromatography and gel filtration in 20 mM Heps, 150 mM KCl, pH 7.4.

Crystallization, Data Collection, and Refinement—siRNA nucleotides (5′-PO₄GGG AAU GCG ACU CAU GAA GCU-3′) were purchased from Ambion and solubilized in 10 mM Hepes, pH 7.4. An equimolar ratio of protein at 10 mg/ml and single-stranded siRNA was mixed together and incubated at 37 °C for 15 min prior to crystallization trials. Crystals were grown by the hanging drop method at 22 °C, with 3.0 M sodium formate, 100 mM Hepes, pH 7.4, as the precipitant. Crystals grew within 2 weeks to a maximum size of ~200 × 200 × 30 μm. For cryoprotection, crystals were transferred into mother liquor containing 25% glycerol (v/v) and immediately frozen in liquid nitrogen. Data were collected at beamline ID23–1 at European Synchrotron Radiation Facility in 1° oscillations. Diffraction was observed to a useful resolution of 3.2 Å and indexed in tetragonal space group I422, with one molecule/asymmetric unit. Data were processed using MOSFLM and SCALA from the CCP4 suite (17). Although the data were weak, with reflections at an I/σ(I) of 2 at 3.6 Å resolution, a plot of the SigmaA versus resolution (supplemental Fig. S1) showed that the weak data were still usable, and thus the refinement utilized all data to 3.2 Å resolution (18, 19).

The structure was solved by molecular replacement, using 1YVU as a search model. Although an unambiguous solution was initially obtained by the CCP4 program MOLREP (17), the density for the PAZ domain was poor. A second run using 1YVU excluding the PAZ domain was done in MOLREP, while placement of the PAZ domain was facilitated by EPMR (20). The model was refined by a combination of rigid body, positional, simulated annealing, and grouped and individual B-factor refinement in CNS (21) with manual rebuilding in Coot (22). B-factor refinement, though it potentially introduces a significantly higher number of parameters, is restrained in CNS and is recommended by the authors of the program, even at moderate resolution (18). Tests of grouped and individual B-factor refinement favored individual B-factor refinement in the early stages and grouped B-factor refinement in the later stages. Data collection and refinement statistics are summarized in Table 1. The choice of refinement protocols, especially in regard to B-factor refinement, is described in greater detail in the supplemental data.

Molecular Dynamics—The final crystallographic model was used for molecular dynamics simulations. First, loops of missing residues in the model were built using MODELLER (23). A 5.5-ns molecular dynamics simulation was then performed using the AMBER 8.0 (24) suite of programs together with the parm99 force field and a modification for backbone torsion angles (25). The starting structure was placed into an orthorhombic periodic box of TIP3P water molecules (26) such that the distance between the edges of the water box and the closest protein atom was at least 10 Å. The particle mesh Ewald method (27) was used to treat long-range electrostatic interactions with a direct space non-bonded cutoff of 9 Å. Bond lengths involving bonds to hydrogen atoms were constrained using SHAKE (28). The time step for the simulation was 2 fs. After minimization, the system was gradually heated to 300 K over a period of 50 ps, after which the solvent density was adjusted during another 50 ps. Harmonic restraints applied during these steps to all solute atoms were then reduced over 50 ps. After an additional 350 ps of simulation at 300 K, the following 5.0 ns were used for analysis, considering snapshots obtained every 1 ps.

Structural changes during the simulation were analyzed in terms of root mean square deviations of Cα atoms from the starting structure and variations in the Cα-Cα distances between particular residues at the top of the M groove. To analyze dynamic properties of the molecule, equal time cross-correlations of atomic fluctuations C(ij) between atoms i and j are obtained (29). For completely correlated motions, C(ij) = 1, and for completely anticorrelated motions, C(ij) = −1. Finally, a principal component analysis of Cα atom motions in Cartesian space was performed, yielding collective variables (also termed “modes”) that describe the essential dynamics of the system (30). The modes most involved in the observed conformational change Δr between 1YVU and the structure determined here were then obtained by calculating the overlap between each mode and Δr according to Ref. 31. An overlap of one indicates that the directions of both kinds of collective Cα displacements are identical. In all cases, only those Cα atoms not contained in the N-terminal or PAZ domain were used for least-squares fitting to avoid the introduction of spurious correlations (32, 33).

