Functional Characterization of *Entamoeba histolytica* Argonaute Proteins Reveals a Repetitive DR-Rich Motif Region That Controls Nuclear Localization

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**ABSTRACT** The RNA interference (RNAi) pathway regulates gene expression in many eukaryotic organisms. Argonaute (Ago) proteins, together with bound small RNAs (sRNAs), are key effectors that mediate gene silencing function. However, there is limited knowledge of Ago proteins and their functions in nonmodel systems. In the protozoan parasite *Entamoeba histolytica*, RNAi is a robust means for stable gene silencing mediated via large populations of antisense sRNAs. Here, we report functional characterization of three Ago proteins in *E. histolytica* (*EhAgo2-1*, *EhAgo2-2*, and *EhAgo2-3*). Our data show that each *Eh* Ago protein has a distinct subcellular localization and binds 27-nucleotide (nt) sRNAs and that the localization of *Eh* Ago proteins is altered in response to stress conditions. Via mutagenesis analyses, we demonstrated that the Ago PAZ (Piwi/Argonaute/Zwille) domain in all three *Eh* Agos is essential for sRNA binding. With mutation of the PAZ domain in *EhAgo2-2*, there was no effect on the nuclear localization of the protein but a strong phenotype and a growth defect. We further show that *EhAgo2-2* contains an unusual repetitive DR-rich (aspartic acid, arginine-rich) motif region which functions as a nuclear localization signal (NLS) and is both necessary and sufficient to mediate nuclear localization. Overall, our data delineate the localization and sRNA binding features of the three *E. histolytica* Ago proteins and demonstrate that the PAZ domain is necessary for sRNA binding. The repetitive DR-rich motif region in *EhAgo2-2* has not previously been defined in other systems, which adds to the novel observations that can be made when studies of the RNAi pathway are extended to nonmodel systems.

**IMPORTANCE** The protozoan parasite *Entamoeba histolytica*, which causes amebiasis and affects over 50 million people worldwide, contains an important RNAi pathway for gene silencing. Gene silencing via the RNAi pathway is mediated by the Argonaute (Ago) proteins. However, we lack knowledge on Ago function(s) in this nonmodel system. In this paper, we discovered that three *E. histolytica* Ago proteins (*EhAgo2-1*, *EhAgo2-2*, and *EhAgo2-3*) all bind 27-nt small RNAs and have distinct subcellular localizations, which change in response to stress conditions. The *Eh* Agos bind small RNA populations via their PAZ domains. An unusual repetitive DR-rich motif region is identified in *EhAgo2-2* that functions as a nuclear localization signal. Our results show for the first time an active nuclear transport process of the *EhAgo2-2* RNA-induced silencing complex (RISC) in this parasite. These data add to the novel observations that can be made when studies of the RNAi pathway are extended to nonmodel systems.

**KEYWORDS** Argonaute proteins, *Entamoeba*, RNA interference, gene expression, transcriptional regulation
involves an sRNA-guided Ago complex, i.e., RNA-induced silencing complex (RISC), to target mRNA for degradation (small interfering RNA [siRNA]-based function), transcriptional repression (microRNA [miRNA]-based function), or chromatin modification (epigenetic silencing function) (4). The complexity and plasticity of RNAi pathways have been demonstrated in higher eukaryotes, which often contain multiple Ago proteins with different sRNA classes (siRNA, miRNA, Piwi-interacting RNA [piRNA], secondary siRNA, etc.) (5). For example, there are eight Ago proteins in Homo sapiens, 10 Ago proteins in Arabidopsis thaliana, and 27 Ago proteins in Caenorhabditis elegans (6). In contrast, single-celled eukaryotes often show a simplified RNAi pathway with fewer Ago proteins. As an example, the yeast Schizosaccharomyces pombe has a single Ago protein (7), whereas no RNAi pathway is found in the related yeast Saccharomyces cerevisiae (7). Among the unicellular protists, the RNAi pathway is often less well studied; Ago proteins exist in Trypanosoma brucei (two Ago proteins), Toxoplasma gondii (three Ago proteins), Giardia lamblia (one Ago protein), and Trichomonas vaginalis (two Ago proteins) but are absent in Plasmodium and Trypanosoma cruzi (6, 8). Small RNA sequencing and experimental data for these organisms revealed functional RNAi pathways for retrotransposon control, gene regulation, and antigenic variation (6, 8).

The protozoan parasite Entamoeba histolytica causes amebiasis and is a major health concern in developing countries (9, 10). The parasite has two life stages: a dormant cyst form and an infective and invasive trophozoite form. The E. histolytica genome encodes several key RNAi machinery components, including three Ago proteins (EhAgo2-1 [EHI_186850], EhAgo2-2 [EHI_125650], and EhAgo2-3 [EHI_177170]), three RNA-dependent RNA polymerases (RdRPs) (EHI_139420, EHI_179800, and EHI_086260), and one protein (EHI_068740) with a single RNase III domain that has double-stranded RNA (dsRNA) cleavage activity but without a dsRNA binding domain (dsRBD) or other domains typically found in canonical Dicer enzymes (11, 12). We have previously reported that E. histolytica EhAgo2-2 has nuclear localization and binds to an atypical 27-nucleotide (nt) sRNA population that has 5′-polyphosphate (polyP) structure (13). We further linked these 27-nt sRNAs with endogenous gene silencing and developed an RNAi-based trigger method to silence genes of interest in this parasite (14). We demonstrated that RNAi-induced TGS in E. histolytica is established via histone modification at H3K27 and with the association of EhAgo2-2 protein (15). Furthermore, 27-nt sRNAs in Entamoeba are involved in regulation of strain-specific virulence genes but do not appear to regulate stage conversion between the trophozoite and cyst stages or the amebic stress response to heat shock or oxidative stress (16, 17). Our attempts using an RNAi-based trigger method to silence the three EhAgo genes failed to downregulate any of these transcripts, despite the fact that antisense sRNAs could be detected in parasites (18), suggesting that these genes may be essential and/or resistant to RNAi-mediated gene silencing. As to EHI_068740, our attempts using an in vitro dsRNA cleavage assay failed to show cleavage activity of this protein under standard experimental conditions (11). However, it partially contributes gene silencing in a heterologous system (S. cerevisiae) along with ScaAgo1 (11).

