Disruption of mouse poly(A) polymerase mGLD-2 does not alter polyadenylation status in oocytes and somatic cells.
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Abstract

The elongation of poly(A) tails in cytoplasm is essential for oogenesis and early embryogenesis in *Xenopus laevis*. mGLD-2 is a mouse homologue of *Xenopus* cytoplasmic poly(A) polymerase xGLD-2. We found an association of mGLD-2 with cytoplasmic polyadenylation components, CPEB and CPSF described in *Xenopus* oocytes. To clarify the role of mGLD-2 in mouse, we produced an mGLD-2 disrupted mouse line by homologous recombination. In spite of the ubiquitous expression of mGLD-2, the disrupted mice were apparently normal and healthy. Moreover, it was demonstrated that mGLD-2 disruption did not affect the poly(A) tail elongation in oocytes using reporter RNAs. Coincide with these observations, the maturation of the oocytes was normal and the mice were fertile. Thus mGLD-2 is dispensable for full-term development and oogenesis. Our results also indicate that there is another source of cytoplasmic poly(A) polymerase in mouse.

Keywords: mGLD-2, poly(A) polymerase, cytoplasmic polyadenylation, CPEB, CPSF, PABP, knockout mice, oocyte.
Introduction

The addition of mRNA poly(A) tails initially occurs in the eukaryotic nucleus. The processing of the mRNA precursors is one of the post-transcriptional gene regulations, because the length of the poly(A) tail is implicated in various aspects of mRNA metabolism, including the transport of mRNAs into the cytoplasm, mRNA stability, and translational control of mRNA [1]. However, the poly(A) tail is not exclusively added to mRNAs in the nucleus but also found to take place in the cytoplasm. The elongation of the poly(A) tail length in the cell cytoplasm are essential for normal gametogenesis [2-4] and early embryogenesis [5]. For example, meiotic maturation of *Xenopus* and mouse oocytes has been reported to require cytoplasmic polyadenylation of c-mos and cyclin B1 mRNAs for synthesis of the proteins [6, 7].

Several poly(A) polymerases responsible for the cytoplasmic polyadenylation are identified. Yeast Cid13 participates in DNA replication and genome maintenance, specifically targeting suc22 mRNA that encodes a subunit of ribonucleotide reductase [8], while yeast Cid1 and *C. elegans* GLD-2 targets a subset of mRNAs controlling the inhibition of mitosis and entry into meiosis [4, 9]. In mammals, we have reported that a testis specific cytoplasmic poly(A) polymerase, TPAP, elongates a poly(A) tail of mRNAs essential for spermatogenesis [3]. However, ploy(A) polymerases that may
function in somatic tissues are not known in mammals. Recently, we cloned the GLD-2 mouse homolog, mGLD-2, by PCR using cDNA prepared from an unfertilized mouse oocyte library as a template [10]. The mGLD-2 was expressed throughout the body and localized both in the nucleus and cytoplasm in somatic cells [10]. However, in oocytes in metaphase I and II stage after germinal vesicle breakdown, mGLD-2 was found exclusively in the cytoplasm [10]. mGLD-2 also possessed a CPE (cytoplasmic polyadenylation element)-specific polyadenylation activity in the ooplasm and a knockdown experiment caused the impairment of oogenesis [10]. Thus mGLD-2 was suggested to control the translation of specific proteins in oogenesis. In the present experiment, we produced an mGLD-2 gene disrupted mouse line to clarify the role of mGLD-2. Here, we report the effects of mGLD-2 on polyadenylation activity in somatic cells and oocytes together with the effects on live mouse including oogenesis.

Materials and methods

**GST pull-down assay**

Recombinant proteins were synthesized *in vitro* in the presence of $^{35}$S-labeled Met (1000 Ci/mmol, MP biomedicals) by a TNT T7 Quick Coupled Transcription/Translation system (Promega). GST fusion proteins immobilized on
glutathione–Sepharose gel (Amersham Biosciences) were mixed with \(^{35}\)S-labeled protein in 50 mM Tris/HCl, pH 7.5, containing 150 mM NaCl, 5 mM EDTA, 0.5% Nonidet P-40, and protease inhibitors for 1 h. After the gel was washed five times with the same buffer, proteins on the gel were treated with a Laemmli buffer, and analyzed by SDS–PAGE.

