Dicer-independent processing of short hairpin RNAs

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ABSTRACT

Short hairpin RNAs (shRNAs) are widely used to induce RNA interference (RNAi). We tested a variety of shRNAs that differed in stem length and terminal loop size and revealed strikingly different RNAi activities and shRNA-processing patterns. Interestingly, we identified a specific shRNA design that uses an alternative Dicer-independent processing pathway. Detailed analyses indicated that a short shRNA stem length is critical for avoiding Dicer processing and activation of the alternative processing route, in which the shRNA is incorporated into RISC and processed by the AGO2-mediated slicer activity. Such alternatively processed shRNAs (AgoshRNAs) yield only a single RNA strand that effectively induces RNAi, whereas conventional shRNA processing results in an siRNA duplex of which both strands can trigger RNAi. Both the processing and subsequent RNAi activity of these AgoshRNAs are thus mediated by the RISC-component AGO2. These results have important implications for the future design of more specific RNAi therapeutics.

INTRODUCTION

RNA interference (RNAi) is an evolutionarily conserved post-transcriptional gene silencing mechanism in eukaryotes that is triggered by double-stranded RNA (dsRNA) (1,2). The key players in the RNAi mechanism are small non-coding dsRNAs of 20–30 basepairs (bp) with a 3′ di-nucleotide overhang, a 5′ monophosphate and 3′ hydroxyl group (3,4). Several classes of small non-coding RNAs exist, including microRNAs (miRNAs) and small interfering RNAs (siRNAs). MiRNAs are transcribed in the nucleus as primary miRNAs (pri-miRNAs) by RNA polymerase (Pol II, occasionally Pol III) and are subsequently processed by the Microprocessor complex, comprising ribonuclease (RNase) III Drosha and subunit DGCR8, into ~70 nucleotide (nt) pre-miRNAs (5,6). The pre-miRNA associates with Exportin 5 that recognizes the 2-nt 3′ overhang and facilitates RNA export to the cytoplasm (7–9). The cytoplasmic complex that contains RNase III Dicer, TAR RNA-binding protein (TRBP) and protein kinase R (PKR) activator (PACT) processes pre-miRNAs into mature miRNA duplexes of 20–24 bp. The miRNA duplex associates with a member of the Argonaute protein family (AGO) within the precursor RNAi-induced silencing complex (pre-RISC) (10,11). Either miRNA strand can mediate gene silencing, but many miRNAs show asymmetry with preferential loading of one of the strand (guide) into RISC. Thermodynamic properties largely determine which strand of the duplex will be incorporated as guide into RISC (12,13). The remaining strand of the duplex (passenger) is cleaved and degraded. The mature RISC contains the single-stranded guide RNA annealed to a complementary target mRNA for post-transcriptional gene silencing (PTGS). The RNAi pathway can also be induced by artificial substrates, transiently by siRNAs that directly feed into RISC or stably by intracellularly expressed shRNA precursors that are processed by Dicer into siRNAs (14–16).

A key component of the RNAi pathway is the AGO protein. Human cells have four Ago genes (Ago1–4), and the siRNAs and miRNA templates are assorted among these AGO proteins (17,18). In worms and flies, the sorting is influenced by prior Dicer processing, the structure of the small RNA duplex (thermodynamic properties) and its terminal nucleotides (19–26), whereas in mammalian cells a strict system for small RNA sorting is lacking (17,27,28). Only the AGO2 protein is known to have ‘slicer’ activity, executed by a catalytically active RNaseH-like domain that cleaves the target mRNA (29). Endonucleolytic cleavage of the target mRNA occurs opposite nucleotide position 10 and 11 of the annealed siRNA guide strand. The mRNA cleavage products are released from RISC and interact with the 5′ → 3′ exonuclease XRN1, leading to decapping and subsequent mRNA degradation (30).