In this study, we have further characterized the three Ago proteins in E. histolytica. We demonstrated that the three EhAgo proteins have distinct subcellular localizations and all bind 27-nt sRNAs. We showed that expression and localization of EhAgo proteins change in response to two stress conditions, indicating a possible role for gene regulation under these conditions. We dissected the PAZ domain function in EhAgo by mutational analysis and demonstrated that EhAgo binding to sRNA was abolished/severely affected upon PAZ mutations; lack of sRNA binding in turn altered cellular localization of EhAgo2-1 and EhAgo2-3 but not EhAgo2-2. Furthermore, we identified an unusual repetitive DR-rich motif region in EhAgo2-2, not previously seen in other systems, which is necessary and sufficient to mediate nuclear localization. Our data provide the first functional analyses of the three E. histolytica Ago proteins, including the novel nuclear localization signal (NLS) function of the repetitive DR-rich motif region in EhAgo2-2, which adds to the known diversity of the structural and functional roles of Ago family proteins.
RESULTS

*Eh*Argonaute proteins have conserved structural PAZ and PIWI domains. RNAi gene regulation pathways and Ago proteins have been found across most eukaryotes (19). Ago proteins are generally conserved for four structural domains (the N terminus, PAZ, middle, and PIWI domains) (20). The *E. histolytica* genome contains genes encoding three Ago family proteins, *EhAgo2-1* (EHI_186850), *EhAgo2-2* (EHI_125650), and *EhAgo2-3* (EHI_177170) (21), with all three Ago proteins showing the conserved PAZ and PIWI domains (Fig. 1). Although the PAZ Superfam domain is annotated for all three *Eh*agos, the PAZ Pfam (PF02170) domain of *EhAgo2-1* and *EhAgo2-2* is shown in green. The PIWI Pfam (PF02171) domain is in blue. Specific PAZ/PIWI mutations generated in this study are shown. Both *EhAgo2-1* and *EhAgo2-2* have 200 aa after the PIWI domain, with an RGG/RR motif (red) and repetitive DR-rich region (gray) identified in *EhAgo2-2*.

**FIG 1** The structural domains (PAZ and PIWI) of three *Eh*agos in *E. histolytica*. The length of proteins is drawn to scale, with the length of PAZ and PIWI based on the annotation of AmoebaDB. We generated *Eh*ago polyclonal antibodies in rabbits using peptide from the N-terminal end of sequence as indicated by Y* (see also Materials and Methods). The PAZ Superfam domain (SSF101690) is shown in orange, and the PAZ Pfam (PF02170) domain of *EhAgo2-1* and *EhAgo2-2* is shown in green. The PIWI Pfam (PF02171) domain is in blue. Specific PAZ/PIWI mutations generated in this study are shown. Both *EhAgo2-1* and *EhAgo2-2* have 200 aa after the PIWI domain, with an RGG/RR motif (red) and repetitive DR-rich region (gray) identified in *EhAgo2-2*.

*The three *Eh*agos have distinct subcellular localizations.* Previously, we have reported that *EhAgo2-2* is localized in the nucleus (23). To characterize *EhAgo2-1* and *EhAgo2-3*, we generated custom-made polyclonal antibodies using selected N-terminal peptide sequences (Fig. 1). The antibodies were first tested by Western blot analysis using total cell lysates from wild-type and Myc-tagged overexpressing cell lines. The expected band size of each Ago was detected (Fig. 2A). However, due to low endogenous expression of both *EhAgo2-1* and *EhAgo2-3* (based on three published data sets using Affymetrix microarray [24, 25] or transcriptome sequencing [RNA-seq] [26], both *EhAgo2-1* and *EhAgo2-3* mRNA expression levels are detectable but are significantly lower than that of *EhAgo2-2*), they could not be detected in wild-type cell.
FIG 2  IFAs for the three EhAgos show distinct subcellular localization in the parasite, and stress conditions affect their expression/localization. (A) Custom-made antibodies to the three EhAgos detect specific bands by Western blotting. Western blot analyses were performed on total cell lysates from (Continued on next page)
lysates but only in lysates from overexpressing cell lines (Fig. 2A). We then performed immunostaining of trophozoites using these antibodies along with their preimmune sera. Immunofluorescence assay (IFA) with preimmune sera showed no specific signal at the background level (data not shown), but IFA with EhAgo2-1 antibody revealed two predominant subcellular localizations, perinuclear ring and cell surface membrane; staining with EhAgo2-3 antibody shows intense signal in both perinuclear and cytosolic localizations (Fig. 2B). Anti-EhFibrillarin is used to label the nucleolus, which was shown to be located at the nuclear periphery in *E. histolytica* (27). Localization of EhAgo2-2 has been previously shown to be in the nucleus (23) and is included for comparison.

In model systems, Ago proteins are found to be localized to many cellular loci depending on their biological functions, including cytoplasm (P-bodies and stress granules) for their PTGS-related functions and perinuclear regions or nucleus for TGS-related functions (20). In *Neurospora crassa*, RNAi components including RdRP, Dicer, and Ago are localized in the perinuclear region, where they function in meiotic silencing by unpaired DNA events (28, 29). In *C. elegans*, Ago proteins CSR-1, WAGO-1, and PRG-1 are localized to germ line perinuclear nuage, where epigenetic inheritance factors interact with Ago and RdRP (30, 31). Our results show that the three EhAgo proteins cover all major loci (cytoplasm, nucleus, and perinucleus), and each EhAgo has a unique localization, indicating that *Entamoeba* could possess diverse RNAi-related roles with each EhAgo protein having a unique function in the different aspects of RNAi.

Expression and localization of EhAgo proteins change in response to stress conditions. In human cells, hAgo2 is localized mainly in the cytoplasm and concentrated in P-bodies and is rapidly distributed to stress granules upon stresses (32), indicating a dynamic intracellular localization change of Ago with stresses. In *E. histolytica*, both cytoplasmic granules under stress condition and P-body-like structures have been reported (33, 34), and our previous transcriptional profiling showed high-level gene expression changes in *E. histolytica* after heat shock and oxidative stress (35, 36). We therefore used fluorescence microscopy to study accumulation/loss of expression of the three EhAgos under two conditions (42°C for 1 h or 1 mM H2O2 for 1 h).