Binding Kinetics—Binding of fluorescently labeled DNA to immobilized AaAgo was detected in real time by total internal reflection fluorescence spectroscopy. This technique was used in combination with reflectance interference for mass-sensitive quantification of the amount of immobilized AaAgo (34). Site-specific immobilization of the protein onto the sensor chip through its His tag was achieved by using multivalent chelator heads coupled to a polyethylene glycol polymer brush, which were loaded with Ni²⁺ ions (35).

100 nM of 21-nt-long ssDNA (5′-CGT ACG CCG AAT ACT TCG ATT-3′), 15-nt-long ssDNA (5′-CGT ACG CCG AAT

### Table 1

Data collection and refinement statistics

| Source | ESRF ID23–1 |
|--------|-------------|
| Wavelength (Å) | 0.9795 |
| Resolution (Å)³ | 3.2 |
| Space group | I422 |
| Unit cell (Å) | a = 125.0, c = 253.5 |
| Rmerge | 0.135 (0.797) |
| Redundancy independent Rmerge &sup3; | 0.145 (0.852) |
| Completeness (%) | 100 (100) |
| Measured reflections | 134396 (19238) |
| Independent reflections | 16989 (2443) |
| Redundancy³ | 7.9 (7.9) |
| (I/σ(I))³ | 4.5 (1.0) |
| Overall (Å)³ | 65 |
| PAZ domain | 101 |
| N-terminal / PIWI / Mid | 61 |
| Rmerge / Rsym (%) | 31.0 / 23.2 |
| Root mean square deviation bond angles (°) | 1.47 |
| Root mean square deviation bond lengths (Å) | 0.009 |
| Ramachandran allowed / generous / disallowed (%) | 97.0 / 2.0 / 1.0 |

³ The threshold for an I/σ(I) of 2 occurs at 3.6 Å resolution.

² Values in parentheses refer to values in the highest resolution shell, 3.37-3.2 Å.
ACT-3), and 9-nt-long (5′-CGT ACG CGG-3′), each 5′-phosphorylated and 3′-labeled with Oregon Green 488, were used for all binding experiments. Purified His-tagged AaAgo in 20 mM Hepes, 150 mM KCl, pH 7.4, was immobilized onto the sensor chip at various concentrations between 1 and 50 nM. 10 mM MgCl₂ was included in the buffer. The fluorescent label was excited with an argon ion laser at 488 nm, at ~100 μW. All experiments were performed at 25 °C with a DNA injection of 36 s, followed by a 200-s wash period. Immobilized protein was washed off the sensor surface by addition of 500 mM imidazole. Kinetic constants were calculated using BIAevaluation 3.1 software.

RESULTS

Structure of AaAgo—AaAgo is a roughly elliptical molecule measuring ~55 × 70 × 90 Å in size. Nearly all of the amino acids are defined in the electron density, with the exception of residues 176–177, 245–247, and 262–271. The domains of AaAgo are divided into three lobes, comprising the N-terminal, residues 176–177, 245–247, and 262–271. The domains of PAZ-RNA complexes show the nucleotide end sandwiched between strand siRNA. X-ray and NMR structures of PAZ-RNA complexes suggest the nucleotide end of the oligonucleotide is hydrophobic and is mediated by conserved aromatic residues. In AaAgo, these residues are Trp-226 and Phe-209, within the α-helical region of the PAZ domain (Fig. 1a, green). Residues 184 and 601 contribute to the PAZ domain. The PIWI domain has an RNase H fold, similar to other endonucleases such as transposases and viral integrases (38, 39) and as noted in structures of other Argonaute species and PIWI proteins. Cleavage of the target mRNA strand likely proceeds through a similar mechanism as other endonucleases, mediated by divalent cations such as Mg²⁺ or Mn²⁺ (12).