Most research on the description of the RNAi machinery and improvement of the RNAi knockdown efficiency
has focused on synthetic siRNAs or cellular miRNAs, but much less on shRNAs (31–36). The sequence and structure of shRNAs have been shown to influence the RNAi activity (16,37–44). However, specific sequences or elements within the shRNAs that determine the processing efficiency remain poorly defined. Recently, evidence has accumulated for non-canonical processing of miRNAs and shRNAs (45–50). Processing of the pre-miRNA-451 has been shown to occur by AGO2 instead of Dicer cleavage (45–47). The short 17 bp stem and 4 nt loop of miR-451 are the major determinants for this alternative processing route (45–47). Two groups previously described chemically synthesized shRNAs of minimal size that potently induced RNAi via mRNA cleavage, yet were not processed by Dicer in vitro and in vivo (34,51). The Siolas report suggested that the siRNA loop could be cleaved by a cellular endonuclease of unknown origin. The Dallas group suggested that AGO2 may be involved in the processing of shRNAs with short stem lengths of \( \leq 19 \) bp. Indirect evidence for AGO2 involvement was obtained by chemical modification and introduction of mismatches at and around the cleavage site, which resulted in decreased processing and knockdown activities (49).

In this study, we generated shRNA variants with different duplex lengths and loop sequences to test the effect on shRNA processing and RNAi activity. Reporter constructs that score guide and passenger strand activity revealed a nearly complete switch from the regular guide to the passenger strand for a certain shRNAs design. Further analyses confirmed an alternative shRNA-processing mechanism that is independent of Dicer, yet carries all hallmarks of AGO2-mediated processing. AGO2 co-immunoprecipitation studies revealed that these alternatively processed shRNAs are indeed incorporated into RISC and processed by AGO2. These findings indicate that the shRNA stem length is a critical factor for Dicer-independent processing. Importantly, the alternatively processed shRNAs induce an exquisite strand-specific RNAi-mediated gene knockdown compared with conventionally processed shRNAs.

**MATERIALS AND METHODS**

**Plasmids**

ShRNA expression plasmids targeting HIV-1 sequences pol47, pol9 and R/T5 were constructed as previously described (16,39,52). The RNA secondary structure of the shRNA transcripts was predicted by the mFold program (53). The firefly luciferase reporter plasmids were constructed by insertion in the 3′ untranslated region (3′UTR) of a 50–70-basepair fragment with the actual 19-nucleotide target in the centre. For this, we used the EcoRI and PstI sites of the pGL-3 plasmid (54). The luciferase reporters with the sense target sequences were described previously (52), whereas the reporters harbouring the antisense target sequences were newly constructed as described above. The wild-type AGO2 protein, the catalytic mutant (D597A) and the N domain mutant (F181A) were expressed from the pRESneo-Flag-haemagglutinin-AGO2 vector and were kindly provided by Y. Tomari (55).

**Cell culture**

Human embryonic kidney 293T adherent cells and HCT-116 cells were grown as monolayer in Dulbecco’s modified Eagle’s medium (Life Technologies, Invitrogen, Carlsbad, CA, USA) supplemented with 10% fetal calf serum (FCS), penicillin (100 U/ml) and streptomycin (100 \( \mu \)g/ml) in a humidified chamber at 37°C and 5% CO\( _2 \).

**DNA transfection**

Co-transfection of pGL-3 (Firefly luciferase reporter) with the shRNA vector was performed in 96-well format. Per well, \( 2 \times 10^4 \) 293T cells were seeded in 100 \( \mu \)l DMEM with 10% FCS without antibiotics. The next day, 25 ng pGL-3, 10 ng shRNA vector and 0.5 ng pRL (Renilla luciferase reporter) were transfected using 0.5 \( \mu \)l Lipofectamine 2000 according to the manufacturer’s instructions (Life Technologies). Cells were lysed 48 h post transfection to measure firefly and renilla luciferase activities using the Dual-Luciferase Reporter Assay System (Promega, Madison, WI, USA) according to the manufacturer’s instructions. The ratio between firefly and renilla luciferase activity was used for normalization of experimental variations such as differences in transfection efficiencies. Transfection experiments were corrected for between-session variations as described previously (56).