The overall changes to the localization of endogenous EhAgo proteins under these conditions are shown in Fig. 2. For EhAgo2-1, the perinuclear ring in untreated cells was almost lost under H2O2 treatment but was unchanged with heat shock. With EhAgo2-2, the nuclear localization remained for both untreated and stress-treated cells; however, signal was greatly increased throughout the cytoplasm under stress conditions compared to almost no cytoplasmic signal in untreated cells (Fig. 2C). There are also some distinct punctate dots with induction of oxidative stress, similar to the stress granules reported earlier for *E. histolytica* (33). However, due to the lack of a marker for these

FIG 2 Legend (Continued)

wild-type and Myc-tagged overexpressing cell lines. The expected band size of EhAgo2-2 was detected in both lysate sample lanes; however, EhAgo2-1 and EhAgo2-3 can be detected only in lysates from overexpressing cell lines and not in wild-type cell lysates, likely due to their low endogenous expression, based on three published data sets using Affymetrix microarray (24, 25) or RNA-seq (26). Anti-actin was used as a loading control. (B) Distinct subcellular localizations are shown for each EhAgo in the parasite by IFAs. *E. histolytica* trophozoites were fixed and immunostained using custom peptide antibodies for EhAgo2-1, EhAgo2-2, and EhAgo2-3. Localization for EhAgo2-2 has previously been published (23) and was used as a control. A ring structure at the perinuclear location was seen for both EhAgo2-1 and EhAgo2-3, contrasting with the nuclear staining for EhAgo2-2. IFA. Additional signals were seen for EhAgo2-1 (cell surface membrane) and for EhAgo2-3 (cytosol). Anti-EhFibrillarin is used as a marker of the nucleolus, which is located at the nuclear periphery in *E. histolytica*. DAPI, 4',6-diamidino-2-phenylindole. (C) Localization of EhAgo proteins changes in response to stress conditions. IFAs of merged image of DAPI and each anti-EhAgo are shown for untreated parasites and parasites treated with two stress conditions (42°C for 1 h or 1 mM H2O2 for 1 h). Compared with the wild-type condition, the EhAgo2-1 localization of the perinuclear ring was almost lost under H2O2 treatment but was largely unchanged with heat shock. The EhAgo2-2 signal completely overlapped with the DAPI signal in untreated parasites but was greatly increased throughout the cytoplasm under both stress conditions, along with noticeable punctate dots, but its nuclear localization signal remained for both stress conditions. The EhAgo2-3 perinuclear localization is no longer observed under both stress conditions, and the major signal of staining is from cytoplasm. All stress condition assays were performed with multiple biological replicates and gave reproducible results; representative images are shown.
granules, we cannot definitively say if the *EhAgo* staining overlaps. To provide confirmation for the increased *EhAgo*-2-2 protein levels in the cytoplasm upon stress, we performed cell fractionation for nuclear and cytoplasmic proteins based on an earlier method (37). Western blot analysis using anti-*EhAgo*-2-2 showed increased cytoplasmic signal compared with untreated cells (see Fig. S2 in the supplemental material). For *EhAgo*-2-3, the perinuclear localization in untreated cells was completely lost under both stress conditions, and the majority of staining was in the cytoplasm. RNAi machinery and its gene regulation mechanism have been indicated for many biological events (38), and Ago can interact with diverse proteins and protein complexes, including RNA processing, maturation, and transport and the regulation of RNA stability and translation (2). Our data indicated that RNAi machinery actively responds to stresses in amebae. However, earlier sequencing of 27-nt sRNA libraries for parasites under these two conditions showed rather similar profiles of sRNA species (17). This may indicate that *EhAgo* localization changes help mediate changes in gene expression as a response to stress.

The three *EhAgo* proteins show specific sRNA binding that is abolished/severely degraded by YY-to-AA mutation in the PAZ domain. Phylogenic analyses for eukaryotic Ago proteins have divided all Ago proteins into three clades: the Ago-like, the Piwi-like, and the WAG0 subfamilies (6, 19, 39). Amebozoan Ago proteins are placed into a PIWI-like clade (6, 19). In order to identify conserved residues, we aligned three *EhAgo* PAZ and PIWI domains with human HIWI and *Drosophila* PIWI domain sequences using the Clustal Omega tool (Fig. S3). It is well documented that the PAZ domain binds the 3′ end of sRNAs with some highly conserved residues, the so-called R/K-F-Y signature sites (20). The alignment of three *EhAgo* PAZs showed that these signature residues are conserved for *EhAgo*-2-1 and *EhAgo*-2-2 as highlighted in the red box in Fig. S3, but not for *EhAgo*-2-3, for which the residues are W-K-Y (of note, two FF residues are right next to K, which may indicate that *EhAgo*-2-3 could have W-F-Y residues). In addition, the mutation of the two tyrosine residues in PAZ was shown to abolish sRNA binding in *C. elegans* (40). We therefore selected these two residues for mutagenesis as indicated in Fig. 1.

One characteristic of Ago proteins is the ability to bind sRNAs. We reported earlier that *EhAgo*-2-2 binds antisense 27-nt small RNAs and is linked to TGS in *E. histolytica* (13, 14). To check if sRNAs are also bound to *EhAgo*-2-1 and *EhAgo*-2-3, we overexpressed Myc-tagged protein for *EhAgo*-2-1 and *EhAgo*-2-3 using the same vector that was used for *EhAgo*-2-2 (13). Additionally, based on the sequence alignment of PAZ, we mutated the conserved YY to AA and generated stable overexpression transfectants for *Myc-EhAgo*-2-1 PAZmut and *Myc-EhAgo*-2-3 PAZmut mutants. We were not able to generate *Myc-EhAgo*-2-2 PAZmut-stably transfected parasites after multiple attempts. We speculated that the overexpression of *Myc-EhAgo*-2-2 PAZmut could be harmful to parasite growth and viability. Thus, to minimize the potential toxicity of *Myc-EhAgo*-2-2 PAZmut, we used a protein destabilization domain approach (41) to establish dihydrofolate reductase (DHFR)-*Myc-EhAgo*-2-2 PAZmut in parasites. The induction of DHFR-*Myc-EhAgo*-2-2 PAZmut protein was detected upon addition of a stabilizing agent (Fig. S4).

