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Target RNA-directed tailing and trimming purifies the sorting of endo-siRNAs between the two Drosophila Argonaute proteins

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ABSTRACT

In flies, 22–23-nucleotide (nt) microRNA duplexes typically contain mismatches and begin with uridine, so they bind Argonaute1 (Ago1), whereas 21-nt siRNA duplexes are perfectly paired and begin with cytidine, promoting their loading into Ago2. A subset of Drosophila endogenous siRNAs—the hairpin-derived hp-esiRNAs—are born as mismatched duplexes that often begin with uridine. These would be predicted to load into Ago1, yet accumulate at steady-state bound to Ago2. In vitro, such hp-esiRNA duplexes assemble into Ago1. In vivo, they encounter complementary target mRNAs that trigger their tailing and trimming, causing Ago1-loaded hp-esiRNAs to be degraded. In contrast, Ago2-associated hp-esiRNAs are 2‘-O-methyl modified at their 3‘ ends, protecting them from tailing and trimming. Consequently, the steady-state distribution of esiRNAs reflects not only their initial sorting between Ago1 and Ago2 according to their duplex structure, length, and first nucleotide, but also the targeted destruction of the single-stranded small RNAs after their loading into an Argonaute protein.

Keywords: Argonaute; RNAi; miRNA; microRNA; small RNA sorting

INTRODUCTION

In Drosophila, two small silencing RNA pathways regulate gene expression in both the germ line and the soma: the microRNA (miRNA) and the small interfering RNA (siRNA) pathways. miRNAs act as post-transcriptional regulators in diverse aspects of animal development and physiology; siRNAs counteract viral infections and ensure genomic stability by targeting transposable elements (Ghildiyal and Zamore 2009). Both miRNAs and siRNAs are cleaved from longer double-stranded RNA (dsRNA) precursors by RNase III family ribonucleases. Dicer-1 liberates 22-nucleotide (nt)-long miRNAs from 65-nt stem–loop precursors (pre-miRNAs), whereas Dicer-2 cuts 21-nt exo-siRNAs off of long dsRNA derived from exogenous sources, such as viruses or 21/22-nt endo-siRNAs from endogenous dsRNA derived from convergently transcribed mRNAs, transposons, or long, partially self-complementary “hairpin” RNAs (Zamore et al. 2000; Bernstein et al. 2001; Hutvágner et al. 2001; Lee et al. 2004; Czech et al. 2008; Ghildiyal et al. 2008; Kawamura et al. 2008; Okamura et al. 2008a,b).

Dicer-1 and Dicer-2 bind to dsRNA-binding proteins that enhance their activities: Dicer-1 partners with the PB isoform of the protein Loquacious (Loqs-PB) to process pre-miRNAs, while Dicer-2 acts with the shorter Loqs-PD isoform to generate at least some classes of endo-siRNAs (Liu et al. 2003; Förstemann et al. 2005; Jiang et al. 2005; Saito et al. 2005; Park et al. 2007; Hartig et al. 2009; Zhou et al. 2009; Marques et al. 2010).

As a consequence of their being cleaved from dsRNA precursors, both miRNAs and siRNAs are born double-stranded, and these duplexes are loaded into members of the Argonaute family of proteins, which mediate their silencing functions. In addition to its role in producing siRNAs, Dicer-2 also participates in loading siRNAs into Argonaute2 (Ago2). For this function, Dicer-2 partners with the protein R2D2, a paralog of Loqs-PB, to form a heterodimer that
binds siRNA duplexes (Liu et al. 2003; Tomari et al. 2004b; Pham and Sontheimer 2005; Liu et al. 2006). Loading of small RNAs into Argonaute proteins also requires the HSP90-chaperone machinery (Iki et al. 2010; Iwasaki et al. 2010; Miyoshi et al. 2010).

*Drosophila* miRNAs typically assemble into Ago1, while siRNAs bind to Ago2—the Argonaute protein that mediates RNA interference—to form the RNA-induced Silencing Complex (RISC) (Okamura et al. 2004). In flies, the production and loading of small RNA duplexes are not coupled (Fürstemann et al. 2007; Tomari et al. 2007). Instead, miRNAs and siRNAs are sorted between Ago1 and Ago2 according to their duplex structure and the identity of their 5′ nucleotide (Ghildiyal et al. 2008; Czech et al. 2009; Okamura et al. 2009; Ghildiyal et al. 2010). Extensive double-stranded character, as is found in siRNAs, leads to assembly into Ago2, whereas central bulges and mismatches, typically found in miRNA/miRNA* duplexes, promote assembly into Ago1 (Tomari et al. 2007; Kawamata et al. 2009).