**siRNA detection by Northern blotting**

Northern blots were performed as previously described (57). Briefly, \( 1.5 \times 10^6 \) HCT-116 cells were transfected with equimolar quantities (5 \( \mu \)g) of shRNA constructs using Lipofectamine 2000. Total cellular RNA was extracted 2 days post-transfection with the mirVana miRNA isolation kit (Life Technologies, Ambion, Austin, TX, USA) according to the manufacturer’s protocol. The RNA concentration was measured using the Nanodrop 1000 (Thermo Fisher Scientific). For Northern blot analysis, 15 \( \mu \)g total RNA or 1 \( \mu \)g of AGO2-immunoprecipitated RNA samples were electrophoresed in a 15% denaturing polyacrylamide gel (precast Novex TBU gel, Life Technologies). RNA molecular weight marker (Life Technologies) was prepared according to the manufacturer’s protocol and run alongside the cellular RNA. To check for equal sample loading, ribosomal RNA was stained with 2 \( \mu \)g/ml ethidium bromide and visualized under UV light. The RNA in the gel was electro-transferred to a positively charged nylon membrane (Boehringer Mannheim, GmbH, Mannheim, Germany) and cross-linked to the membrane using UV light at a wavelength of 254 nm (1200 \( \mu \)J/cm\(^2 \)). Hybridizations were performed at 42°C with radiolabelled locked nucleic acid (LNA) oligonucleotides in 10 ml ULTRAhyb hybridization buffer (Life Technologies, Austin, TX, USA) according to the manufacturer’s instructions. LNA oligonucleotide probes were 5′-end labelled with the kinaseMax kit (Life Technologies) in the presence of 1 \( \mu \)l [\( \gamma \)-32P] ATP (0.37 MBq/\( \mu \)l, Perkin Elmer). To remove unincorporated nucleotides, the
probes were purified on Sephadex G-25 spin columns (Amersham Biosciences) according to the manufacturer’s protocol. We used the following oligonucleotides to detect the antisense strand of the siRNA (LNA positions underlined): 5'-ATGCGAGGAAGACGGGAG-3' (R/T5), 5'-GGTAGGGGCTGATGATAT-3' (pol47) and 5'-TGGCAGGAAGATGGCCAGT-3' (pol9). To detect the sense strand of the siRNA, the following oligonucleotides were used (LNA positions underlined): 5'-CTCGCGCTTCTTCTGCCCAT-3' (R/T5), 5'-ATTACTACGTGCCCTTACC-3' (pol47) and 5'-CTGCGGATCTTCTGTGCTA-3' (pol9). The signal was detected by autoradiography and quantified using a phosphorimager (Amersham Biosciences).

**AGO2 co-immunoprecipitation of small RNAs**

HCT-116 cells (5 × 10⁶ cells) were co-transfected with 5 µg Flag-tagged AGO2 plasmid and 20 µg of the various shRNA vectors. At 36 h post-transfection, cytoplasmic cell extracts were prepared by the treatment of cells on ice for 20 min with IsoB-NP-40 [10 mM Tris-HCl (pH 7.9), 150 mM NaCl, 1.5 mM MgCl₂, 1% NP-40] followed by a centrifugation at 12000 g for 10 min at 4°C. The supernatant was incubated with 75 µl of anti-FLAG M2 agarose beads (Sigma) with constant rotation overnight at 4°C. The beads were washed three times in NET-1 buffer [50 mM Tris-HCl (pH 7.5), 150 mM NaCl, 2.5% Tween 20]. Small RNAs associated with AGO2 were isolated by phenol chloroform extraction followed by DNase treatment using the TURBO DNA-free kit (Life Technologies). The RNAs were subsequently analysed on Northern blots (as described above) or cloned and sequenced.

**Cloning of small RNAs**

The purification and cloning strategy of small RNAs was previously described (58). Briefly, small RNAs were polyadenylated using the A-Plus™ Poly(A) Polymerase Tailing Kit (EPICENTRE Biotechnologies) according to the manufacturer’s protocol, and a 5’ adapter was ligated to the polyadenylated RNAs. Next, the 5’ adaptor-small RNA-poly(A)n molecule was reverse transcribed into cDNA followed by amplification of the cDNA by PCR. PCR products were gel-purified and subsequently used for TOPO-TA cloning (TOPO-TA cloning kit, Life Technologies). Positive clones were sequenced with the T7 or M13rp primer using the BigDye Terminator v1.1 Cycle Sequencing Kit (Applied Biosystems).