We used the cell lines described above to determine sRNA binding. For all three *EhAgo* Myc-tagged overexpressed cell lines and their PAZ mutants, we performed anti-Myc immunoprecipitation (IP) using whole-cell lysate. Both IP RNA and IP protein were collected; IP RNA was labeled by radioactive labeling after RNA separation on a denaturing gel; IP proteins were subjected to Western blot analysis. The data demonstrate that for *Myc-EhAgo*-2-1, the bound sRNAs form a smear at the 20- to 27-nt range, which disappears in the PAZ mutant (Fig. 3A). For *Myc-EhAgo*-2-2, a distinct 27-nt sRNA population is shown, as we previously observed (13); the PAZ mutation of *EhAgo*-2-2 shows no intact 27-nt sRNA but a severely degraded pattern of sRNA down to 18 nt. For *Myc-EhAgo*-2-3, there appear to be two sRNA populations at 27 nt and a smear of 20 to 24 nt, and upon generation of a PAZ mutant, there is no bound sRNA. Western blotting of IP proteins in the lower panels in Fig. 3A shows that wild-type and mutant lines pulled down similar levels of protein, indicating that the sRNA difference is not due to
FIG 3 Characterization of three EhAgos and their PAZ mutants for sRNA binding and protein localization. (A) Specific sRNA populations are bound to each Myc-EhAgo, and PAZ domain mutations abolished or severely degraded sRNA binding. Overexpressing Myc-tagged cell lines of wild-type EhAgos and PAZ mutants were used. For EhAgo2-2 PAZ mutant, a protein destabilization domain approach was used to express mutant protein upon addition of TMP. Shown in the upper panel is the pCp-labeled IP sRNA, showing specific sRNA populations bound to each EhAgo protein, which is absent in the control IP of the wild-type line or the uninduced Myc-DHFR-EhAgo2-2 PAZmut line. While Myc-EhAgo2-2 binds to a distinct 27-nt sRNA population, sRNA bands bound to Myc-EhAgo2-1 and Myc-EhAgo2-3 are rather smeary, composed of a noticeable 27-nt band along with its streak down to 20 nt. The PAZ mutations abolish the sRNA binding in Myc-EhAgo2-1 PAZmut and Myc-EhAgo2-3 PAZmut. However, the 27-nt band in Myc-EhAgo2-2 turned into a greatly degraded pattern in Myc-DHFR-EhAgo2-2 PAZmut. The lower panel is the anti-Myc Western blot analysis detecting specific Myc pulldown of Myc-EhAgo as well as its counterpart for the PAZ mutant. Both IP RNA and IP protein samples were made from the same anti-Myc IP pulldown experiment by splitting beads into half-and-half samples as described in Materials and Methods. (B) Anti-Myc IFAs show that the perinuclear ring localization is lost for mutant Myc-EhAgo2-1 PAZmut in comparison with wild-type Myc-EhAgo2-1. (C) Anti-Myc IFAs with and without TMP demonstrate that the mutant Myc-DHFR-EhAgo2-2 PAZmut has nuclear localization similar to wild-type Myc-EhAgo2-2. (D) Anti-Myc IFAs show that the perinuclear ring localization is lost for mutant Myc-EhAgo2-3 PAZmut in comparison with wild-type Myc-EhAgo2-3. (E) Growth kinetics for the Myc-DHFR-EhAgo2-2 PAZmut cell line. A significant reduction in the growth rate of the mutant cell line at days 4 and 5 was observed compared with the no-TMP control. Error bars are calculated as standard error from duplicate samples. We have extensively tested the effect of TMP on parasite growth in multiple Entamoeba strains and species and found no/minimal effect on growth rates (41).
FIG 3 (Continued)
differences in IP efficiency or protein expression level. In conclusion, the Myc IP for three PAZ mutant EhAgo lines failed to pull down specific sRNAs in EhAgo2-1 and in EhAgo2-3 mutants and severely degraded sRNA in the EhAgo2-2 mutant, indicating that the PAZ domain provides both binding and protection for sRNA, possibly on its 3’ end. Without it, sRNA is subjected to exonucleases for either complete degradation (in the case of EhAgo2-1 and EhAgo2-3 PAZ mutants) or partial degradation (in the case of the EhAgo2-2 PAZ mutant). These data confirm that PAZ RNA recognition is well conserved, and the binding mechanism of the 3’ end of sRNA of Ehagos likely uses the same mechanism as all other Agos in model systems (42).

Further, we performed fluorescence microscopy assay for these cell lines and observed a significant change in the localization of mutant proteins in Myc-EhAgo2-1 PAZmut and Myc-EhAgo2-3 PAZmut. The perinuclear localization of both Myc-EhAgo2-1 and Myc-EhAgo2-3 was absent in both PAZmut mutants (Fig. 3B and D). However, in DHFR-Myc-EhAgo2-2 PAZmut, the nuclear localization was unchanged compared to wild-type parasites (Fig. 3C). These data indicate that the PAZ domain is critical for sRNA binding for each of the three EhAgo proteins and that the protein/sRNA binding appears to affect localization for EhAgo2-1 and EhAgo2-3 but not for EhAgo2-2. As sRNA is a guide for RISC to find its target in different subcellular compartments, this may explain how PAZ mutants of EhAgo2-1 and EhAgo2-3 lost their specific protein localization. However, localization of the EhAgo2-2 PAZ mutant is unchanged despite severely degraded sRNA, indicating that either severely degraded sRNA can still function as a guide for RISC or/and there is a specific nuclear importing process for this protein that functions through the NLS pathway.

**The EhAgo2-2 PAZ mutant has a growth defect.** Given that we could not establish an EhAgo2-2 PAZ mutant without using the destabilization domain approach, we wondered whether the EhAgo2-2 PAZ mutation conferred a growth disadvantage on the parasites. The protein destabilization domain approach provides a way to study protein function for a short period of time upon adding a stabilizing agent. In order to identify the effect of the PAZ mutation on cell growth, we monitored the growth kinetics for the DHFR-Myc-EhAgo2-2 PAZmut cell line upon adding the stabilizing agent trimethoprim (TMP) and inducing the PAZ mutant protein. We observed a significant reduction in the growth rate of the DHFR-Myc-EhAgo2-2 PAZmut cell line at days 4 and 5 compared to dimethyl sulfoxide (DMSO) controls (Fig. 3E). Of note, the stabilization of EhAgo2-2 PAZ mutant protein is in the context of endogenous wild-type EhAgo2-2 expression. We have extensively tested TMP on parasite growth on different lines of culture and found no/minimal growth reduction (41); thus, the growth phenotype indicates that DHFR-Myc-EhAgo2-2 PAZmut may be functioning in a dominant negative manner when overexpressed. Further characterization of the amebic RISC will shed light on the mechanistic action of EhAgo2-2 function.