The affinity of a small RNA duplex for the Dicer-2/R2D2 heterodimer, a component of the complex that loads small RNAs into Ago2, determines the extent to which it loads into Ago2 (Liu et al. 2003; Tomari et al. 2004a,b, 2007). R2D2 binds to the more stable 5′ end of the small RNA duplex, orienting the siRNA so that the less tightly paired 5′ end becomes the guide strand bound to Argonaute (Tomari et al. 2004b). Central mismatches reduce the affinity of the small RNA duplex for Dicer-2/R2D2, antagonizing Ago2 loading and promoting loading into Ago1 (Fürstemann et al. 2007; Tomari et al. 2007; Kawamata et al. 2009). In addition to structure, the sequence of a small RNA guide influences sorting in many organisms (Lau et al. 2001; Ghildiyal et al. 2008; Mi et al. 2008; Montgomery et al. 2008; Hu et al. 2009; Ghildiyal et al. 2010). In flies, miRNAs bound to Ago1 usually start with uridine (U), whereas Ago2-bound miRNAs, miRNA* strands, and exo-siRNAs tend to start with cytidine (C) (Ghildiyal et al. 2010). The 5′ phosphate group of small RNA guides binds to a pocket in the MID (middle) domain of eubacterial and archaeabacterial Argonaute proteins (Ma et al. 2005; Parker et al. 2005). The equivalent domain in eukaryotic Argonautes may "read" the identity of the first nucleotide of a small silencing RNA, as the MID domain of human Ago2, which is closely related in sequence to fly Ago1, preferentially binds to adenosine and uridine 5′ monophosphate (Frank et al. 2010). The molecular basis for the 5′ nucleotide preference of *Drosophila* Ago2 is not yet known and could reflect a preference for C of the Ago2-loading machinery, Ago2 itself, or both.

While most exo- and endo-siRNA precursors are believed to comprise long, fully complementary sense and anti-sense RNAs, the siRNA precursors transcribed from the six *Drosophila* "structured" loci (esi-1/hp-CG18854, esi-2/hs-CG4068, hp-CG32207, hp-CP32205, hp-pncr009/pncr009: 3L, and an intergenic region adjacent to CG4770) contain mismatches, bulges, and wobble base pairs, characteristics usually associated with miRNA precursors (Fig. 1; Supplementary Figs. S1, S2; Czech et al. 2008; Kawamura et al. 2008; Okamura et al. 2008b). The duplexes of the "hp-esiRNAs" derived from these hairpin transcripts resemble miRNAs rather than canonical siRNAs (Fig. 1A). Yet hp-esiRNAs accumulate mainly in Ago2. Why?

We recently reported that the binding of an Ago1-bound miRNA to an artificial, highly complementary target RNA triggers its tailing and 3′-to-5′ trimming, ultimately decreasing the steady-state concentration of the miRNA (Ameres et al. 2010). Hen1, the methyltransferase that adds a 2′-O-methyl group to the 3′ end of Ago2-bound small RNAs (Horwich et al. 2007; Pelisson et al. 2007), protects Ago2-bound siRNAs against such tailing and trimming (Hutvágner and Zamore 2002; Haley and Zamore 2004; Ameres et al. 2010). Here, we show that, in vitro, hp-esiRNA duplexes assemble into Ago1 or Ago2 according to their sequence, structure and length: Ago1 preferentially binds to 22 nt, bulged small RNAs starting with A or U; Ago2 associates with
21-nt, double-stranded small RNAs beginning with C or U. Like exo-siRNAs and miRNA* strands, hp-esiRNAs that begin with C bind Ago2 in vivo. In contrast, hp-esiRNAs that start with U bind mainly Ago1 in vitro yet accumulate in Ago2 in vivo. We present evidence that in vivo these hp-esiRNAs encounter complementary target mRNAs that trigger their degradation via the tailing and trimming pathway. Since Ago2-associated hp-esiRNAs bear a protective 2′-O-methyl modification at their 3′ ends, they escape tailing and trimming and accumulate. Thus, the steady-state distribution of hp-esiRNAs between Ago1 and Ago2 reflects not only their partitioning during the loading of Argonaute proteins, but also the target-directed destruction of specific hp-esiRNA species after their assembly into Ago1 complexes.

RESULTS AND DISCUSSION

Hp-esiRNAs predictably partition between Ago1 and Ago2 in vitro

Their partially double-stranded structure predicts that hp-esiRNAs should load into Ago1, yet in vivo they accumulate in Ago2 (Kawamura et al. 2008; Czech et al. 2009; Okamura et al. 2009). To test whether hp-esiRNAs violate the previously established rules for Drosophila small RNA sorting, we examined their loading into Ago1 and Ago2 in Drosophila embryo lysates. The esi-1 (hp-CG18854; Figure S1) and esi-2 (hp-CG4068; Figure S2) loci produce 98% of all mature hp-esiRNAs in fly heads and 96% in S2 cells (Supplemental Table S1; Czech et al. 2008; Kawamura et al. 2008; Okamura et al. 2008b). Among the most abundant hp-esiRNAs derived from these two loci, esi-2.1 and esi-1.1 are 22 nt long and begin with U; esi-2.1 contains a mismatch at position 8 of its predicted guide/passenger duplex (Fig. 1B, top row). These are all features of Ago1-loaded small RNAs (Ghildiyal et al. 2010). Other hp-esiRNAs resemble Ago2-associated small RNAs. For example, esi-1.2 is 21 nt long, begins with C, and has a paired central region.