**Results**

**Reversal of strand selection for a particular shRNA design**

To study whether we could improve the shRNA design, we tested several variants of shPol47 (Figure 1) with the anti-HIV guide sequence encoded on the 3’ side of the hairpin duplex that targets a conserved sequence in the HIV-1 Pol gene. These shPol47 variants differ in length of the basepaired stem and terminal loop sequence. Several shRNA variants have been described in a previous study where we studied the impact of the terminal loop, varying in size and structure, on shRNA-induced gene silencing (39). We renamed the shRNAs according to the stem length and loop size, e.g. shPol47 with a stem length of 21 bp and a loop sequence of 5 nt was named 21/5. The 21/5A design that was originally proposed by Brummelkamp and co-workers is commonly used in many laboratories (16,52,59–62). We tested both the knockdown activity mediated by the 5’ and 3’ side of the shRNA duplex by co-transfection with luciferase reporter constructs encoding the respective targets. An irrelevant shRNA (shNef) served as negative control that resulted in maximal luciferase expression (set at 100%). Most shPol47 variants efficiently inhibited gene expression of the luciferase reporter with the sense HIV-target and exercised little activity on the antisense reporter, indicating that the 3’ strand of the shRNA is indeed selected as guide (Figure 2A). However, the 19/5 design showed a significant reduction in sense luciferase knockdown with a concomitant increase in antisense luciferase knockdown. Moderate activity on the antisense reporter was also observed for three other variants with a shortened 19 bp duplex (19/9 A, B and C), but not for shRNAs with 21 bp. Thus, the 19 bp duplex seems to result in a shift in guide strand selection from the 3’ to the 5’ side of the shRNA.

To investigate this phenomenon in more detail, Northern blot analyses were performed to analyse the processing of the shPol47 variants. We used LNA oligonucleotides to detect shRNA products derived from the 3’ and 5’ side (Figure 2B, upper and lower panel, respectively). An irrelevant shRNA (shGag) served as negative control. We observed 3’ side-derived siRNAs (~21 nt) for most shRNA variants, except for the 19/5 variant and to a lesser extent for the other 19/9 designs. This nearly complete loss of the 3’ signal (Figure 2B, top) correlated for construct 19/5 with reduced knockdown activity on the sense luciferase reporter (Figure 2A, top). A low level of regular ~21 nt fragments was detected for all constructs with the 5’ side probe, reflecting the passenger strand. Interestingly, a new and abundant RNA fragment of ~30 nt was observed for the 19/5 variant (marked * in Figure 2B, bottom), which coincides with the knockdown activity gained on the antisense luciferase reporter (Figure 2A, bottom). The size of this RNA product is not consistent with regular Dicer processing.

To study whether this phenomenon also applies to unrelated shRNAs with a completely different stem sequence, we tested several stem/loop variants of the validated shRT5 and shPol9 HIV-1 inhibitors for knockdown activity on the two reporters, including the special 19/5 design (Figure 3A). As expected, for most shRNAs, we observed more potent inhibition on the sense reporter (normal guide or 3’ side shRNA activity) than the antisense reporter (passenger or 5’ side shRNA activity). However, the activity of the 19/5 shRNA variants was shifted from 3’ towards the 5’ side, indicating a reversal of strand selection. Moreover, we again observed the loss of the regular 3’ side RNA fragment and appearance of a new ~30 nt 5’ side RNA fragment for the 19/5 shRNAs (Figure 3B). These ~30 nt RNA
fragments do not resemble regular Dicer-processed RNA products. They seem to consist of 19 nt from the 5' side of the shRNA, 5-nt loop and ~6 nt from the upper 3' side.

The hairpin stem length is critical for alternative shRNA processing

The shRNA mutant data indicate that variation in the stem and/or loop can induce alternative shRNA processing. An additional set of shRNAs was designed to critically test this idea. The stem of shRT5 was shortened or extended at the bottom of the hairpin (Figure 4, stem variants 15/5–23/5). The hairpin loop was varied between 3 and 8 nt (Figure 4, loop variants 19/3–19/8), and in addition, we reversed the 5-nt loop sequence between 3 and 8 nt (Figure 4, stem variants 15/5–23/5). The hairpin loop was varied extended at the bottom of the hairpin (Figure 4, stem variants 15/5–23/5). The hairpin loop was varied extended at the bottom of the hairpin (Figure 4, stem variants 15/5–23/5). The hairpin loop was varied extended at the bottom of the hairpin (Figure 4, stem variants 15/5–23/5).

The shRNAs of at least 20 bp do show the regular 3' side activity on the sense reporter (20/5–23/5). The loop size does clearly modulate the 5' side activity on the antisense reporter, with optimal activity for 19/3 and a gradual decrease towards larger loops. In fact, 19/7 and 19/8 seem to regain some regular 3' side activity at the expense of 5' side activity.