RNA Binding Sites—The architecture of this three-lobed structure results in a Y-shaped system of positively charged grooves that runs throughout the protein (Fig. 1b). The “stem” of the structure is flanked by the PAZ and MID/PIWI domains and contains the catalytic amino acids. Because of the location of the active site and the positively charged amino acids lining the stem, the dsRNA intermediate, composed of the guide strand siRNA and the target mRNA, likely binds here (13). Crystal structures of PIWI bound to dsRNA strongly support this model (37, 40). In our structure, the groove stem measures ~17–19 Å in diameter. Notably, the side chains of Arg-184 and Arg-601 approach within 4.5 Å of one another, thus requiring a conformational change or side chain rearrangement to properly accommodate the dsRNA substrate. The PAZ lies in an orientation that encloses part of the groove, forming a short tunnel.

During the course of this work, a crystal structure of AaAgo was reported (14), crystallized in a different space group from ours (henceforth referred to as 1YVU). The differences between the structures are most pronounced in the relative orientations of the PAZ domain. This movement arises from a gradual shift of the L1 β-sheet connecting the N-terminal and PAZ domains. Analysis of the conformational changes between our structure and 1YVU using DynDom (42) shows a ~24° shift in the position of the PAZ in our structure, away from the N-terminal domain (Fig. 1c). Consequently, this domain shift in our structure exposes the binding site for the 3′-end of the guide strand siRNA. The orientation of the PAZ domain observed in our
structure resembles the conformation of the *Pyrococcus furiosus* Argonaute apoenzyme, which also adopts a three-lobed structure with prominent grooves between the N-terminal domain and the PAZ and PIWI domains (13). The relatively high average B-factors of the PAZ domain, 101 Å² compared with 61 Å² for the remainder of the molecule, further support a tendency of the domain to be flexible. As the B-factors in the PAZ domain are relatively high, with an average amplitude of thermal vibration \( u = 1.13 \) Å (\( B = 8 \pi^2 <u^2> \)), we calculated a composite omit map to investigate how well the structure of the PAZ domain is defined by the density map. The map was created by omitting a fixed 5% volume of the model in the asymmetric unit and subjecting the remainder to simulated annealing refinement. The reassembled map shows clear density for the PAZ domain (supplemental Fig. S2), demonstrating that the position of the PAZ domain is well defined in our structure. We also note that in the crystal the PAZ domain is relatively unhindered by crystal contacts, which may be the source of the higher B-factors. In contrast, the PAZ domain of 1YVU is involved in extensive crystal contacts with the N-terminal, MID, and PIWI domains. As the conformational differences in the two AaAgo structures alternatively expose or occlude the entrance to the nucleotide binding site in PAZ, this implies a functional flexibility of the PAZ domain in RISC function. An analysis of the surface accessibility of the two AaAgo structures shows that Lys-37 and Trp-40 in the \( \alpha 1 \) of the N-terminal domain are exposed in our structure relative to 1YVU (91.1 and 110.6 Å² in our structure versus 68.7 and 64.6 Å², respectively, in 1YVU). Likewise, residues Glu-215, Arg-216, and Thr-218, lining the entrance to the nucleotide binding pocket in PAZ, are also comparatively exposed in our structure (131.9, 152.9, and 115.7 Å² in our structure versus 109.7, 59.9, and 9.8 Å², respectively, in 1YVU).