We used 254-nm UV-crosslinking to measure the partitioning of synthetic, 5′-phosphorylated hp-esiRNA duplexes between Ago1 and Ago2 (Fig. 1B). Esi-2.1 and esi-1.1 crosslinked efficiently to Ago1, although at lower levels than the let-7/let-7* duplex. Crosslinking to Ago2 was detectable but much less efficient than a canonical let-7 siRNA. In contrast, esi-1.2 crosslinked to Ago2 but not Ago1, just like a canonical siRNA (Fig. 1B). We conclude that, in vitro, hp-esiRNAs follow the established rules for small RNA sorting in flies, partitioning according to their duplex structure and first nucleotide (Czech et al. 2008; Okamura et al. 2009; Ghildiyal et al. 2010).

5′ Nucleotide identity and length influence small RNA partitioning between Ago1 and Ago2

In plants, the 5′ nucleotide of a small silencing RNA determines how it partitions among the different Argonaute proteins (Mi et al. 2008; Montgomery et al. 2008). A similar mechanism operates in some animals (Ghildiyal et al. 2008; Frank et al. 2010; Ghildiyal et al. 2010). In flies, Ago2-bound small RNAs, including exo-siRNAs and miRNA* species, typically begin with C, whereas Ago1-associated small RNAs (mostly miRNAs) begin with U (Ghildiyal et al. 2008, 2010). To disentangle the influence of structure and 5′ nucleotide identity, we examined the loading of four variants of the let-7/let-7* duplex and four variants of an siRNA in which the guide strand corresponded to let-7 (Fig. 2A). To ensure selection of let-7 as the guide strand, the first nucleotide of let-7 was unpaired in all duplexes (Schwarz et al. 2003; Tomari et al. 2007).

As observed previously (Ghildiyal et al. 2010), the let-7/let-7* duplex more efficiently assembled into Ago1 when
let-7 began with U than when it began with C (Fig. 2A, left; Supplemental Fig. S3). The let-7/let-7* duplex that began with A also loaded Ago1 efficiently, whereas loading of a let-7/let-7* duplex beginning with G was as inefficient as the duplex in which let-7 began with a C. Experiments in which each of the four let-7/let-7* variants was used as a competitor for loading a 5′-32P-radiolabeled canonical let-7/let-7* duplex confirmed these results (Supplemental Fig. S4). Conversely, a let-7 siRNA duplex that began with a pyrimidine (C or U) was efficiently loaded into Ago2, whereas the same siRNA duplex, but starting with a purine (G or A), loaded into Ago2 less efficiently (Fig. 2A, right; Supplemental Fig. S3). Among pyrimidines, Ago2 preferred C over U.

Distinct lengths of small RNAs are bound to Ago1 and Ago2 in flies: Ago1-associated small RNAs are 22–23 nt, while nearly all small RNAs bound to Ago2 are 21 nt (Czech et al. 2008; Ghiildiyal et al. 2008). The distinct RNA lengths likely reflect the intrinsic characteristics of the Dicer proteins that produce the small RNAs bound to Ago1 versus Ago2: Dicer-1 generates ~22–23 nt miRNAs, which typically load into Ago1, while Dicer-2 produces 21 nt siRNAs, which bind to Ago2 ( Zamore et al. 2000; Bernstein et al. 2001; Hutvágher et al. 2001; Lee et al. 2004). Does the propensity of Dicer-1 to cleave pre-miRNAs into 22–23 nt miRNA/miRNA* duplexes reflect a specific preference of Ago1 to load longer small RNA duplexes than Ago2? To test this idea, we compared the loading of four let-7/let-7* duplexes and four let-7 siRNA duplexes into Ago1 and Ago2. Each duplex comprised either two 21-nt or two 22-nt RNAs, with the let-7 strand beginning with either C or U. Figure 2B shows that the 22-nt miRNA/miRNA* duplexes were more efficiently loaded into Ago1, whereas the 21-nt siRNA duplexes were more efficiently loaded into Ago2.

For Ago1, the favored 22-nt length takes precedence over the disfavored 5′ C: a 22-mer miRNA starting with C loaded Ago1 as efficiently as a 21-mer miRNA starting with U (Fig. 2B, left). In contrast, Ago2 preferred a 21-nt duplex—even one in which the guide strand began with U—over a 22-nt duplex beginning with either C or U (Fig. 2B, right). These data are consistent with our finding that in Drosophila embryo lysate, esi-1.1 and esi-2.1—both 22 nt long—preferentially assemble into Ago1, whereas the 21-nt long esi-1.2 predominantly associates with Ago2 (Fig. 2B).