**EhAgo2-2 has an unusual repetitive DR-rich motif that mediates nuclear localization.** An interesting structural difference between Ehagos and model system human HIWI and DmPIWI is the presence in EhAgo2-1 and EhAgo2-2 of extended long tail sequences (~200 amino acids [aa]) after the PIWI domain (Fig. 4A). While a BLAST search for EhAgo2-1 tail sequences failed to show any meaningful homologous hits, the EhAgo2-2 C-terminal tail sequence is strikingly DR rich and comprises 12 repeats of RRDDRD motif sequences (Fig. 4A). Searches using both AmoebaDB and the NCBI BLAST tool found that this structure is not recognized as any known annotated functional domain; instead, it was designated a low-complexity segment. In *E. histolytica*, a single additional protein with this repeat was found, a U1 snRNA protein with likely nuclear localization (EHL_153670). Performing BLASTP on all eukaryotic genomes, we identified ~200 hits with ≥5 repeats of the RRDDRD motif (Table S3). Interestingly, most of these hits are from nonmodel systems, and Gene Ontology (GO) analysis indicated that many are either nuclear or RNA binding (Table S2). Out of all published Agos, there is only one hypothetical Piwi protein (EGT41773) from *Caenorhabditis brenneri*, with the DR-rich motif stretch appearing before the Piwi domain (Table S3).
FIG 4 An unusual repetitive DR-rich motif region in *EhAgo2-2* mediates nuclear localization. (A) The C-terminal repetitive DR-rich motif region in *EhAgo2-2*. Structural domains (PAZ and PIWI) of *EhAgo2-2* with two PIWI proteins in higher eukaryotes (*Hs*HIWI [33563234] and *Dm*PIWI [17136736]) are drawn to scale for comparison, showing that *EhAgo2-2* has an extended 200-aa region after PIWI. The C terminus of *EhAgo2-2* consists of a possible RGG/RG motif (green) and 12 repetitive RRDDR sequences that range from 8 to 14 aa. The dashed box lists the repeat sequences of C-terminal *EhAgo2-2*, arranged into separate lines for each RRDDR motif with highlights of arginine (R) (red) and aspartic acid (D) (blue). The underlined sequence is the predicted NLS. Two deletion mutants are made at positions as shown. A chimeric protein, *EhAgo2-3* with the DR-rich motif region from *EhAgo2-2*, is illustrated to scale. (B) Two C-terminal deletion mutants (Myc-*EhAgo2-2* DR-deletion and Myc-*EhAgo2-2* NLS-DR-deletion) cause Myc-*EhAgo2-2* protein localization to the cytosol. Anti-Myc IFA for both wild type and two deletion mutants are shown. The wild-type parasites have signal in the nucleus, which is in contrast with the deletion mutants, which are largely cytosolic. (C) The *EhAgo2-2* DR-rich motif region directs *EhAgo2-3* to the nucleus. Anti-Myc IFA shows that the chimeric protein *EhAgo2-3* is localized in the nucleus, compared with its wild-type localization, which is perinuclear and cytoplasmic. Anti-Myc IFA shows that the chimeric protein *EhAgo2-3* is localized in the nucleus, compared with its wild-type localization, which is perinuclear and cytoplasmic. (D) sRNA binding ability is unaffected either in the two *EhAgo2-2* deletion mutants or in the *EhAgo2-3* chimeric protein. The upper panel shows pCp-labeled IP sRNA for all cell lines, indicating that the bound sRNAs are in each line except control. The lower panel shows Western blot analysis for efficiency of Myc pulldown of proteins. With the addition of the DR-motif region tail, the *EhAgo2-3* chimeric protein migrates to a higher molecular weight as expected.
FIG 4 (Continued)
We further used several online NLS prediction programs (see Materials and Methods) and found a weak NLS sequence (GGRFGG) at the $Eh$ Ago2-2 C terminus, which is embedded within a stretch of sequences with a possible RGG/RG motif (Fig. 4A). This RGG/RG motif involves protein methylation at arginine (R), which is important for protein functioning in signaling, sorting, and transport and transcription activation (43). In order to study the function of this putative NLS and the DR-rich motif, we generated two deletion mutants: $Eh$ Ago2-2 DR-deletion and $Eh$ Ago2-2 NLS-DR-deletion (Fig. 4A). We overexpressed Myc-tagged protein for both deletion cell lines and used Western blotting to detect the predicted protein size (Fig. S5). Both deletion mutant proteins of $Eh$ Ago2-2 localized to the cytoplasm (Fig. 4B). IFAs using anti-$Eh$ Ago2-2 antibody showed that localization of endogenous $Eh$ Ago2-2 is unaffected (Fig. S6). As the $Eh$ Ago2-2 DR-deletion cell line still contains the predicted NLS sequence, it indicates that deletion of the DR-rich motif itself is sufficient to direct the change of $Eh$ Ago2-2 localization and does not require the predicted weak NLS sequence.

We reasoned that $Eh$ Ago2-2 could use its DR-rich tail region for nuclear localization. In order to test this idea, we made a chimeric $Eh$ Ago2-3 protein with the tail region from $Eh$ Ago2-2 fused to its C terminus (Fig. 4A), thus achieving a similar layout structure as in $Eh$ Ago2-2. IFA shows that the $Eh$ Ago2-3 chimeric protein is now localized to the entire nucleus (instead of its wild-type location in the perinuclear and cytosolic region) (Fig. 4C). This nuclear localization is similar to the localization for $Eh$ Ago2-2, indicating that the fused tail region of $Eh$ Ago2-2 is sufficient to localize another Ago protein to the nucleus. Last, we checked sRNA binding using anti-Myc IP for these deletion mutants and the chimeric protein of $Eh$ Ago2-3. The data show that both $Eh$ Ago2-2 deletion mutant proteins bind 27-nt sRNA comparably to the wild-type $Eh$ Ago2-2 (Fig. 4D). However, $Eh$ Ago2-3 chimeric protein shows a slightly different pattern; the 20- to 24-nt smear sRNA is no longer as obvious as its wild-type counterpart, possibly due to an exonuclease environment change of its changed localization.

In Entamoeba, molecular determinants for the protein trafficking into the nucleus are not well studied (44). The typical classical nuclear localization sequences (NLS of the simian virus 40 [SV40] large T antigen) do not appear to work for amebic parasites (reference 44 and data not shown). The previously described NLS from an $E$. histolytica protein, $Eh$ NCABP166 (45), did not work for several proteins in our experience (data not shown). Our data on the $Eh$ Ago2-2 tail region and its resulting NLS effect indicate that nuclear transport pathways are active in $E$. histolytica and that small regulatory RNAs and RISC are transported to the nucleus.

$Eh$ Ago2-2 PIWI mutants have no effect on protein localization and sRNA binding. The PIWI domain folds like RNase H and has endonucleolytic activity in some Agos for target mRNA cleavage (39). The crystal structure of Ago proteins further reveals a D-D-X triad motif (where X is D/E/H/K) for Ago-based Slicer. However, the $Eh$ Ago PIWI domain sequences appear to be very divergent, and the alignment of $Eh$ Ago PIWI domain with model systems shows that none have a complete D-D-X catalytic triad (Fig. S7), suggesting that all three $Eh$ Agos are likely non-Slicer Agos. The only partially aligned triad position for $Eh$ Ago2-2 is D-G-N, which we analyzed by mutagenesis as indicated in Fig. 1.

