Antisense morpholino targeting just upstream from a poly(A) tail junction of maternal mRNA removes the tail and inhibits translation

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ABSTRACT
Gene downregulation by antisense morpholino oligonucleotides (MOs) is achieved by either hybridization around the translation initiation codon or by targeting the splice donor site. In the present study, an antisense MO method is introduced that uses a 25-mer MO against a region at least 40-nt upstream from a poly(A) tail junction in the 3′-untranslated region (UTR) of maternal mRNA. The MO removed the poly(A) tail and blocked zebrafish cdk9 (zcdk9) mRNA translation, showing functional mimicry between miRNA and MO. A PCR-based assay revealed MO-mediated specific poly(A) tail removal of zebrafish mRNAs, including those for cyclin B1, cyclin B2 and tbp. The MO activity targeting cyclins A and B mRNAs was validated in unfertilized starfish oocytes and eggs. The MO removed the elongated poly(A) tail from maternal matured mRNA. This antisense method introduces a new application for the targeted downregulation of maternal mRNAs in animal oocytes, eggs and early embryos.

INTRODUCTION
The antisense method of gene downregulation classically uses single-stranded (ss) DNA to form a DNA–RNA duplex through complementary base pairing, leading to RNase H-mediated cleavage of the target mRNA in vivo (1,2). However, endonucleases exist that efficiently digest ssDNAs, thereby decreasing their antisense activities. Morpholino oligonucleotides (MOs) are frequently used because they are endonuclease resistant (3). In contrast to antisense DNAs that cause target mRNA digestion, MOs do not mediate the RNase H-dependent digestion of mRNAs (3–5). Duplex formation between MOs and mRNA prevents translation through MO hybridization near the mRNA translation initiation codon and disrupts correct splicing by targeting the splice donor site (4,5).

The activation of gene expression coincides with transcriptional activation; gene activation is closely correlated with an increased accumulation of mRNA. However, specialized cells, such as oocytes and neural cells, have a transcription-independent mechanism by which the expression of dormant mRNAs is strictly controlled by specific RNA-binding proteins (6,7). In this mechanism, cytoplasmic elongation of the poly(A) tail leads to translational stimulation of maternal mRNAs in Xenopus oocytes and early embryos (6,7).

This article describes a new method for gene downregulation. The availability and specificity were confirmed by targeting the maternal mRNAs of zebrafish cdk9, tbp, cyclin B1, cyclin B2, as well as starfish cyclins A and B. The key features of this method are as follows: (i) duplex formation of the MO with the mRNA 3′-untranslated region (UTR) not only prevents poly(A) tail elongation, but also induces deadenylation like miRNA (8) or removal of the entire tail and (ii) this method blocks target mRNA translation.

MATERIALS AND METHODS
Synthesis
DNA and morpholinos were synthesized by Operon Biotechnologies and Gene Tools, respectively.

Embryos, oocytes and eggs
All zebrafish and embryos were maintained at 28°C and staged as described previously (9). Immature oocytes or

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unfertilized mature eggs of the starfish *Asterina pectinifera* were prepared as described (10). Immature oocytes were treated with 1 μM 1-methyladenine (1-MeAde) to induce meiotic maturation at 23°C in artificial seawater (Jamarin Laboratory).

**Microinjection of MO**

Zebrafish wild-type embryos were injected at the one- or two-cell embryo stage with 2.5 pmol of MO. The MOs used in this study are shown in Supplementary Table S1. When a mixture of two MOs was tested, 1.25 pmol of each MO was injected into the embryo.

Starfish immature oocytes or unfertilized mature eggs were injected with 20 fmol of morpholino antisense oligonucleotides against either starfish *cyclin A* or *B* (Supplementary Table S1). Microinjection was performed as described (11). In Figure 6E, a mixture of sfccya MO (10 fmol) and sfccyb MO (10 fmol) was used.

**Preparation of extracts of embryos, oocytes and eggs**

Ten zebrafish embryos were frozen in liquid nitrogen and thawed in 200 μl of RIPA buffer [150 mM NaCl, 1% NP-40, 0.5% deoxycholate, 0.1% sodium dodecyl sulfate, 50 mM Tris–HCl (pH 8.0)]. After thorough sonic- ation, supernatants (100 μl) were collected by centrifuga- tion at 12000g for 1 min and mixed with 40 μl of 4× Laemmli sample buffer. For western blot analysis, 14 μl of the sample, corresponding to a half-embryo, were analyzed.