We analyzed high-throughput sequencing data of small RNAs that co-immunoprecipitated with Ago1 or Ago2 in S2 cells and of small RNAs from fly heads that co-immunoprecipitated with Ago1 or were inferred to be associated with Ago2 because of their resistance to oxidation (Czech et al. 2008; Ghiildiyal et al. 2010). In S2 cells (Fig. 2C), Ago1-bound hp-esiRNAs tended to start with U (42%) or A (32%) rather than C (15%) or G (11%), whereas Ago2-bound hp-esiRNAs tended to start with C (40%) or U (36%) rather than A (13%) or G (11%). Similarly, in fly heads, the hp-esiRNAs bound to Ago1 tended to start with U (58%), A (19%), or C (20%), but not G (3%), whereas those bound to Ago2 began with U (49%) or C (31%), but less often with A (13%) or G (7%). Small RNA assembly in vitro (Figs. 1B, 2A) accurately recapitulated the distribution of 5′ nucleotide identities observed in vivo: Ago1 preferentially loaded small RNAs starting with U or A, but not C or G; Ago2 associated predominantly with small RNAs beginning with C or U but not G or A.

Mutations in the RNAi pathway change the hp-esiRNA repertoire

We analyzed the genetic requirements for hp-esiRNA accumulation from each of the six Drosophila hairpin loci. As reported previously, wild-type hp-esiRNA accumulation in fly heads required dicer-2, r2d2, and ago2 (Fig. 3A; Supplemental...
Hp-esiRNAs are generally more abundant in Ago2 than in Ago1: We analyzed high throughput sequencing data sets for the small RNAs co-immunoprecipitated with Ago1 and Ago2 in S2 cell extracts, as well as from fly heads in which Ago1-bound small RNAs were isolated by co-immunoprecipitation with Ago1 and Ago2-bound small RNAs inferred from their resistance to oxidation. In S2 cells, hp-siRNAs were twice as abundant in Ago2 than in Ago1; in fly heads, hp-siRNAs were about four times more abundant in Ago2 than in Ago1 (Figure S7).

Individual hp-esiRNA species have a common requirement for dicer-2, but distinct requirements for r2d2 and ago2 (Fig. 3C). Esi-2.1, esi-1.1, and esi-1.2 each decreased 13- to 26-fold in dicer-2ΔH16X mutant fly heads, consistent with the idea that Dicer-2 cleaves hp-esiRNAs from their precursor hairpin transcripts. The concentration of esi-1.2, which in vitro was assembled into Ago2 (Fig. 1B), declined >19-fold in ago2Δ44 and ~10-fold in r2d2Δ mutant flies (Fig. 3C; Supplemental Fig. S6), yet esi-2.1 and esi-1.1, which in vitro were loaded predominantly into Ago1 (Fig. 1B), decreased only threefold (esi-2.1) or twofold (esi-1.1) in ago2Δ14 mutants (Fig. 3C; Supplemental Fig. S6). These data suggest that esi-1.2 is destroyed when it cannot be loaded into Ago2, whereas when esi-2.1 and esi-1.1 cannot be loaded into Ago2, they accumulate bound to Ago1. The >30-fold decrease of esi-1.1 in the absence of R2D2 might reflect a role for R2D2 in dicing esi-1.1 from its precursor (see below).

The hp-esiRNAs that persist in ago2Δ14 mutants were more likely to begin with U (77%), compared to those in wild-type Oregon R fly heads (49%) (Fig. 3D). Conversely, the hp-esiRNAs that persist in ago2Δ14 mutants were less likely to begin with C (13% in ago2Δ14 versus 28% in wild type) or A (4% in ago2Δ14 versus 17% in wild type). The fraction of hp-esiRNAs starting with U (54%) also increased in mutant flies homozygous for r2d2Δ compared to their heterozygous siblings (42%), and the fraction beginning with C decreased (17% in r2d2Δ homozygotes versus 33% in heterozygotes). Thus, loss of the RNAi machinery causes not only a decrease in hp-esiRNAs, but also changes the hp-esiRNA repertoire from one that bears the hallmarks of Ago2-loaded small RNAs to a population with the characteristics of Ago1-bound small RNAs.