We next investigated the processing of these shRT5 variants by Northern blot analyses. We observed regular 3' strand siRNA production (~21 nt) for hairpins with a stem length of 20–23 bp, with the original 21/5 A design being the most optimal (Figure 6, upper, left panel). These shRNAs also showed some passenger 5' strand ~21 nt siRNA production (Figure 6, lower, left panel). Consistent with the luciferase knockdown data, we observed no 3' strand siRNA production for the 19/3–19/6 loop mutants, but a modest regain of regular siRNA production (~21 nt) was apparent for 19/7 and especially 19/8 (Figure 6, upper, right panel). Loss of regular 3' side product coincides with the appearance of the new ~30 nt product from the 5' side (Figure 6, lower, right panel). Note that the size of this new product gradually increases for the loop variants (19/3–19/8), demonstrating that the loop nucleotides are an integral part of this new processing product. The 17/5 and 18/5 variants also produced some 5' strand-derived RNAs of ~30 nt. The 20/5 and 21/5 A variants showed weak expression of the ~30 nt RNA fragment (Figure 6, lower, left panel). These shRNAs are therefore likely processed in different ways by the conventional and a novel processing route. The shRT5 variants of 15 and 16 bp did not show any 3' or 5' siRNA expression, consistent with the lack of knockdown activity.

These combined results indicate that an shRNA with minimal duplex length (19 > 18 > 17 bp) in combination with a small loop (3 > 4 > 5 > 6 > 7 nt) favours an alternative shRNA-processing route. The alternative processing route produces an RNA fragment that does not resemble conventional Dicer-processed shRNA products and most likely consists of 19 nt 5' side, the terminal loop and ~6–8 nt from the upper 3' side of the shRNA.

AGO2-mediated processing of 17–19 basepair shRNAs

The results described above indicate that the new shRNA-processing products of ~30 nt are active in the RNAi mechanism and are causing efficient knockdown of reporter gene expression, which implies that they end up in the mature RISC complex and associate with AGO2. Recent miRNA and shRNA studies implicated AGO2 in the processing step (45–50). To test this for our 19/5 shRNA design and to determine the exact nature of the processed RNA fragments, we performed immunoprecipitation and sequencing of the AGO2-associated small RNAs. HCT-116 cells were co-transfected with FLAG-tagged AGO2 plasmid and a vector encoding the shRT5 variants 19/5 or 21/5 A. At 36 h post transfection, cytoplasmic cell extracts were prepared and the AGO2 complexes were captured with an anti-FLAG M2 agarose resin. The RNA content of purified AGO2 complexes was extracted, cloned and subsequently...
Figure 2. Knockdown activity of the 3′/5′ strands of the shPol47 variants. (A) The knockdown activity of the 3′ and the 5′ strands of the different shRNAs was determined by co-transfection of a luciferase reporter encoding either the sense- or antisense-target sequence, respectively. 293T cells were co-transfected with 25 ng of the respective firefly luciferase reporter plasmid, 0.5 ng of renilla luciferase plasmid and 5 ng of the corresponding shRNA constructs. An irrelevant shRNA (shNef) served as negative control, which was set at 100% luciferase expression. (B) Processing of the 3′ strand (upper panel) and 5′ strand (lower panel) of shPol47 was analysed by Northern blot analysis. HCT-116 cells were transfected with 5 μg of the shRNA constructs. An irrelevant shRNA (shGag) was used as negative control. Northern blot analysis was performed on total cellular RNA. The RNA size marker (nt) is shown on the left. The regular 21 nt siRNA products and the new ~30 nt product are indicated (marked as *).

Figure 3. Knockdown activity of the 3′/5′ strands of several shRT5 and shPol9 variants. (A) The knockdown activity of the 3′ and the 5′ strands of the indicated shRNAs was determined by co-transfection of a luciferase reporter encoding the sense (white bars) or antisense target sequence (black bars), respectively, in 293T cells. See Figure 2A for details. (B) Processing of the 3′/5′ strands of the indicated shRNAs was analysed by Northern blot analysis. We used LNA oligonucleotides to detect the 3′ (upper panel) and 5′ strand (lower panel) of the siRNA. An irrelevant shRNA (shGag) was used as negative control. The regular 21 nt products are marked and the * indicates the ~30 nt RNA products.
Sequences of the small RNA sequences were prepared from cells expressing the different AGO2 constructs and shRT5 variant 19/5 or 21/5A. The AGO2 complexes were immunoprecipitated, and the small RNAs were isolated for Northern blot analysis. We first probed for the regular 3' side-derived shRNA products. Moderate ~21 nt 3' side siRNA and precursor shRNA signals were detected for 21/5A in association with the wt and mutant AGO2 proteins (Figure 7, upper panel). The highest amount of ~21 nt RNAs was detected in association with the catalytically deficient AGO2 mutant. For the 19/5 hairpin, neither precursor nor 3' side-processed siRNAs could be detected in association with the wt-, N- or cat-AGO2 protein. These results are consistent with the processing pattern previously observed for 19/5 shRNAs (Figures 2B, 3B and 6).