Four starfish oocytes were recovered in 5 μl of seawater and 7 μl of 2× Laemmli sample buffer were added for western blot analysis (12).

**Purification of total RNA from embryos and oocytes**

Total RNA was extracted from 10 zebrafish embryos or 20 starfish oocytes with Sepasol-RNA I super (Nacalai Tesque).

**Poly(A) test assays**

The PAT assays were performed essentially as described (13), except for minor modifications. Total RNA (300 ng) was incubated at 65°C for 5 min in the presence of a mixture (total 20 ng) of phosphorylated oligo (dT) primers, which were 12- to 18-mer poly (dT) primers. After incubation for 1 h at 42°C with T4 DNA ligase (350 U) (Takara), the samples were further incubated at 12°C for 1 h in the presence of 200 ng (dT)3-2-anchor primer (5'-GGAGCTCCCG GGGCCTTGGTT TTTTTT TTTT-3') to generate the 3'-end of the poly (dT) primer, followed by incubation with SuperScript III reverse transcriptase (200 U) (GE Healthcare) for 1 h to make PAT complementary DNAs (cDNAs). Finally, the products of PCR using a gene-specific primer and the (dT)12-anchor primer were subjected to 2.2% agarose gel electrophoresis and visualized by ethidium bromide staining. The bands were cut out, the DNA fragments were eluted and the fragments were cloned into a pMD20-T vector. After blue–white selection of clones on agar plates containing ampicillin, plasmid DNA was recovered from each clone. Insert regions were confirmed by restriction enzyme digestion. More than 10 clones of each sample were selected for further DNA sequencing analysis.

**RT–PCR**

RT–PCR was performed with SuperScript III transcriptase (200 U). Random and oligo (dT) primers (Invitrogen) were used as reverse primers. The gene-specific primer sets (forward and reverse primers) used for RT–PCR are shown in Supplementary Table S3.

**Western blot analysis**

To generate anti-zCdk9 antibodies, the recombinant protein of zCdk9 was expressed in bacteria (*Escherichia coli*) and fractionated by disk preparative electrophoresis. A rabbit was immunized with the Cdk9 protein, according to the standard protocol of Operon Biotechnologies. The polyclonal anti-zCdk9 antibody was affinity purified from the immunoserum as described (14). H-169, which reacted with human Cdk9 and anti-actin (clone C4) antibodies, was purchased from Santa Cruz Biotechnology and Chemicon, respectively. Immunoblotting was performed as described previously (15). Blots were developed with the ECL system (GE Healthcare).

In the case of starfish, the primary antibodies used were anti-starfish cyclin B (10) and anti-MAPK (Upstate), and the secondary antibodies were HRP-conjugated anti-rabbit IgG (GE Healthcare), anti-mouse IgG (Dako) and AP-conjugated anti-rabbit IgG (Dako). Immunocomplexes were detected by the BCIP/NBT phosphatase substrate system (KPL) or ECL Plus (GE Healthcare) and were visualized with LAS4000 IR multi color (Fuji Film).

**Determination of the 3'end sequence of mRNA**

We performed reactions according to the manufacturer’s manual supplied with the small RNA Cloning Kit (Takara) with minor modifications. Three hundred nanograms of total RNA isolated from embryos were treated with bacterial alkaline phosphatase (BAP). A biotinylated RNA/DNA 3' adaptor was ligated to the 3'-end of BAP-treated RNA. Streptavidin-conjugated magnetic beads were used to collect the adaptor-ligated RNA. After washing the beads, reverse transcription reactions were performed with PCR-R & RT-primer (5'-GTCTCTTAGC C TGCAGGATCG ATG-3'). Synthesized cDNAs were recovered from the beads by alkaline treatment, followed by PCR with zcdk9 PAT primer and PCR-R & RT-primer. The PCR products were subjected to agarose gel electrophoresis, and DNA bands were visualized by ethidium bromide staining. The bands were cut out, the DNA fragments were eluted and the fragments were cloned into a pMD20-T vector. After blue–white selection of clones on agar plates containing ampicillin, plasmid DNA was recovered from each clone. Insert regions were confirmed by restriction enzyme digestion. More than 10 clones of each sample were selected for further DNA sequencing analysis.