FIGURE 4. The RNAi loading complex loads hp-esiRNAs starting with C into Ago2. (A) Correlation analysis of the change in abundance of hp-esiRNAs in r2d2Δ mutants and ago2Δ14 mutants (r = 0.47; P = 0.0002). The most abundant hp-esiRNA guide (red) and passenger strands (blue), as well as a trimmed species of esi-2.1 (green) are indicated. (B) Same data as in A, but with 5′ nucleotide identity indicated by color: green, 5′ A; blue, 5′ C; yellow, 5′ G; red, 5′ U. (C) The 5′ nucleotide composition of the hp-esiRNAs species in (B) ranked by the change in abundance (<; top panel; >; bottom panel) in ago2Δ14 and r2d2Δ. (<;SD) Below the limit of detection.
whereas the corresponding passenger strands, which are normally destroyed during RISC maturation, accumulated (Fig. 4A). Small RNAs derived from the esi-2 locus, but not the esi-1 locus, correlated well between ago2 and r2d2 mutants (Supplemental Fig. S8B,C). For esi-1.1 both the guide and passenger strands decreased in r2d2 mutants, suggesting that R2D2 might act in its biogenesis (Fig. 4A). Nevertheless, hp-esiRNAs from the esi-1 locus decreased less in r2d2 mutants compared to ago2 mutants and their decrease in r2d2 mutants does not correlate well with dicer-2 mutants, which would be expected if R2D2 assisted Dicer-2 in the processing of esi-1 hairpins (Supplemental Fig. S8B).

To determine if the RISC loading complex plays an active role in selecting for specific 5’ nucleotides when loading a small RNA guide into Ago2, we analyzed the change in abundance of hp-esiRNAs in ago2^{214} and r2d2 mutants as a function of the 5’ nucleotide of the small RNAs (Fig. 4B,C). Since the small RNAs that increased in abundance in r2d2 mutants tend to be passenger strands (Fig. 4A), we analyzed only small RNAs that declined in r2d2. Hp-esiRNA species that decline in both ago2 and r2d2 mutants exhibited a mixed 5’ nucleotide composition (33% A, 24% C, 12% G and 31% U). Among those, the species that decreased most in r2d2 and ago2 mutants tend to start with C: 35% of hp-esiRNAs that decreased more than fourfold and 56% of those that decreased more than 16-fold in both mutants begin with C (Fig. 4C, bottom panel). These data suggest that the RISC-loading complex specifically loads into Ago2 those hp-esiRNAs that bear a 5’ C. We obtained similar results for Ago2-enriched miRNAs and miRNA* strands in ago2^{214}, r2d2 and dicer-2^{m16}X mutants (Figure S9).

In contrast, hp-esiRNAs that were less dependent on R2D2 and Ago2 tended to begin with U: 53% of species that decreased in both ago2^{214} and r2d2 mutants less than 16-fold and 83% of those that changed less than fourfold started with U (Fig. 4C, top panel). 5’ U small RNAs constitute 49% of Ago2-bound hp-esiRNAs in wild-type flies and 36% in S2 cells (Fig. 2C). The increase in 5’ U small RNAs in r2d2 and ago2^{214} mutants (Fig. 3D) is therefore likely a result of their ability to load into Ago1 (58% of small RNAs bound to Ago1 in fly heads and 42% in S2 cells start with U) (Fig. 2C), compensating partially for the loss of Ago2-bound small RNAs.

Esi-1.1 provides an example of 5’ nucleotide-dependent accumulation of small RNAs in Ago1 and Ago2: esi-1.1 was originally annotated as a 22-nt small RNA starting with a U (henceforth, 5’ U esi-1.1) (Czech et al. 2008), but also exists as an abundant 21-nt isoform, 5’ C esi-1.1, that begins with the second nucleotide of the annotated species (Supplemental Fig. S10). The two isoforms are differentially expressed in fly heads, ovaries, and S2 cells (Supplemental Fig. S10): 5’ C-esi-1.1 is more abundant in heads, whereas 5’ U esi-1.1 predominates in ovaries. In S2 cells, the two isoforms are equally abundant. Although the guide/passenger duplex structure of the two isoforms is essentially identical, they differentially partition between Ago1 and Ago2: 5’ U esi-1.1 assembled in vitro into both Ago1 and Ago2 and was bound in vivo to both Ago1 and Ago2 (Supplemental Fig. S10). In contrast, 5’ C esi-1.1 assembled in vitro almost exclusively into Ago2 and in vivo was only detected bound to Ago2 (Supplemental Fig. S10). Moreover, 5’ C esi-1.1 decreased 23-fold in ago2^{214} mutants, but 5’ U esi-1.1 decreased only twofold (Fig. 4A; Supplemental Fig. S11). The length of the two esi-1.1 isoforms likely contributes to their differential partitioning between Ago1 and Ago2, since in vitro, Ago1 favors 22 nt RNAs, while Ago2 favors 21 nt RNAs (Fig. 2).

### Highly complementary target RNAs affect the stability of Ago1-bound hp-esiRNAs

Given the inherent preference for loading 22-nt RNAs into Ago1 and 21-nt RNAs into Ago2, we anticipated that the median hp-esiRNA length would increase in r2d2 and ago2^{214} mutant flies. This was not the case: 22- and 21-nt hp-esiRNAs decreased in both mutants (2.6- and 3.3-fold for 22-mers and 2.1- and 3.1-fold for 21-mers in r2d2 and ago2^{214} mutants, respectively) (Fig. 3A). Paradoxically, full-length hp-esiRNA isoforms (21 and 22 nt) decreased in ago2^{214} and r2d2 mutants, while the abundance of isoforms =20 nt generally was essentially unchanged (ago2^{214}) or increased slightly (1.4-fold; r2d2), constituting >20% of all hp-esiRNAs reads in those mutants (Fig. 3A; Supplemental Fig. S12). Such shorter species correspond to 3’ trimmed versions of full-length hp-esiRNAs.