One branch of the Y-shaped groove is lined by the N-terminal and PIWI domains (Fig. 2a, S groove) and measures ~17–19 Å in diameter. The size of the groove is very similar in 1YVU as well as *P. furiosus* Argonaute. The S groove is lined with basic amino acids, which may serve to recognize the backbone of dsRNA. The stretch of sequence corresponding to the PIWI box, residues 615–657, a highly conserved element present in eukaryotic Argonaute proteins, lies at one end of the S groove. Although the corresponding amino acids of the PIWI box in human Ago2 (residues 730–772) are not conserved, the region is solvent-exposed and is structurally localized at the base of the groove defined by the PIWI and N-terminal domains. A direct interaction between Dicer and Argonaute was observed, localized to an interaction between the PIWI box and the second RNase III domain of Dicer (43, 44), while subsequent studies have shown that Argonaute is responsible for the cleavage of the sense strand of Dicer-processed siRNA (9, 10). As such, these results constrain the location of Dicer within the RISC complex and suggest that the S groove may act as a channel for...
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Helices α1 and α2 act as a putative RNA recognition face of the N-terminal domain, spanning both the M and S grooves (Fig. 2c). Although AaAgo was crystallized in the presence of a single-stranded siRNA, only disordered, unassignable electron density was observed in the 5′-nucleotide binding pocket, with no discernible density observed in the 3′-nucleotide binding site in PAZ. This may be explained by the preference of AaAgo for ssDNA ($K_d \sim 0.01 \mu M$) relative to ssRNA ($K_d \sim 0.97 \mu M$) as measured by filter binding assays (14).

Molecular Dynamics—To further investigate the potential flexibility of the PAZ domain within Argonaute, we performed a 5.5-ns molecular dynamics simulation. Visual inspection of snapshots along the trajectory reveals a tendency for the PAZ domain, and to a lesser extent, the N-terminal domain, to change conformations, leading to a widening and narrowing of the M groove. This is corroborated by a time series of root mean square deviation of C$_\alpha$ atoms from the experimental structure (Fig. 3a). Whereas the MID/PIWI domains only show moderate deviations that remain below 2.5 Å, the N-terminal and PAZ domains show major conformational changes of up to 4.3 and 5.5 Å, respectively. Importantly, there was a tendency for both the PAZ and N-terminal domains to swing back and forth during the course of the simulation, with two and a half cycles observed for the PAZ and at least four cycles for the N-terminal domain (Fig. 3b).

The motions lead to considerable changes of the distances between both domains, as depicted in Fig. 3b for those three pairs of residues at the “tips” that show closest interactions in our structure (Thr-218-Ser-73 or 1YVU (Arg-216-Glu-76) or Thr-218-Trp-40), respectively. Changes in the C$_\alpha$-C$_\alpha$ distances of up to 10 Å can be observed, which relate to changes in the diameter of the M groove of more than 5 Å.

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Orientational correlations between motions of C$_\alpha$ atoms are displayed color-coded in Fig. 3c. The map shows strongly correlated motions within the PAZ and N-terminal domains, indicative of a rigid body type of motion of the domains.

Finally, a principal component analysis of the motions of the C$_\alpha$ atoms was computed yielding collective variables (or modes) for the description of the essential molecular dynamics

passage of the Dicer-cleaved siRNA product into the active site of Argonaute.

The second branch of the Y-shaped groove is bordered by the N-terminal and PAZ domains (Fig. 2b, M groove), and measures ~17 Å in diameter. The closest interactions between the N-terminal and PAZ domains in our structure occur between the side chains of residues Ser-73 and Thr-218 (7.3 Å), which lie at the top of the groove. Importantly, in our structure, the 3′-nucleotide binding pocket in PAZ is exposed to the M groove. In contrast, 1YVU shows a closed conformation, characterized by a salt bridge between the side chains of residues Arg-216 and Glu-76 (2.9 Å), as well as a hydrogen bond between the indole nitrogen of Trp-40 and the carbonyl oxygen of Thr-218 (3.2 Å). Our structure shows no interactions between these residues, with a Arg-216/Glu-76 distance of 17.8 Å and a Thr-218 distance of 10.2 Å. Notably, although the size of the M groove shows significant variation between our structure and 1YVU, the size of the S groove is the same.

A sequence alignment of residues lining the S groove between AaAgo and human Ago1 and Ago2 reveals a number of conserved residues, including Lys-34, Arg-52, Arg-95, from the N-terminal domain, and Arg-573, Arg-576, Lys-642, and Arg-644 from the PIWI domain, which constitute the majority of the basic amino acids in the S groove (Fig. 2c).