**Messenger RNA synthesis and in vitro translation**

To generate a plasmid for starfish *cyclin B* [pcDNA3.3sFCycB(Xhol)], a fragment of the starfish *cyclin B* mRNA-coding region was PCR-amplified with the
RESULTS

Injection of MO targeting the 3′-UTR terminus of zcdk9 mRNA into zebrafish early embryos inhibits polyadenylation and subsequent translation

While studying the mechanistic aspects of zebrafish kinase cdk9 mRNA expression at the midblastula transition (17), we noticed that elongation of the poly(A) tail plays a critical role in translation stimulation. To address this finding, we performed an antisense experiment with the low-toxicity nucleoside analogs, MOs (3). We postulated that duplex formation between an MO and the 3′-UTR of mRNA would prevent poly(A) tail elongation. In this study, a nucleotide at the junction of the poly(A) tail is defined as −1, and numbers are assigned as indicated in each figure. The junction information was obtained from the NCBI nucleotide database.

We prepared cdk9 MO consisting of 25 bases that were completely complementary to the terminal sequence of the zcdk9 mRNA 3′-UTR (Figure 1A). To examine whether MOs against the 3′-UTR affect polyadenylation, a PAT assay (13) was performed, in which the PCR products reflect the poly(A) tail lengths present on a specific mRNA. When injected into fertilized embryos, cdk9 MO markedly reduced the production of slowly migrating bands compared to that of untreated embryos (WT), indicating that cdk9 MO inhibited polyadenylation (Figure 1B). The cdk9 MO also strongly reduced the amount of cdk9 PCR product, without affecting the products of tbp and cyclin B1. This result was specific for cdk9 MO, because embryos treated with cdk9m MO carrying 5-base mismatches acted no differently from untreated embryos (Figure 1B). These results suggest that cdk9 MO specifically affects cdk9 mRNA.

We next assessed the effect of MOs at the protein level. A western blot assay was performed with two independent antibodies, including our original anti-zCdk9 antibody against zebrafish Cdk9 and the commercially available H-169 antibody against human Cdk9. Anti-actin antibodies were used as a control. The cdk9 MO did not affect zCdk9 accumulation in 3 hpf embryos, but blocked the zCdk9 increment that began at 4 h after fertilization (4 hpf) (Figure 1C). In contrast to cdk9 MO, there was no detectable effect of cdk9m MO on zcdk9 translation. We compared this method with the traditional method targeting the translational start site. Figure 1D shows that this method targeting the 3′-UTR of mRNA works as well as the traditional antisense method. Taken together, these results indicate that the cdk9 MO injection inhibited the translation of zcdk9 mRNA during early development.

Since the levels of cdk9 products in the PAT assay were decreased by the injection of cdk9 MO, we next investigated the effect of cdk9 MO using RT–PCR. Reverse transcription was performed with either a random primer or oligo-dT primer, followed by PCR amplification of the zcdk9-coding region. To confirm an experimental time course, we examined the expression of the zygotic transcription marker gene, bhlk, which was first detected at 3 hpf and began showing mRNA accumulation at 4 hpf (Figure 1E and F) (18). Thus, zygotic transcription appeared to occur at around 3 hpf, corresponding approximately to the beginning of cleavage cycle 10. This result was consistent with previous reports (17–19).

RT–PCR with the random primer yielded the same amount of zcdk9 products in the RNA of all embryos, regardless of cdk9 MO injection (Figure 1E), which indicates that cdk9 MO did not decrease the amount of zcdk9 mRNA. In contrast, the oligo-dT primer showed a low-level amplification of zcdk9 when cdk9 MO-injected embryos were tested (Figure 1F). The oligo-dT primer hybridized with the poly(A) tail, such that the short tail consisting of a few adenosines was insufficient for hybridization and resulted in low amplification. Therefore, the low-level amplification may have been due to poly(A) tail shortening, caused by deadenylation activity (see below). These results indicate that the injection of cdk9 MO into embryos induces deadenylation rather than mRNA degradation.