**Generation of mutant mice lacking mGLD-2**

Genomic clones encoding mGLD-2 were isolated from a mouse 129/SvJ genomic DNA library in \(\lambda\)FIXII (Stratagene) using \(^{32}\)P-labeled DNA fragments as probes. Of the ten clones isolated, a clone, termed MGLD4, was used for construction of a targeting vector carrying a neomycin-resistance (neo) expression cassette flanked by a 1.5- and 9.7-kbp genomic region of \(mGLD-2\). For negative selection, an \(HSV-TK\) (\(tk\)) cassette was inserted at the 5’-end of the targeting vector (Fig. 1C). The construct was designed to replace a part of second exon and the whole third exon with the neo expression cassette. Following electroporation of the targeting vector into D3 mouse ES cells, homologous recombinants were selected by using G418 and gancyclovir. Ten ES cell clones containing the targeted mutation were selected from 186 clones, and injected into C57BL/6 mouse blastocysts. Chimeric male mice were crossed to
C57BL/6 females (SLC Inc., Japan) to establish the heterozygous mutant lines. All animal experiments were carried out according to the Guide for the Care and Use of Laboratory Animals in the University of Tsukuba.

**Blot hybridization**

Genomic DNA was prepared from the mouse tail, digested by *Bgl*II and *Pst*I, separated by agarose gel electrophoresis, and transferred onto Hybond-N⁺ nylon membranes (Amersham Biosciences). Total cellular RNA was prepared from liver using Isogen (Nippon Gene, Japan) [3]. The RNA samples were glyoxylated, separated by agarose gel electrophoresis, and transferred onto the nylon membranes. The blots were probed by ³²P-labeled DNA fragments, and analyzed by a BAS-1800II Bio-Image Analyzer (Fuji Photo Film, Japan).

**Antibodies and immunoblot analysis**

Affinity-purified anti-mGLD-2 antibody was prepared as described previously [10]. The antibody against β-tubulin was purchased from Sigma. Mouse tissues were homogenized at 4°C in 20 mM HEPES/KOH, pH 7.5, containing 150 mM NaCl, 1 mM EDTA, 1.0% Triton X-100, 0.5% deoxycholate, 0.1% SDS. Oocytes were lysed at
4°C in 10 mM Tris/HCl, pH 8.0, containing 1.5 mM MgCl₂, 10 mM KCl, 1.0% Triton X-100, 0.5% deoxycholate, 0.1% SDS. Proteins in the supernatant solution were subjected to Western blot analysis. The immunoreactive proteins were visualized by an ECL Western blotting detection kit (Amersham Biosciences).

**Immunoprecipitation**

Nuclear extract of mouse liver was prepared as previously described [10]. Protein A-agarose beads (50% slurry, Pierce), bound with the affinity purified anti-mGLD-2 antibody, were mixed with the nuclear extracts (0.5 mg protein), and then incubated at 4°C overnight. After centrifugation, the pellet was washed five times with the binding buffer, and then subjected to the measurement of polyadenylation activity.

**Assay of polyadenylation activity**

Poly(A) polymerase activities were determined by measuring incorporation of AMP from [α-³²P]ATP (400 Ci/mmol, MP biomedicals) into oligo(A)₁₂ RNA primer in the presence of MnCl₂ [10]. The reaction mixtures were sequentially spotted onto Whatman DE-81 paper, washed with 0.1 M phosphate buffer, pH 7.0, and then measured by liquid scintillation counting.
Polymerase chain reaction (PCR)