FIGURE 3. Periodic motions of the PAZ and N-terminal domains. a, time series of root mean square deviations of C$_\alpha$ atoms of the MID/PIWI domains (red), the N-terminal domain (green), and the PAZ domain (blue) from the starting structure over 5.0 ns of molecular dynamics simulation. The equilibration phase is not included. b, time series of C$_\alpha$-C$_\alpha$ distances between residues at the tips of the PAZ and N-terminal domain: Thr-218-Ser-73 (red), Arg-216-Glu-76 (green), Thr-218-Trp-40 (blue). c, cross-correlation map of C$_\alpha$ atomic fluctuations of AaAgo. The two axes refer to residue indices. Positive correlations are indicated in red, negative ones in blue. Positions of the N-terminal domain and the PAZ domain are indicated by bold lines. d, the cumulative summation of the overlap (29) between the first 20 modes determined by principal component analysis of the 5.0-ns MD trajectory and the vector related to the conformational change between our structure and 1YVU as a function of the mode number. Only C$_\alpha$ atoms were considered.
A consequence, closed conformations of Argonaute may already exist in the ensemble of the unbound protein.

Kinetic Behavior of ssDNA Binding to Argonaute—To further examine the interaction between ssDNA and Argonaute, we performed kinetic binding experiments using immobilized AaAgo. We used surface-sensitive fluorescence detection by total internal reflection fluorescence spectroscopy. The fluorescently labeled ssDNA is excited by the evanescent wave emanating from the surface. The interaction of 3’-labeled 21-nt ssDNA containing a 5’-phosphate with immobilized AaAgo shows biphasic association and dissociation kinetics (Fig. 5a). To exclude that the signal was due to ssDNA binding to the surface, a control experiment with 100 nm ssDNA in the absence of immobilized protein was run, showing minimal fluorescence signal (Fig. 5a). The interaction kinetics was independent of the immobilized protein concentration (0.06, 0.12, and 0.23 ng/mm², corresponding to 2, 5, and 10 nm protein injected onto the chip) (Fig. 5b). The slow dissociation phase could be fit with a single exponential decay, with a mean dissociation rate constant of 0.004 ± 0.002/s, corresponding to very slow dissociation of the ssDNA from Argonaute (Fig. 5c). No systematic dependence of the on the surface concentration of AaAgo was observed, which excludes the possibility that rebinding or multivalent interaction of the ssDNA with adjacent immobilized protein molecules was the source of the slow dissociation kinetics.

Under simple binding conditions involving a protein and ligand, the is the ratio of the association and dissociation rate constants (k_a/k_d). The kinetic behavior observed in the case of AaAgo and ssDNA precludes calculating a true K_d for the interaction. Nevertheless, a range of “K_d” values could be calculated based on the fitting of the two components of the association and dissociation curves. The K_d was estimated to be between 2.9 nm (fast association, slow dissociation) and 270 nm (slow association, slow dissociation), in relative agreement with the K_d measured by Yuan et al. (14) by filter binding (10 nM). We note that the identical nucleotide sequence and buffer conditions were used for these kinetic experiments with immobilized AaAgo.

We then examined the kinetics of binding as a function of the length of ssDNA, using 9- and 15-mer ssDNA in addition to the 21-mer in the previous experiments. Surprisingly, the kinetic behavior is nearly identical regardless of the length of ssDNA used (Fig. 5d). One interpretation of this finding is that the “hotspot” of binding energy is localized to the 5’-region of the ssDNA.