In flies, a target mRNA that is highly complementary to a small RNA bound to Ago1 (but not Ago2) can trigger 3’-to-5’ exonucleolytic trimming and 3’ tailing of a small RNA (Ameres et al. 2010). Might the 3’ trimmed hp-esiRNAs we detected in ago2^{214} and r2d2 mutants be produced by 3’ tailing and trimming of Ago1-bound esiRNAs in response to complementary target mRNAs? In wild-type flies, such target-directed tailing and trimming could deplete the hp-esiRNA population of Ago1-bound small RNAs, explaining why Ago2-bound hp-esiRNAs are normally more abundant than those bound to Ago1 (Supplemental Fig. S7; Czech et al. 2008; Kawamura et al. 2008; Okamura et al. 2008b). In fact, esi-2.1, which loads into Ago1 in vitro but accumulates in Ago2 in vivo (Figs. 1C, 5C), is highly complementary to the mus308 mRNA (Supplemental Fig. S13; Czech et al. 2008; Okamura et al. 2008b). Perhaps esi-2.1 is initially loaded into both Ago1 and Ago2, but the portion of esi-2.1 loaded into Ago1 is then destroyed via the target-directed tailing and trimming pathway.

To test this idea, we compared the esi-2.1 isoforms in S2 cells that are bound to Ago1 with those that are bound to Ago2 (Fig. 5A, top row). Nearly all the Ago2-bound esi-2.1 was 22 nt, whereas the most abundant isoform of Ago1-bound ei-2.1 bound was 20 nt. In contrast, 90% of Ago1-bound miRNAs were at least 21 nt, 67% were 21 (32%) to 22 nt (35%), and <10% were <20 nt or shorter (data not shown). Target-directed trimming is accompanied by tailing, the
nontemplated addition of U or A to the 3' end of small RNAs bound to Ago1 (Ameres et al. 2010). If the shorter forms of Ago1-bound esi-2.1 reflect target-directed trimming, then we would expect to also detect tailed esi-2.1 species. Indeed, the fraction of esi-2.1 RNAs in Ago1 that had nontemplated nucleotides added to their 3' ends was three times greater than those bound to Ago2 (Fig. 5A, bottom). The most abundant 3' nontemplated nucleotide for Ago1-bound esi-2.1 was U (73%).

In S2 cells, esi-2.1 was approximately sixfold enriched in Ago2 versus Ago1, whereas all other esi-2-derived hp-esiRNAs accumulated equally in Ago1 and Ago2 (Fig. 5B). Unlike esi-2.1, the other Ago1-bound hp-esiRNAs derived from the esi-2 locus showed little evidence of trimming (Fig. 5C). The size distribution of these esi-2 species between Ago1 and Ago2 reflects the expected preference for 22-nt RNAs to load into Ago1 and 21-nt RNAs to load into Ago2 (Fig. 5C). We suggest that in S2 cells, the partitioning of esi-2.1 between Ago1 and Ago2 is purified by tailing and trimming directed by *mus308* mRNA and possibly other highly complementary target RNAs (Supplemental Fig. S13), leading to the destruction of Ago1-bound esi-2.1. Because there are no mRNA targets with similarly high complementarity for the other hp-esiRNAs derived from the esi-2 locus, they accumulate in both Ago1 and Ago2. Fly heads express additional target mRNAs with considerable complementarity to esi-2.1: *stat92E*, *gatA*, and *CG7359* (Supplemental Fig. S13B). Consistent with this observation, the abundance of full-length esi-2.1 decreased in *ago2* nonsense and *r2d2* mutants, while the abundance of 3' trimmed and tailed esi-2.1 isoforms increased (Fig. 3C; Supplemental Figs. S12, S14).

Hp-esiRNAs derived from the esi-1 locus are complementary to the CG8289 mRNA (Supplemental Fig. S13; Czech et al. 2008; Okamura et al. 2008b), and, like esi-2.1, those esi-1 RNAs that were associated with Ago1, but not those bound to Ago2, were trimmed and contained 3' nontemplated U's and A's (Fig. 5D).