We generated three cell lines, $Eh$ Ago2-2 PIWID545A, $Eh$ Ago2-2 PIWIT747A, and $Eh$ Ago2-2 PIWID545A;N747A. All Myc-tagged mutants were checked by Western blotting for expression, and the sRNA binding profile was detected by radioactive pCp labeling of RNA prepared from anti-Myc IP samples. Analysis of Ago-bound sRNAs revealed that the 27-nt sRNA population is associated with both the wild type and PIWI mutants, indicating that sRNA binding is intact in these mutants (Fig. 5A). IFAs of these mutants using anti-Myc antibody further showed almost exclusively nuclear localization, similar to the wild-type parasites (Fig. 5B). These results indicate that $Eh$ Ago2-2 appears to be tolerant in terms of sRNA binding and localization on its PIWI mutants, implying that $Eh$ Ago2-2 may be indeed a non-Slicer.
DISCUSSION

The RNAi pathway regulates gene expression and is mediated by Ago and its bound small RNAs. Here, we report functional characterization of three Ago proteins in *E. histolytica*, *EhAgo2-1*, *EhAgo2-2*, and *EhAgo2-3*. We have defined their subcellular localization and showed how environmental stress impacts their localization. Mutagenesis analyses of the *EhAgo* proteins demonstrated that the PAZ domain is essential for *Eh* Ago sRNA binding, indicating a conserved role of the PAZ domain in sRNA loading in this organism. We further dissected the functional role of an *EhAgo2-2* repetitive DR-rich motif and found that this region can function as a nuclear localization signal and is sufficient to mediate nuclear localization of another Ago protein (*EhAgo2-3*). The DR-rich motif region in *EhAgo2-2* has not previously been defined in other systems and adds to the novel observations that can be made when studies of the RNAi pathway are extended to nonmodel systems.

Our data revealed that *EhAgo* proteins have distinct localizations: the nucleus (*EhAgo2-2*), perinuclear ring (*EhAgo2-1* and *EhAgo2-3*), cytosol (*EhAgo2-3*), and cell surface membrane (*EhAgo2-1*). The nucleus and perinuclear ring localizations are indicative of cellular locations for the TGS pathway that we and the others reported earlier in this parasite (14, 46–48). Whether or not cytoplasmic *EhAgo2-3* has a functional role in the PTGS process in this organism is currently unknown. The apparent cell surface membrane localization of *EhAgo2-1* is quite interesting, as some Agos in model systems have been reported to be present in both soluble and membrane fractions. Membrane-bound Ago proteins are often found either in the vicinity of the Golgi complex and endoplasmic reticulum or at endosomes/multivesicular bodies (MVBs), and their functions are thought to be related to sRNA loading or RISC assembly (49). Recently, studies on exosomes have shown that they contain cargos including Ago and a specific subset of miRNAs and mediate cell-to-cell communication or host-pathogen interaction (50). Thus, localization of *EhAgo* to the membrane could indicate that this parasite can use a similar strategy to assemble RISC or to release exosomes to deliver nucleic acid to surrounding parasites or to host cells. Future efforts will be important to pursue this avenue of investigation.

Our data also indicated that three *EhAgo* undergo localization changes in response to stress conditions. *E. histolytica* survives under harsh environmental conditions as well...

**FIG 5** *EhAgo2-2* PIWI mutants have no effect on sRNA binding or protein localization. (A) sRNA binding ability is unaffected in three PIWI mutants. Overexpressing Myc-tagged cell lines of three PIWI mutants (*EhAgo2-2* PIWID545A, *EhAgo2-2* PIWIN747A, and *EhAgo2-2* PIWID545A;N747A) were used for anti-Myc IP pulldown experiments. The upper panel shows pCp-labeled IP sRNA for all cell lines, indicating that the bound sRNAs are unchanged for PIWI mutants. The lower panel shows anti-Myc Western blot analysis for the IP proteins. (B) *EhAgo2-2* localization is unchanged in the overexpressing cell lines of three PIWI mutants. Anti-Myc IFAs for all three PIWI mutants show signal in the nucleus.
as inside host tissues, and previous studies revealed genome-wide gene regulation changes under stress conditions (35, 37, 51). In \textit{E. histolytica}, two cytoplasmic granules, P-bodies and stress granule-like structures, have been reported (33, 34), and these organelles have previously been shown as important sites for some RNAi pathways in model systems (2). We speculate that the parasite may use a dynamic network of \textit{EhAgo} complexes, which vary in composition and localization at distinct cellular sites of action under different cellular stimuli. Further determination of how the induction of cell stress can lead to altered Ago-containing complexes and the biological consequences of these changes is needed.

Our mutagenesis analysis results are summarized in Table 1. For \textit{EhAgo} PAZ domains, all three \textit{EhAgo} proteins abolished/severely degraded sRNA binding upon mutations on the conserved residues in the PAZ domain, confirming a conserved role of binding the 3’ end of RNA among all Ago proteins (19). The PAZ mutants also altered the perinuclear ring localization for \textit{EhAgo}2-1 and \textit{EhAgo}2-3 but did not alter \textit{EhAgo}2-2 nuclear localization. We have also identified an unusual DR-rich motif in \textit{EhAgo}2-2 which appears to control the localization of this protein to the nucleus; two deletion mutants indicated that the DR-rich motif does not alter sRNA binding but is sufficient to mediate nuclear localization of another \textit{EhAgo} protein. These findings are suggestive that this DR-rich motif may serve as an NLS. For \textit{EhAgo} PIWI domains, PIWI mutants of \textit{EhAgo}2-2 did not affect sRNA binding or localization; thus, the exact role of the PIWI domain of \textit{Ehagos} needs further future study. Future studies using these mutants for an \textit{in vitro} Slicer activity assay will help to elucidate specific PIWI domain function and determine if any of the \textit{EhAgos} contain Slicer function in this organism.

With the data presented in this paper as well as previous work (13–15, 23), including a strong growth defect shown by a PAZ mutant of \textit{EhAgo}2-2, a working model for the mechanism of TGS in \textit{E. histolytica} is proposed (Fig. 6). In this model, the polyP 27-nt sRNAs (probably generated by RdRP using a low level of mRNA as the template) are loaded into \textit{EhAgo}2-2, which needs the PAZ domain for sRNA 3’-end binding and provides full protection of sRNA. After sRNA loading, the RISC is translocated into the nucleus, using the tail region with the DR-rich motif as an NLS. The RISC targets specific loci for TGS using sRNA as a base-pairing guide, which leads to specific histone H3 modification for long-term silencing; we have previously shown that gene silencing in