Total cellular RNAs were extracted from metaphase II-arrested oocytes using Isogen (Nippon Gene) [10]. For RT-PCR, first-strand cDNA was synthesized from total RNAs by a SuperScript III reverse transcriptase (Invitrogen) using oligo dT\textsubscript{20} as a primer. A portion of the synthesized cDNAs was subjected to PCR using specific primer sets. A set of primers, 5’-TCGGCCCTTCGGTGACG-3’ (sense) and 5’-TGTAAGGACACACGCTCTAGAC-3’ (antisense), corresponding to the sequences in the first and third exons of \textit{mGLD-2}, respectively, was used for assessment of the third exon replacement by \textit{neo}. For the PCR-based poly(A) test, total RNA from oocytes were ligated to a 32-mer RNA oligonucleotide, 5’-UACGCAUACGCUUGGGCGUACCUGUA-3’ (60 pmol), and reacted at 42°C for 50 min with SuperScript II reverse transcriptase (Invitrogen), using a PAT1 oligonucleotide, 5’-ACAAGGTACGCCACAGCGTATG-3’, as a primer. A portion of the reaction mixture was subjected to first round PCR using a set of PAT1 and a gene-specific primer. Second round PCR was carried out using a nested set of primers, PAT2, 5’-GGCTCGAGGTACGCCACAGCGTATG-3’, and another gene-specific primer. Details of the PCR conditions, including other primer sequences are described.
previously [10].

**RNA synthesis in vitro**

DNA fragments including the CPE/HEX and mutCPE/HEX of cyclin B1 mRNA were amplified by PCR from a mouse testis cDNA library [11] using primer sets, CYBP7/9 and CYBP8/9, respectively. The following oligonucleotides were used as primers: CYBP7 (5’–AAGTCGACTTTAATTATACATCTGATATCAAG–3’), CYBP8 (5’–AAGTCGACTTTGGATTTATACATCTGATATCAAG–3’), and CYBP9 (5’–TAAAGCTTTCCACCAATAAATTTTATCCAA–3’). The PCR products were introduced between EGFP and 45 adenosine residues of a pEGFPA45 vector. The resulting plasmids were linearized by cutting with BsmBI. RNAs were synthesized with T3 polymerase using an mMESSAGE mMACHINE T3 Kit (Ambion).

**Microinjection**

Oocytes at metaphase II stage were collected from the oviducts of female mice that had been superovulated by pregnant mare’s serum gonadotropin (5 units, ASKA Pharmaceutical, Japan) followed by human chorionic gonadotropin (5 units, ASKA Pharmaceutical) 48 h later. A mixture of equal volumes of RNAs (1 mg/ml) and
TRITC-labeled dextran (50 μM) was injected into the oocytes using a Piezo-driven micromanipulator (Prime Tech Ltd., Japan). Approximately 10 pl of the RNA solution was introduced into the oocytes. The oocytes were incubated in drops of kSOM medium [10] covered with mineral oil (Sigma) at 37°C under 5% CO₂ in air. The oocytes were observed using an Olympus IX-70 inverted microscope equipped with a SPOT RT camera (Diagnostic Instruments). The 16-bit digital images were analyzed by using a Metamorph software (Universal Imaging Corp.).

**Results and discussion**

*Interaction of mGLD-2 with the cytoplasmic polyadenylation complex*

GLD-2 in *C. elegans* encodes the catalytic moiety of a cytoplasmic poly(A) polymerase (PAP) that is associated with a regulation of mitosis/meiosis decision and other germline events [4]. xGLD-2 from *Xenopus* is also known to function in ooplasm which implies that cytoplasmic PAPs are fundamental factors to regulate the translation of mRNA [12]. The most prominent and well investigated site for cytoplasmic polyadenylation is the oocytes. In *Xenopus*, the translational levels of the mRNAs are controlled by xGLD-2 together with polyadenylation regulatory factors such as CPEB (CPE-binding protein) and CPSF (cleavage and specificity factor)
during oogenesis and early embryogenesis [12]. We have already shown the CPE (cytoplasmic polyadenylation element)-specific polyadenylation activity of mGLD-2 [10]. To further investigate the roles of mGLD-2 in vitro, we produced recombinant mGLD-2 and examined its association with various GST-tagged proteins included in cytoplasmic polyadenylation machinery. As shown in Fig. 1A, mouse CPEB1 [13] and cytoplasmic poly(A) binding protein 1, PABPC1 [14], interacted with mGLD-2. CPEB2 [15], which is expressed in mouse testis was also found to interact with mGLD-2. To analyze the binding ability of mouse CPSF160 [16], a subunit of CPSF, to mGLD-2, we produced GST-tagged mutant mGLD-2 (D215A), which has almost no polyadenylation activity [10], and mixed it with recombinant CPSF160 (Fig. 1B). This was necessary due to the low yield of GST-tagged CPSF160 and mGLD-2. Moreover, the association of recombinant CPEB1, CPEB2, and PABPC1 with GST-tagged D215A was also demonstrated (Fig. 1B). These data indicate the involvement of mGLD-2 in the regulation of mRNA translation by associating with the cytoplasmic polyadenylation complex which is formed on the poly(A) tail.