DISCUSSION

Domain Motions in Argonaute—Assuming that a range of PAZ orientations is represented by our crystal structure and 1YVU, our molecular dynamics simulation demonstrates that the unliganded form of Argonaute can assume a number of conformations (Figs. 1c, 3, a and b, and 4). The molecular dynamics simulation thus lends further support to the notion of a flexible PAZ domain and, to a lesser extent, the N-terminal domain. Furthermore, this is evident from the different conformations of AaAgo seen in our structure and 1YVU, as well as the unliganded structure of P. furiosus Argonaute. Thus the
FIGURE 5. Kinetics of binding of ssDNA to AaAgo. a, sensorgram of kinetics of ssDNA binding to immobilized AaAgo. The surface concentration of AaAgo was estimated to be 0.23 ng/mm², corresponding to an injection of 10 nM AaAgo (blue). A control with 100 nM ssDNA shows minimal binding (black). b, sensorgram of protein concentration dependence of ssDNA binding, with 0.06 (black), 0.12 (blue), and 0.23 ng/mm² (pink) surface concentration of AaAgo. c, single exponential fit, with residuals, of the slow dissociation at three protein concentrations (colors). d, length dependence of kinetics of ssDNA binding to AaAgo, for 9- (red), 15- (green), and 21-mer (blue) oligonucleotides. Curves were normalized to the peak fluorescence intensity.
closed and open forms of Argonaute may represent low energy conformations that become fixed in specific orientations upon nucleotide binding.

The biphasic kinetics of ssDNA binding to AaAgo is further suggestive of a conformational rearrangement in the protein. We note that the fluorescence intensity observed is primarily due to the interaction of the ssDNA with the protein and not due to the ssDNA alone (Fig. 5a). The total internal reflection fluorescence spectroscopy measurements show little difference in the association and dissociation kinetics of 9-, 15-, and 21-mer ssDNA nucleotides (Fig. 5d). We conclude the following from this observation. First, this suggests that the majority of the interaction energy is localized to the so-called seed region at the 5’-end and that the DNA beyond this point may be only loosely associated with Ago. Secondly, the kinetics of the ssDNA/AaAgo interaction are also independent of the length of ssDNA substrate used and the surface concentration of immobilized AaAgo. Together with the data from AaAgo crystal structures and the molecular dynamics simulation, the kinetic behavior points toward domain rearrangements during binding and dissociation, as reflected by the changes in DNA fluorescence (Fig. 5a). These data are fully consistent with the previously reported crystal structures of the Archaeoglobus fulgidus PIWI-dsRNA complex, which shows well ordered nucleotide density for the so-called “seed region” (37, 40).

**Functional Implications of PAZ Flexibility**—The structural, biochemical, and computational data presented here, in concert with the 1YVU and P. furiosus Argonaute structures, present an unprecedented opportunity to look at potential conformational changes in Argonaute during the cycle of mRNA recognition and cleavage by RISC. As the previous two structures of Argonaute have come from two different species with limited sequence identity (25%), the structure reported here allows for the first time the comparison of two Argonaute structures from the same species. In the context of the RISC, which has recently been identified as a complex composed of Dicer, Argonaute, and TRBP (11), our structure and biochemical data place constraints on the entrance and exit paths for the loaded Dicer-processed siRNA as well as the mRNA.

It is plausible that the initial loading of the Dicer-processed siRNA may proceed through the S groove prior to Argonaute cleavage of the passenger strand. The position of the PIWI box proximal to the S groove and the interaction between the PIWI box and the catalytic domains of Dicer support this hypothesis. The positively charged residues of the S groove may therefore contribute to the recognition of the siRNA in the “initial loading” complex.

Following cleavage of the passenger strand in the initial loading complex, the single-stranded guide strand-Argonaute complex is now in the catalytically active state. We have shown that unliganded Argonaute is flexible, most notably in the PAZ domain. Upon formation of the complex, the flexibility of the PAZ domain may be restricted. This may allow for recognition of the incoming target mRNA as well as discrimination between perfect and imperfect base pairing in the seed region. The mRNA strand may be directed to the seed region and the enzyme active site through two possible pathways, either along the M groove between the PAZ and N-terminal domains or along the S groove between the N-terminal and PIWI domains. This will have to be tested by further biochemical and structural studies.

The structure presented here, together with our molecular dynamics and kinetic studies, has highlighted a critical role for PAZ flexibility in the function of Argonaute. Future biochemical and structural studies will be necessary to further clarify the cycle of siRNA loading and mRNA recognition and cleavage by Argonaute.

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