### Table 1

| Cell line | Mutation(s) | Protein localization | sRNA binding |
|-----------|-------------|----------------------|--------------|
| Wild type |             |                      |              |
| Myc-\textit{EhAgo}2-1 | None | Cytoplasm/perinuclear ring | Yes |
| Myc-\textit{EhAgo}2-2 | None | Nucleus | Yes |
| Myc-\textit{EhAgo}2-3 | None | Cytoplasm/perinuclear ring | Yes |
| PAZ mutants |             |                      |              |
| Myc-\textit{EhAgo}2-1PAZ-mut | Y267A, Y268A | Cytoplasm | No |
| DHFR-Myc-\textit{EhAgo}2-2PAZ-mut | Y267A, Y268A | Nucleus | No |
| Myc-\textit{EhAgo}2-3PAZ-mut | Y259A, Y260A | Cytoplasm | No |
| \textit{EhAgo}2-2 C-terminal mutants |             |                      |              |
| Myc-\textit{EhAgo}2-2Δ3DR | 783–937 deletion | Cytoplasm | Yes |
| Myc-\textit{EhAgo}2-2ΔNLS-DR | 761–937 deletion | Cytoplasm | Yes |
| \textit{EhAgo}2-2 PIWI mutants |             |                      |              |
| Myc-\textit{EhAgo}2-2DS45A | DS45A | Nucleus | Yes |
| Myc-\textit{EhAgo}2-2N747A | N747A | Nucleus | Yes |
| Myc-\textit{EhAgo}2-2DS45A,N747A | DS45A, N747A | Nucleus | Yes |
| Chimeric protein |             |                      |              |
| Myc-\textit{EhAgo}2-3 plus C-terminal tail region of \textit{EhAgo}2-2 | Full-length \textit{EhAgo}2-3 fused with \textit{EhAgo}2-2 tail (761–937) | Nucleus | Yes |

*Listed are cell lines, specific mutations, protein localizations determined by anti-Myc IFA, and sRNA binding ability determined by anti-Myc IPs and pCp labeling of sRNAs.*
amebae is linked to a histone H3K27Me2 marker at both episomal and chromosomal loci through \( \text{EhAgo2-2} \) (15). A similar pathway was also observed in \( \text{C. elegans} \), where the Ago protein (NRDE3) uses a bipartite NLS that actively transports siRNAs from the cytoplasm to the nucleus for targeting nascent RNA transcripts and building repressive chromatin marks (40, 52). At present, we do not know the exact roles of \( \text{EhAgo2-1} \) and \( \text{EhAgo2-3} \) other than their localization and sRNA binding abilities. What small RNAs they bind and whether they also participate in this working model or possess their own pathways need to be further studied.

Overall, our data provide the first comprehensive functional analyses of the three \( \text{E. histolytica} \) Ago proteins. Identification of the unique DR-rich domain in \( \text{EhAgo2-2} \) that functions as an NLS adds to the mechanisms of protein localization. Future studies on identification of the sRNAs that bind to the three \( \text{EhAgo} \) proteins and the Ago protein silencing complex components will give more insights on the specific roles of the Ago proteins in amebic biology.
MATERIALS AND METHODS

Parasite culture and cell lines. *E. histolytica* trophozoites (HM-1:MMSS) were grown axenically under standard conditions in glass culture tubes with TYI-S-33 medium at 36.5°C as described previously (13, 53). Three *Eh* Ago cell lines constitutively overexpressing N-terminally Myc-tagged Ago protein were made by using the plasmid pKT3M as a backbone and cloned full-length sequences of the three Ago genes (EHI_186850, EHI_125650, and EHI_177170). The PCR products of three Ago genes were amplified from genomic DNA using primers listed in Table S1 in the supplemental material and cloned with SmaI and XhoI sites. The resulting constructs are pKT3M-Myc-EhAgo2-1, pKT3M-Myc-EhAgo2-2 (reported in reference 13 and used as a control), and pKT3M-Myc-EhAgo2-3. To make EhAgo PAZ and PIWI domain mutants, we used the NEB Gibson assembly cloning kit (New England Biolabs). The two fragments were PCR amplified for each EhAgo using primers listed in Table S1 and assembled with pKT3M-Myc backbone using protocols provided in the kit; the resulting constructs are pKT3M-Myc-EhAgo2-1PAZ-mut, pKT3M-Myc-EhAgo2-2PAZ-mut, and pKT3M-Myc-EhAgo2-3PAZ-mut. To make the EhAgo2-2PAZ-mut destabilization domain construct, we inserted the dihydrofolate reductase (DHFR) gene in-frame at the SmaI site between 3×Myc and EhAgo2-2PAZ-mut based on pKT3M-Myc-EhAgo2-2PAZ-mut plasmid, resulting in pKT3M-Myc-DHFR-EhAgo2-2PAZ-mut. To make EhAgo2-2 C-terminal deletion mutant constructs, we used the forward primer of EhAgo2-2 and the reverse primers of del-DR and del-NLS-DR as listed in Table S1; the resulting constructs are pKT3M-Myc-EhAgo2-2 DR-deletion and pKT3M-Myc-EhAgo2-2 NLS-DR-deletion; the primer sets amplify EhAgo2-2 for amino acids 1 to 783 and 1 to 761, respectively. To make EhAgo2-3 plus the DR-rich motif region of EhAgo2-2, we first performed site-directed mutagenesis on pKT3M-Myc-EhAgo2-2 using the QuickChange Lightning kit (Agilent) to generate a unique AvrII site at amino acid position 762 before the DR-rich motif region, and then full-length EhAgo2-3 was subjected to PCR using primers (forward (EhAgo2-3) and its AvrII reverse primer). The PCR product was cloned using Smal and AvrII sites. The three EhAgo2-2 PIWI mutants were also constructed using the NEB Gibson assembly cloning kit (New England Biolabs) using primers listed in Table S1. All plasmid constructs were confirmed by sequencing, and parasite transfectants were maintained at 6 μg/ml G418.

Polycyclonal antibody production and immunofluorescence assays. EhAgo polyclonal antibodies were generated in rabbits. The antigen peptide used for EhAgo2-2 was reported in reference 23. For EhAgo2-1, peptide sequence C-KKLYDLTYQYQSYFVNYFE-NH2 corresponding to the N-terminal amino acids 31 to 50 of EhAgo2-1, was used. For EhAgo2-3, peptide sequence C-ODPYPKEFTDLPQ2YEDVDO-NH2, corresponding to the N-terminal amino acids 25 to 44 of EhAgo2-3, was used. The antibodies were affinity purified on a peptide column (Open Biosystems). For IFA, we fixed cells using methanol-acetone (1:1) and followed previously reported protocols (23). Primary antibody (custom-made polyclonal antibodies for three *Eh* Agos and anti-EhFibrillarin [27] [a gift from Sudha Bhattacharya] or commercial anti-Myc) was used at a 1:250 dilution and incubated at 4°C overnight. Alexa 488 (anti-rabbit or anti-mouse) (Molecular Probes) was used as secondary antibody at a 1:1,000 dilution. Slides were mounted with Vectashield mounting medium (Vector Laboratories, Inc.), and images were collected using a Leica CTR6000 microscope using a BD CARVII confocal unit.