Production of mGLD-2-deficient mice

Mutant mice lacking mGLD-2 were produced by homologous recombination in ES
cells, using a targeting vector containing neo and tk expression cassettes (Fig. 1C). A part of the second exon and the whole third exon were replaced by the neo cassette. The genotypes of wild-type (mGLD-2^+/+), heterozygous (mGLD-2^+/−), and homozygous (mGLD-2^−/−) mice for the null mutation of mGLD-2 were identified by Southern blot analysis of genomic DNA (Fig. 1D). Mating of mGLD-2^+/− male and female mice yielded the expected Mendelian frequency of mGLD-2^−/− mice (mGLD-2^+/+: mGLD-2^+/−: mGLD-2^−/− = 10 (17%) : 34 (58%) : 15 (25%) for 59 offspring from 6 litters). Both mGLD-2^−/− males and females were apparently normal in behavior, body size, and health condition. Northern blot analysis indicated the absence of mGLD-2 mRNA in the liver of mGLD-2^−/− mice (Fig. 1E). Moreover, protein extracts of the mGLD-2^−/− liver completely lacked mGLD-2 (Fig. 1F). These data demonstrate the successful gene disruption of mGLD-2^−/− in our mutant mouse line.

Analysis of mGLD-2-deficient somatic cells

mGLD-2 localizes both in the nucleus and in the cytoplasm [10]. We prepared mGLD-2 from the nuclear extract of mouse liver cells by immunoprecipitation and assayed the polyadenylation activity. As a result, it was demonstrated the polyadenylation activity in mGLD-2^+/− mice are about half that of the mGLD-2^+/+ mice
and disappeared in \textit{mGLD-2}^{+/--} mice (Fig. 2A). However, when we observed the total polyadenylation activity using nuclear or cytoplasmic extract, we could not find any significant differences between \(mGLD-2^{+/--}\) and \(mGLD-2^{+/--}\) mouse extracts (Fig. 2B). The results indicate that the disruption of mGLD-2 did not affect the overall poly(A) length in the liver.

\textit{Analysis of mGLD-2-deficient oocytes}

We previously reported that over expression of mGLD-2 in mouse oocytes could selectively elongate the poly(A) tail length of cyclin B1 and Mos mRNAs \cite{10}. This may indicate that mGLD-2 is involved in oogenesis by controlling the translation of key factors. Thus we next examined the effect of mGLD-2 disruption on oocytes. Naturally, the expression of mGLD-2 was demonstrated to disappear totally in \(mGLD-2^{+/--}\) mouse oocytes in mRNA and also in protein levels (Figs. 3A and 3B). However, as observed in the liver, no significant decrease in overall polyadenylation activity was evident in \(mGLD-2^{+/--}\) mouse oocytes (Fig. 3C). For more detailed analysis, we examined the change in poly(A) tail length of cyclin B1 and Mos mRNA by amplifying the poly(A) tails by PCR using two primer sets described in the materials and methods. However, we could not find any significant difference in the poly(A)
tails in germinal vesicle, metaphase I and metaphase II-stage oocytes from mGLD-2/- mice (Fig. 3D).