The active region for polyadenylation by MO exists at 40 nt from the terminus of the zcdk9 3′-UTR

To address whether the 3′-UTR terminus plays an important role in MO-mediated repression, three antisense MOs were created, including MO-2 (−26 to −50), MO-3 (−51 to −75) and MO-4 (−76 to −100) (Supplementary Figure S1). The PAT assay and western blot analysis indicated that, of the five MOs listed in Supplementary Figure S1, only cdk9 MO exerted an inhibitory effect on both polyadenylation and translation. This finding suggests that the most terminal portion of the 3′-UTR is critical for inhibition. To verify this result, additional antisense MOs (Figure 2) were created, namely MO-5 (−6 to −30), MO-6 (−11 to −35) and MO-7 (−16 to −40), which showed inhibitory effects on both polyadenylation and translation that were greater than or similar to those of cdk9 MO. A modest effect on both polyadenylation and translation was apparent when MO-8 (−21 to −45) was examined (Figure 2B and C). These results indicate that
the terminal 40 nt of the 3'-UTR act as the active region for MO-mediated inhibition.

Antisense MO removes the poly(A) tail from zcdk9 mRNA

As shown above, duplex formation of MO with the mRNA 3'-UTR terminus prevents proper polyadenylation. To determine precisely how many adenosine residues remain in the poly(A) tail after injection of MO, we performed DNA sequence analysis of the 3'-terminal region of MO-injected embryos. PCR products were analyzed on a 2.2% agarose gel. Right side, length markers in bases. (C) Specific translation inhibition of zcdk9 mRNA by cdk9 MO, but not cdk9m MO. Embryos were collected at the indicated times. Western blotting was performed with extracts from untreated (lanes 1, 4, 7 and 10), cdk9 MO-injected (lanes 2, 5, 8 and 11) and cdk9m MO-injected embryos (lanes 3, 6, 9 and 12) by using the antibodies indicated. HeLa cell nuclear extracts (NEs) (lane 13) served as a control. Blots were probed with H-169, anti-zCdk9 and anti-actin. (D) The cdk9m MO (lanes 1 and 2), cdk9 MO (lanes 3 and 4) and 5'-cdk9 MO targeting a region around the first codon (lanes 5 and 6)-injected embryos were collected at 3 and 5 hpf. The extracts were subjected to western blotting by using antibodies indicated. (E and F) Reverse transcription using random (E) or oligo-dT primers (F), followed by PCR with primer sets for cdk9, biklf, tpb and actin, and analysis on a 2.2% agarose gel. Embryos were collected at the indicated times. Total RNA was extracted from untreated (WT: lanes 1–3), cdk9 MO-injected (cdk9 MO: lanes 4–6) and cdk9m MO-injected embryos (cdk9m MO: lanes 7–9).
upstream relative to the MO-mRNA hybrid position. This tendency of the 3′-end to move upstream was seen with MO, MO-5, MO-6, MO-7 and MO-8. This finding suggests that the hybridization position between MO and mRNA affects the 3′-end determination of each mRNA. This effect may be due to deadenylase and exonuclease activity or to the action of an endonuclease recognizing the hybrid.

Investigation of the effects of MO on four maternal mRNAs

To validate the effect of MO on polyadenylation, three kinds of gene-specific MOs were created, including cyclin B1 MO, cyclin B2 MO and tub MO, in addition to cdk9 MO (Figure 4). To determine the MO specificity, the total RNA derived from 5 hpf embryos was examined, and a PAT assay was performed (Figure 4B and C). We tested a mixture of cdk9 MO plus cyclin B2 MO (Figure 4A and B). Zebrafish has the cyclin B orthologs B1 and B2, and their 3′-UTRs exhibit sequence similarity; indeed, there were 10-base homologies between cyclin B1 MO and cyclin B2 MO (Figure 4A). Cyclin B1 MO and cyclin B2 MO each affected its own target without oligonucleotides interference (Figure 4B). We also observed the specific action of MO when cdk9 MO and tub MO were examined (Figure 4C). The effects of MO on the polyadenylation of mRNAs were confirmed by using RNA samples collected from 2 to 5 hpf embryos (Supplementary Figure S2A and S2B). In addition, each MO binds to a target 3′-UTR in a specific manner (Supplementary Figure S3). These results indicate that antisense MOs targeting the 3′-UTR of mRNAs affect polyadenylation in early zebrafish embryos in a specific manner.