Stress stimuli and conditions. Both stress conditions were the same as previously published, H2O2 stress (1 mM H2O2 for 1 h) (35) and heat shock (42°C for 1 h) (31). We used *E. histolytica* HM-1:MMSS culture tubes for stress treatments. After treatment, tubes were shaken gently on the rim of an ice bucket to release any remaining attached cells, and the cells were collected by spinning at 1,000 rpm for 5 min and washed once with 1× phosphate-buffered saline (PBS). The cells were then transferred to Eppendorf tubes for methanol-acetone (1:1) fixation, followed by incubation of primary and secondary antibodies as described above; after a final wash, cells were mounted on the coverslip for imaging.

Cell fractionation for cytoplasmic and nuclear lysates. We followed the same protocol used in the lab (37). Briefly, both untreated and stress-treated cells were collected; resuspended in buffer A (10 mM HEPES, pH 7.9, 1.5 mM MgCl2, 10 mM KCl, and 0.6% [g/g]) with protease inhibitors, 1 μM leupeptin, 1 μM E-64-d, and 1× HALT protease inhibitor mixture (Pierce); and incubated on ice for 20 min. After low-speed centrifugation for 10 min at 1,000 × g at 4°C, the supernatant was collected as cytoplasmic fraction lysate. The pellet was then washed and resuspended in buffer C (20 mM HEPES, pH 7.9, 420 mM NaCl, 1 mM EDTA, and 1 mM EGTA supplemented with the same protease inhibitor mixture) on ice for 30 min. The supernatant from the last centrifugation at 18,000 × g for 20 min was collected as nuclear fraction lysate.

Cell lysate, immunoprecipitation (IP), and RNA isolation. We used a protocol similar to that in reference 54 with modifications. The lysis buffer (20 mM Tris-HCl [pH 7.5], 1 mM MgCl2, 10% [vol/vol] glycerol, and 50 mM NaCl) contained 0.5% (vol/vol) Nonidet P-40 (NP-40), plus 1 mM NaF, 1 mM dithiothreitol (DTT), 1 mM phenylmethylsulfonyl fluoride (PMSF), 2× of 100× EDTA-free protease inhibitors (100× stock diluted to 2× as final concentration; Thermo Scientific), and 1× RNase inhibitor, 1 U/μl. The cells from one T25 flask were lysed in 600 μl of lysis buffer and incubated on ice for 15 min. Lysate was centrifuged at 14,000 rpm for 30 min in a benchtop centrifuge at 4°C, and supernatant was saved at −80°C.

For IP with anti-Myc beads (Thermo Scientific), 20 μl packed beads was rinsed twice with cold 1× Dulbecco PBS (DPBS) with 0.01% Tween 20. After washing, 500 μl whole-cell lysate (1 to 2 μg/μl) was added and incubated for 2 h with rotation at 4°C, and the beads were then pelleted and washed six times (3× each) in basic lysis buffer containing PMSF, 0.1% (vol/vol) Tween 20, and 0.1% (vol/vol) NP-40 at 4°C. We collected both IP RNA and IP protein samples for each IP sample by splitting the bead-wash mixture at the last wash step. The IP RNA was isolated using TRIzol (Invitrogen) reagent according to standard protocol, and the IP proteins were released by adding 30 μl 1× reducing lane marker buffer (Thermo Scientific) and heated at 95°C for 5 min.
**RNA pCp labeling.** We followed similar procedures as in reference 13 for RNA labeling at 3′ termini. Typically, 2 μL RNA was ligated to [32P]-pCp using T4 RNA ligase (New England Biolabs), and the reaction mixture was incubated at room temperature for 2 h. The ligation reaction mixture was resolved directly on a denaturing 7 M urea-containing 15% polyacrylamide gel, and signal was exposed to a phosphor screen and imaged on a Personal molecular imager (Bio-Rad).

**Western blot analysis.** Standard Western blotting technique was used. We used 5% milk as a blocking reagent, and primary antibody was used at a 1:1,000 dilution, with incubation at 4°C overnight. The appropriate secondary antibody was used at a 1:10,000 dilution and incubated at room temperature for 1 h. Signal was developed with ECL (+ GE Healthcare, USA) and detected by the Kodak Image Station 4000R (Kodak, USA) or with a film processor (GE Healthcare). Antibodies used in Western blot analysis were anti-Myc antibody (Cell Signaling), anti-actin antibody (Cell Signaling), and the custom-made EhAgo polyclonal antibodies.

**Protein stabilization conditions and growth kinetics.** For Western blot analysis, trophozoites expressing Myc-DHFR-EhAgo2-2PNA-mma were collected after adding stabilization compound (10 μM TMP) for 24 h, and protein lysates were made and analyzed by Western blotting using anti-Myc antibody (Cell Signaling). For determination of growth kinetics, parasites were seeded at 40,000 cells/tube, no-drug control along with stabilizing compound (10 μM TMP) was set up in duplicate, and the number of parasites was counted every 24 h for 5 consecutive days after exposure to stabilizing compound.

**Sequence alignment, phylogenetic analysis, and motif search.** The three EhAgo sequences and their annotations, including Superfamily and Pfam domain regions, are based on www.amoebadb.org/amoeba. To search for orthologs in other ameba species, we used the orthology and synteny functional tab that is a built-in function for each gene in AmoebaDB. We then used full sequences to build a phylogenetic tree using a web tool (www.phylogeny.fr). The Clustal Omega tool was used for PAZ and PIWI alignment analysis (www.ebi.ac.uk/Tools/msa/clustalo/). For NLS prediction, we used several online tools, including NLS mapper (http://nls-mapper.iba.keio.ac.jp/cgi-bin/NLS_Mapper_form.cgi), PredictProtein (www.predictprotein.org), and NucPred (https://nucpred.bioinfo.se/nucpred/). For motif search, we used the built-in motif search function in AmoebaDB and NCBI BLASTP. The RRDDDR motif was used to search all eukaryotic proteins with the following parameters: E value, 1,000; word size, 2; window size, 40. Proteins with 5 or more copies of the repeat were identified, and conserved domains were identified (http://pfam.xfam.org).

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mSphere.00580-19.

**FIG S1**, TIF file, 0.1 MB.

**FIG S2**, PDF file, 1.3 MB.

**FIG S3**, PDF file, 0.1 MB.

**FIG S4**, PDF file, 0.1 MB.

**FIG S5**, PDF file, 0.3 MB.

**FIG S6**, PDF file, 0.9 MB.

**FIG S7**, PDF file, 0.2 MB.

**TABLE S1**, PDF file, 0.1 MB.

**TABLE S2**, PDF file, 0.1 MB.

**TABLE S3**, TXT file, 0.2 MB.

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