Unlike Ago1-loaded small RNAs, Ago2-loaded small RNAs are protected from tailing and trimming by a 2′-O-methyl group deposited on the 3′ end of the small RNA guide by the methyltransferase Hen1 (Horwich et al. 2007; Pelisson et al. 2007; Ameres et al. 2010). In *hen1* nonsense mutant fly heads, the abundance of full-length esi-2.1 and full-length (21–22 nt) hp-esiRNA species derived from the esi-1 locus each decreased approximately twofold; the abundance of tailed and trimmed hp-esiRNA 2.1 and hp-esiRNAs derived from the esi-1 locus each increased ~17-fold and approximately fivefold, respectively, in the *hen1* mutant (Fig. 6A). Additionally, loss of Hen1 changed the nature of the hp-esiRNA repertoire (Fig. 6B): the hp-esiRNAs that persisted in *hen1* nonsense mutants tended to start with U (56% versus 44% in *hen1* heterozygotes), whereas hp-esiRNAs that begin with C were reduced (22% versus 31% in *hen1* heterozygotes), as would be expected if Ago2-bound hp-esiRNAs are tailed and trimmed in the absence of Hen1.

**FIGURE 5.** Ago1- but not Ago2-bound hp-esiRNAs are destabilized by highly complementary target RNAs. (A) Size distribution of genome-matching (top) and prefix-matching (bottom) esi-2.1 sequences associated with Ago1 (left) or Ago2 (right) in S2 cell extracts. The most abundant 3' nontemplated, added nucleotides are noted. (B) The abundance of esi-2.1 (top) or all other small RNAs from the esi-2 locus (bottom) associated with Ago1 (red) or Ago2 (blue) in S2 cell extracts. (C) Size distribution of small RNAs, other than esi-2.1, derived from the esi-2 locus and associated with Ago1 (top) or Ago2 (bottom) in S2 cell extracts. (D) Size distribution of genome-matching (top) and prefix-matching (bottom) small RNAs from the esi-1 locus associated with Ago1 (left) or Ago2 (right) in S2 cell extracts. The most abundant 3' nontemplated, added nucleotides are noted.
A model for target-directed purification of small RNA sorting in flies

Our data suggest that the sorting of hp-esiRNAs between Ago1 and Ago2 follows the previously established RISC assembly rules: hp-esiRNAs starting with U or A and containing mismatches in the central region of the small RNA duplex preferentially associate with Ago1, whereas those that begin with C or U and possessing a paired center are loaded into Ago2. We propose that hp-esiRNAs that begin with C are loaded into Ago2 by the RISC loading complex, but we do not yet know whether the bias toward C reflects a preference of the loading machinery, of Ago2, or of both. As the human homolog of Ago1 prefers to bind a 5'-U (Frank et al. 2010), we suspect that fly Ago1 intrinsically prefers small RNAs that start with U. Moreover, our results reinforce the view that in flies, small RNA production and Argonaute loading are not coupled. Just as miRNAs are made by Dicer-1 but can assort between Ago1 or Ago2 (Fürstemann et al. 2007), so hp-esiRNAs are made by Dicer-2 and can be loaded into either fly Argonaute protein (Czech et al. 2009).

The steady-state accumulation of individual small silencing RNA species in Ago1 and Ago2 reflects not only their initial sorting during the Argonaute-loading process, but also their subsequent purification by mRNA targeted tailing and trimming (Fig. 7): specific, single-stranded hp-esiRNA species, bound to Ago1, are destabilized when they bind to highly complementary targets. The result is that these hp-esiRNAs accumulate in Ago2, despite their initial loading into both Ago1 and Ago2.

The mechanism underlying this purification is similar to that which leads to loss of specific miRNAs upon the artificial expression of a fully complementary target RNA or antisense oligonucleotide, which triggers 3'-to-5' exonucleolytic trimming and 3' tailing of the small RNA (Ameres et al. 2010). Ago2-bound hp-esiRNAs are protected from this process because they bear a protective 2'-O-methyl modification. Why does high complementarity between a target RNA and an Ago1-, but not an Ago2-bound, small RNA trigger small RNA destruction? Ago1 is a poor enzyme compared to Ago2 (Fürstemann et al. 2007), and is therefore likely specialized to repress expression of partially complementary miRNAs through mechanisms that do not require endonucleolytic cleavage of the target (Iwasaki et al. 2009). Target-directed purification therefore might serve to ensure that highly complementary target RNAs are regulated only by Ago2, an enzyme adept at catalyzing multiple rounds of cleavage (Haley and Zamore 2004).

More puzzling still is why hp-esiRNAs are generated from partially complementary precursors that predispose them to load into Ago1 in the first place. While Ago2-bound hp-esiRNAs may generally repress high complementary target miRNAs, such as mus308, perhaps Ago1-bound hp-esiRNAs are required at a particular developmental stage or in a specific tissue to regulate less complementary target RNAs by a miRNA-like mechanism. Curiously, 5'-U esi-1.1, which is rare in S2 cells and fly heads, is more abundant than 5'-C

FIGURE 6. Hen1 stabilizes hp-esiRNAs. (A) The size distribution of genome-matching (top) and prefix-matching (bottom) sequences corresponding to esi-2.1 (left) or derived from the esi-1 locus (right) in the heads of hen1f00810 homozygous (red) or hen1f00810/CyO heterozygous (black) flies. The most abundant 3' nontemplated, added nucleotides are noted. (B) The distribution of 5' nucleotides among genome-matching hp-esiRNAs in hen1f00810 heterozygotes or homozygous mutant fly heads.