MO antisense method targeting the 3′-UTR of mRNAs in starfish oocytes: sfccyB MO downregulates expression of cyclin B maternal mRNA in a specific manner

Recently, we observed that the poly(A) tail elongation of cyclins A and B mRNAs during meiotic maturation of oocytes is induced by 1-MeAde (20). Therefore, we verified our method further in starfish. By using the
protocol described in Figure 5A, immature oocytes were injected with sfycB MO and treated with 1-MeAde. We prepared sfycB MO that has a complementary sequence in its 3'-UTR terminus for starfish cyclin B mRNA (Figure 5B), and used cdk9m MO as a control. After meiosis II (MII), total RNA was isolated from oocytes (Figure 5A) and a PAT assay was performed to determine the effect of MO on poly(A) tail elongation. As seen in Figure 5C, 1-MeAde-induced poly(A) tail elongation (≈100 residues) of both maternal cyclins A and B mRNAs (lanes 2 and 6), and sfycB MO injection inhibited mRNA polyadenylation of cyclin B (lane 7), but not cyclin A (lane 3). Control MO injection did not affect the polyadenylation of either cyclin A or B mRNAs (lanes 4 and 8). These results indicate that antisense MOs targeting the 3'-UTR of maternal mRNA specifically inhibit polyadenylation in starfish oocytes.

To determine the effect of MO on translation, western blotting was performed with antibodies against the cyclin B protein and MAPK. The 1-MeAde treatment of immature oocytes led to activation of MAPK and cyc-lic production/degradation of the cyclin B protein (Figure 5F) (10,21). Disappearance of the cyclin B protein at 60 min after 1-MeAde addition and its

Figure 3. Determination of the mRNA 3'-UTR end. (A) Schematic drawings indicate the method employed to determine the mRNA 3'-UTR end. (B) Experiments were performed with the total RNA used in Figure 2B. PCR products were visualized by ethidium bromide staining. (C) Results of DNA sequencing analysis of each cDNA derived from MO-injected embryos. Five clones were selected and the DNA sequences of their 3'-termini were aligned. A vertical line indicates a poly(A) tail junction of mRNA. The junction information was obtained from the NCBI nucleotide database (NM_212591.1).
Appearance at 70 min indicated that the M phase of MII began ~70 min after 1-MeAde addition (Figure 5F). Furthermore, re-disappearance of cyclin B protein at 90 min indicated an exit from M phase. Importantly, injection of sfycB MO blocked accumulation of the cyclin B protein at 70 and 80 min (Figure 5F), but the injection of control cdk9m MO did not (data not shown; Figure 5F), which indicates a specific inhibitory effect of sfycB MO on the translation of maternal cyclin B mRNA during MII.

To investigate whether sfycB MO targets cyclin B mRNA in a specific manner, a rescue experiment was performed with in vitro synthesized capped mRNAs (Figure 5D). The zebrafish cyclin B1 3'-UTR was employed to support the cell cycle-dependent translation of starfish cyclin B. The sequence of sfycB MO did not match a complementary sequence within the zebrafish cyclin B1 3'-UTR, suggesting that sfycB MO cannot make a duplex with zebrafish cyclin B1 3'-UTR. Capped starfish cyclin B mRNA lacking the 3'-UTR (Figure 5D) also was prepared. The synthetic individual RNAs were coinjected with sfycB MO into immature oocytes, and maturation was induced by the addition of 1-MeAde. Western blot analysis showed that the cyclin B proteins were translated efficiently from both synthetic RNAs in vitro (Figure 5E). In oocytes, however, the two RNAs exhibited different results. When chimeric mRNA was injected, a significant increase in cyclin B protein accumulation was observed that peaked 70 min after 1-MeAde addition, which was 10 min faster than that of untreated oocytes (Figure 5F). This 10-min difference may be due to the difference in the 3'-UTR between endogenous and injected cyclin B mRNAs. Regardless, oocytes produced the cyclin B protein efficiently from the chimeric RNA carrying the zebrafish cyclin B1 3'-UTR at MII. In contrast, the cyclin B mRNA without the 3'-UTR showed a low level of translation compared to untreated oocytes (Figure 5F), indicating that the 3'-UTR is required for proper cyclin B mRNA translation at MII. These results indicate that the zebrafish cyclin B1 3'-UTR can partly complement the function of the starfish cyclin B 3'-UTR in regulating cyclin B mRNA translation during oocyte maturation.

Taken together, these results demonstrate that sfycB MO downregulates expression of cyclin B maternal mRNA in a specific manner.

**Duplex formation just upstream of the poly(A) tail induces poly(A) tail deadenylation in mature starfish oocytes**

We next examined whether the antisense MO targeting the 3'-UTR of mRNA stimulated deadenylation activity. The protocol shown in Figure 6A was used, because it allowed the injection of MO into pronucleus (PN) stage eggs, in which the cell cycle is arrested at the G1 phase after the completion of MII. Elongation of the poly(A) tail in cyclins A and B mRNAs occurs during meiotic maturation, and the elongated tails are maintained in arrested oocytes (20).