Using a probe to detect the 5' side of the shRNAs, we exclusively detected alternatively processed 33 nt RNA fragments for the 19/5 shRNA in association with wt- and N-AGO2 proteins (Figure 7, lower panel), whereas nearly no 33nt RNA was found in association with cat-AGO2. Notably, this product was present in whole cell extracts of cat-AGO2-transfected cells (data not shown). This observation indicates that AGO2 slicer activity is required for generation of the alternatively processed 33 nt RNA fragment. The weak 33 nt siRNA signal may represent processing by the endogenous wt AGO2 protein and subsequent uptake by the FLAG-tagged cat-AGO2 mutant. For the 21/5A construct, we detected the precursor shRNA, the alternatively processed 37 nt RNA fragment and the regular ~21 nt 3' side siRNAs in association with wt- and N-AGO2. This mixed profile of regular and alternative shRNA processing was previously observed for different 21/5A constructs (Figures 3B, 6 and 7). The catalytically dead AGO2 mutant contained little precursor 21/5A and 3' siRNAs of ~21 nt, whereas no 37 nt siRNAs were detected. This result confirms that AGO2 slicer activity is required for alternative shRNA processing. For unknown reasons, the catalytic AGO2 seems to favour the loading of ~21 nt 3' siRNAs (Figure 7, upper panel) derived from the 21/5A over 5' siRNAs. Notably, mutation of the unwinding domain did not hinder alternative processing for both 19/5 and 21/5A shRNAs (Figure 7, lower panel).

**DISCUSSION**

The sequence and structure of natural miRNAs are important features that influence their processing and RNAi activity. Exact features in man-made shRNAs that determine the processing efficiency have not been examined in detail. The shRNA loop sequence is likely to have an effect on Dicer-mediated recognition and processing (37,39,63,64), but also the stem length is an important determinant for RNAi activity (51,65,66). We tested a variety of shRNAs and identified a specific shRNA design with a minimized stem length and a small loop that triggers an alternative processing route to form a new characteristic RNA fragment of 33 nt. The new cleavage event half-way the 3' side of the shRNA duplex suggested a role for AGO2, which cleaves target mRNA or the passenger strand of an siRNA duplex between nt 10 and 15/5–23/5. In addition, shRNA terminal loops ranging in size from 3 to 8 nt were designed (19/3–19/8) and the loop sequence was reversed (19/9–19/15). The original shRT5 21/5A variant was also included.
Sequencing of AGO2-associated small RNAs confirmed this exact cleavage event. Such non-canonical processing has recently been reported for miR-451 and has been suggested for several other miRNAs and shRNAs (45–50). It would be interesting to compare the processing kinetics of AgoshRNAs versus conventional shRNAs and the underlying binding activities for Dicer and AGO2.

Production of the 33 nt RNA fragments was abolished in the presence of a catalytic AGO2 mutant, demonstrating that processing is mediated by AGO2 slicer activity. The minimized shRNAs of 17–19 bp are likely too small to be recognized by Dicer (34) and end up in AGO2 for alternative processing and subsequent RNAi silencing of an appropriate mRNA (Figure 8, middle panel). The new shRNA design that elicits this non-canonical processing pathway is termed AgoshRNA. We confirmed this specific AgoshRNA design for other shRNAs, demonstrating the universal value of the new shRNA design (19 bp stem and a small loop). Notably, this AgoshRNA processing yields only a single RNAi-active RNA fragment, whereas the conventional shRNA design potentially yields two active siRNA strands of ~21 nt (Figure 8). This AgoshRNA feature is important to restrict RNAi-induced off target effects via the passenger strand of the siRNA.

The frequently used shRNA 21/5A design (16) exhibited a mixed processing profile according to the conventional Dicer and the alternative AGO2 mechanism. This may relate to the fact that the upper 2 bp of this shRNA design are relatively weak (U-G and U-A). Thus, this hairpin may exist in solution as a mixture of structures with 19, 20 and 21 bp. The conformers with 19 bp stem will be alternatively processed, whereas conformers with an extended stem of 20 or 21 bp will be processed conventionally. A recent report by Yang et al. (48) showed that a conventional hairpin of 21 bp with similar design was alternatively processed in a Dicer-independent manner.