FIGURE 7. A model for the sorting of Drosophila hp-esiRNAs between Ago1 and Ago2 and their subsequent purification by target-directed tailing and trimming.
esi-1.1 in ovaries, raising the possibility that esi-1.1, bound to Ago1 instead of Ago2, serves to regulate one or more mRNAs specific to egg production (Supplemental Fig. S10).

MATERIALS AND METHODS

General methods

Synthetic small RNAs (Dharmacon) were PAGE purified and phosphorylated using T4 polynucleotide kinase (New England Biolabs); for sequences of synthetic RNAs see Supplemental Table S2. Small RNA annealing, preparation of lysates from Drosophila melanogaster 0–2 h embryos, and RISC loading reactions were as previously described (Haley et al. 2003).

UV crosslinking analysis

UV crosslinking was as described previously (Ameres et al. 2010). Briefly, RNA duplexes (20 nM) were incubated with lysate in a standard RNAi reaction at 25°C (Haley et al. 2003). At the indicated times, the reaction was stopped by incubation on ice and then irradiated with 254 nM UV light for 5 min using a Stratalinker (Stratagene) with the sample ~1 cm below the light bulbs. The photocrosslinked proteins were then resolved by 4%–20% gradient SDS-polyacrylamide gel electrophoresis (Criterion precast gels; BioRad), dried, and exposed to storage phosphor screens (Fuji). Signal intensities were determined using ImageGauge V4.22 (Fuji). Relative crosslinking signals were determined by normalizing to the maximal crosslinking of a 5’ U guide to Ago1 for miRNAs and to a 5’ C guide to Ago2 for siRNAs. Crosslinking of 5’ U miRNA to Ago1 reached saturation after 30 min and 5’ C siRNAs to Ago2 after ~15 min. Crosslinking of small RNAs to the respective Argonaute protein was normalized to total small RNA in the reaction as determined by 32P scintillation counting (LS6500 Multi-Purpose Scintillation Counter, Beckman Coulter).

Bioinformatic analyses

Small RNA libraries

Published small RNA libraries used in this study were Ago1 and Ago2 immunoprecipitated small RNAs (Czech et al. 2008), 19–24-nt small RNAs from S2 cells (Zhou et al. 2009), Ago1 immunoprecipitated small RNAs from Oregon R heads, 18–30-nt small RNAs from heads of flies heterozygous or homozygous for r2d21, ago2414, and Cs i R N A st oA g o 2 a f t e r 15 min. (Haley et al. 2003). At the indicated times, the reaction was stopped by incubation on ice and then irradiated with 254 nM UV light for 5 min using a Stratalinker (Stratagene) with the sample ~1 cm below the light bulbs. The photocrosslinked proteins were then resolved by 4%–20% gradient SDS-polyacrylamide gel electrophoresis (Criterion precast gels; BioRad), dried, and exposed to storage phosphor screens (Fuji). Signal intensities were determined using ImageGauge V4.22 (Fuji). Relative crosslinking signals were determined by normalizing to the maximal crosslinking of a 5’ U guide to Ago1 for miRNAs and to a 5’ C guide to Ago2 for siRNAs. Crosslinking of 5’ U miRNA to Ago1 reached saturation after 30 min and 5’ C siRNAs to Ago2 after ~15 min. Crosslinking of small RNAs to the respective Argonaute protein was normalized to total small RNA in the reaction as determined by 32P scintillation counting (LS6500 Multi-Purpose Scintillation Counter, Beckman Coulter).

Extraction of sequences

Genome-matching sequences were extracted as described (Ghildiyal et al. 2010). Prefix matching reads were determined as described (Ameres et al. 2010); sequences with ambiguous base calls were excluded from the analysis.

Filtering and normalization of immunoprecipitation data

To determine the relative abundance of hp-esiRNAs in Ago1 and Ago2, we divided the hp-esiRNA reads in each immunoprecipitated or oxidized sample by a normalizing factor, computed with a subset of small RNAs that were sequenced at least once in the respective data set, as well as in the total RNA sample. To avoid ambiguity, we required these small RNAs to be detectable only in the total RNA and in the appropriate immunoprecipitated/oxidized sample and map uniquely to one location in the genome. The normalizing factor was set to the ratio of the total reads of these small RNAs in the appropriate immunoprecipitated/oxidized sample and their total reads in the total RNA sample.

5’ Nucleotide analysis

To compute the first nucleotide composition, hp-esiRNAs were weighted by their abundance.

Statistical analysis

Pearson correlation analysis was performed using Prism V5.0b (GraphPad Software Inc.).

SUPPLEMENTAL MATERIAL

Supplemental material can be found at http://www.rnajournal.org.

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