The sfycA MO was used against the starfish cyclin A 3'-UTR in addition to sfycB MO (Figure 6B). Eggs (PN210) were collected at 90 min after injection of MO into arrested mature eggs (PN120). The total RNA isolation from the oocytes (Figure 6A) was used in a PAT assay. Both cyclins A and B mRNAs in mature oocytes had longer poly(A) tails than those of mRNAs in immature oocytes (Figure 6C; lanes 1 and 2), indicating that meiotic maturation induced the poly(A) tail elongation of mRNAs, as we reported previously (20). Injection of sfycA MO reduced the amount of PCR products from cyclin A, but not from cyclin B, and sfycB MO affected only PCR products from cyclin B (Figure 6C). In contrast, the control cdk9m MO had no effect on either PCR product. These results indicate that a specific duplex formation upstream of the poly(A) tail leads to poly(A) tail shortening by stimulating deadenylation activity.

Next, we examined whether maternal cyclins A and B mRNAs carrying shortened poly(A) tails could stimulate...
translation. Eggs were treated with the MAPK inhibitor U0126, which stimulates translation of cyclins A and B mRNAs (Figure 6D) (20). U0126 inhibited MAPK and stimulated translation in normal eggs (Figure 6E). In contrast, eggs that were injected with a mixture of MOs against cyclins A and B showed very weak stimulation of translation (compare lanes 3 and 4 in Figure 6E), indicating that MOs targeting the 3′-UTR of mRNAs repress the expression of maternal mRNAs.

DISCUSSION

We report the MO-mediated repression of maternal mRNAs in zebrafish early embryos and starfish oocytes and eggs. This repression was accomplished by the inhibitions of polyadenylation and translation; however, we do not know whether the same mechanism operates in zebrafish and starfish. Duplex formation of MO with mRNAs at the junction of the poly(A) tail is thought to be necessary for MO-mediated polyadenylation inhibition (Figure 2 and Supplementary Figure S1). Notably, cdk9 MO-2, which possibly hybridizes with the region from 26 to 50 bases upstream from the end of the zcdk9 3′-UTR, had no effect on the poly(A) tail (Figures 2 and 3). This result suggests that the responsible cis-element(s) exist within 25 bases from the 3′-UTR terminus.

Twenty-five bases of zcdk9, cyclins B1 and B2 mRNAs and 30 bases of tbp mRNA have a canonical
Polyadenylation signal (AAUAAA) that binds 10 cleavage-polyadenylation specificity factor (CPSF), a component of the cytoplasmic polyadenylation RNP complex (7). In *Xenopus* oocytes, the RNP complex consists of cytoplasmic poly(A) ribonuclease (PARN) and the cytoplasmic poly(A) polymerase (Gld-2), in addition to CPSF and cytoplasmic polyadenylation factors, including CPEB, Pumilio and Musashi (7). Therefore, it is plausible that hybridization of MO to the junction may prevent proper binding of CPSF to the polyadenylation signal, which possibly affects RNP complex function. This process may lead to perturbation of the balance between Gld-2 and PARN activities, resulting in poly(A) tail shortening (22).

Hybridization between MO and mRNA led to removal of the 3'-terminal region and the entire poly(A) tail from mRNA in zebrafish embryos. A few nucleotides at the 3' region remained just downstream from the hybrid position. Further downstream, nucleotides, including the poly(A) tail, were removed from the mRNA (Figure 3). This removal may be caused by a deadenylase-dependent cutting of the poly(A) tail, followed by exonuclease action to remove several 3'-terminal nucleotides. Alternatively, hybridization may stimulate the activity of a certain endonuclease, leading to the digestion of the mRNA at the region just downstream of the hybrid.

There are three important similarities between miRNA and MO behaviors: a target region is located in the 3'-UTR, deadenylation or poly(A) tail removal is induced and translation inhibition occurs (8). It will be of considerable interest to determine the protein components associated with MO-mediated inhibition.

Finally, this method using MO is limited to maternal mRNAs and is not a new tool for repression of gene expression in mammalian cells.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online: Supplementary Tables 1–3 and Supplementary Figures 1–3.

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