Figure 5. Knockdown activity of the shRT5 variants. RNAi activity of the 3′ side (lower panel) and 5′ side (upper panel) of the shRT5 was determined by co-transfection of a luciferase reporter encoding the sense or antisense target sequence, respectively, in 293T cells. See Figure 2A for details.
However, this study only addressed the required specifics to induce alternative processing for pre-miRNAs and not for shRNAs.

We described the specific AgoshRNA design that bypasses the Dicer endonuclease and instead uses AGO2 for processing. During processing of the AgoshRNA, AGO2 cleaves in the 3' strand of the hairpin and the short 3' end product is likely degraded. The remaining 33 nt RNA is used as a guide to target the complementary mRNA for cleavage. The determinants for AGO2 loading of the unprocessed AgoshRNA hairpin are the stem lengths and loop sizes. Our results further support the notion that RISC loading does not require prior Dicer-mediated processing of the RNAi inducer (17, 68).

The loop size of the AgoshRNA molecules has a modest influence on its activity with higher knockdown efficiencies observed for smaller loops. Increasing the loop size from 3 to 7 or 8 nt caused a partial return to regular Dicer processing as detected by Northern blot analysis, Figure 6.

Table 1. AGO2-associated small RNAs from transfected HCT-116 cells

<table>
<thead>
<tr>
<th>AGO2 product</th>
<th>shRT5 19/5</th>
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<tr>
<td>5' 21 nt siRNA</td>
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<td>3' 21 nt siRNA</td>
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<tr>
<td>~30 nt siRNA</td>
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<td>1x (37 nt)</td>
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<td>miRNA</td>
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<td>tRNA</td>
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<tr>
<td>Other RNAs (rRNA, mitRNA)</td>
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*A total of 48 clones were sequenced.

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The loop size of the AgoshRNA molecules has a modest influence on its activity with higher knockdown efficiencies observed for smaller loops. Increasing the loop size from 3 to 7 or 8 nt caused a partial return to regular Dicer processing as detected by Northern blot analysis,
suggesting that the two processing routes are in competition. We hypothesize that a large loop (e.g. >7 nt) sterically hinders the efficient uptake of AgoshRNA by AGO2 (Figure 8, middle panel). Consistent with our findings that increasing the loop size of the 19 bp shRNAs can cause a shift from alternative to conventional processing, pre-miRNAs appear to be more efficiently processed by Dicer when they contain larger loops (69).

It seems that alternatively processed AgoshRNAs can be more active in RNAi knockdown experiments than conventional shRNAs. More potent target knockdown was observed in the luciferase reporter assays with the 17–19 bp minimized AgoshRNA compared with the 21 bp regular shRNAs, but a direct comparison remains difficult, as different guide strands are generated that are probed on different reporter constructs. The design of such optimized AgoshRNA therapeutics may allow one to reduce the RNA dosage, thus reducing the chance of adverse effects, e.g. due to off-targeting (69). Increased activity may be related to the production of only a single guide strand, which will also reduce the chance of off-target effects.

AgoshRNAs may have additional theoretical benefits as AgoshRNAs form smaller hairpin duplexes than regular shRNAs. The former may exhibit a better safety profile concerning activation of the dsRNA-induced protein kinase R (PKR)/interferon pathways (71). Because AgoshRNAs do not mature via Dicer, they also do not compete with this aspect of miRNA biogenesis. AGO2-mediated processing may also yield more distinct RNA molecules as Dicer creates imprecise ends (72). Finally, AgoshRNAs are attractive molecules to silence target genes in Dicer-compromised cells, and they represent the only option in Dicer-deficient cells, which relates to a growing number of human diseases and cancers (73–75) and monocytes that lack Dicer expression (76).

In conclusion, we describe a specific shRNA design that yields only a single active strand (guide), as the other strand (passenger) is destroyed by AGO2 cleavage. This feature is an important property to restrict off-target effects induced by the passenger strand. Experiments are ongoing to test whether these AgoshRNAs are safer than conventional shRNAs and whether conversion of the
regular loop sequences into anti-HIV sequences will generate more potent and possibly escape-proof antivirals.

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