Noninvasive measurement of the cytoplasmic polyadenylation activity

In order to quantify the CPE dependent polyadenylation activity in oocytes, we produced synthetic EGFP RNA with 3’-UTR of cyclin B1 which contained CPE and AAUAAA followed by a 45 bp length poly(A) tail (Fig. 4A). Since, the elongation of cyclin B1 poly(A) tail is reported to take place in metaphase I and II-stage oocytes [7], we injected the reporter RNA into oocytes at metaphase II stage. Six hours after the introduction of the RNA, a tremendous increase of fluorescence from EGFP was observed (Fig. 4B). On the other hand, the oocytes did not become fluorescent when the reporter RNA which contained mutated CPE was injected (Figs. 4A and 4B). Using this system, we measured the cytoplasmic polyadenylation activity in mGLD-2 disrupted oocytes. Different from our expectation, as shown in Figs. 4B and 4D, the mGLD-2 null oocytes became fluorescent by the injection of the reporter RNA. Concomitant with this observation, the elongation of the poly(A) tail was observed in the reporter RNA including the CPE sequence (Fig. 4C). Reflecting the normal cytoplasmic polyadenylation in mGLD-2 knockout mouse oocytes, the transition rate of
mGLD-2\(^{−/−}\) mouse oocytes from germinal vesicle to metaphase II stage \textit{in vitro} was comparable to that of mGLD-2\(^{+/−}\) oocytes (\(+/−, 80 \pm 13\%\), \(n = 3\); \(-/−, 77 \pm 11\%\), \(n = 3\)). Moreover, the spindle formation in the mGLD-2 disrupted oocytes was normal when oocytes were observed after staining with anti-β-tubulin antibody and Hoechst 33342 (data not shown). The lack of apparent phenotype was reassured by the normal number of pups obtained from mGLD-2\(^{−/−}\) female mice (7.9 ± 3.0, \(n = 13\)) compared to the mGLD-2\(^{+/−}\) female mice (8.9 ± 2.7, \(n = 8\)) when mated with wild-type mice.

In \textit{Xenopus}, there is a notion that xGLD-2 is the responsible enzyme for CPE-dependent cytoplasmic polyadenylation, but this was not applicable in mouse since mGLD-2 was indicated not to be an exclusive enzyme for the polyadenylation. In mammalian cells, six PAPs have been identified: canonical PAP (PAPI and PAPII) \cite{17}, TPAP \cite{3}, neo-PAP \cite{18}, PAP\(γ\) \cite{19}, GLD-2 \cite{4}, and nuclear-encoded mitochondrial PAP, mitoPAP \cite{20}. These mammalian PAPs, except TPAP, GLD-2, and mitoPAP, are exclusively localized in the nucleus. Nuclear PAP localizes in the cytoplasm during metaphase where the nuclear envelope disappears. However, canonical PAP has been reported to be inactivated via phosphorylation by p34cdc2/cyclinB complex during metaphase \cite{21}. Mitochondria is another source of PAP, but mitoPAP localizes only
inside the mitochondria [20]. Thus PAP that functions in cytoplasm is not known in mouse except mGLD-2. The result of our RNAi experiment suggested an important role of mGLD-2 in mouse oocyte maturation [10]. Contrary to our expectation, we could not find any phenotype in mGLD-2 deficient mice. This might be a result of some compensation effect (up regulation) of a yet unknown mouse cytoplasmic poly(A) polymerase. However, the compensation was not effectively functional in the RNAi experiment [10], which may relate to the nature of the sudden impairment of mGLD-2 activity in RNAi.

Combining the results from other cases, where distinctive phenotypes are observed such as disruption of GLD-2 in C. elegans, Cid1 and Cid13 in yeast, [4, 8, 9, 22] and testis specific PAP (TPAP) in mouse [3], the lack of phenotype in the mGLD-2 disrupted mouse line may indicate the existence of a yet unknown cytoplasmic PAP. We presume that it would be worth to quest a new cytoplasmic PAP to elucidate the regulation of mRNA translation leading to dramatize various characteristics of living cells.

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References


Cid13 is a cytoplasmic poly(A) polymerase that regulates ribonucleotide reductase mRNA, Cell 109 (2002) 563-573.


**Figure Legend**

**Figure 1**

Direct interaction of mGLD-2 with components in polyadenylation complex and generation of mGLD-2-deficient mice. (A) GST pull-down assays were carried out using $^{35}$S-labeled mGLD-2 and each polyadenylation components fused with GST. (B) GST pull-down assays were carried out using $^{35}$S-labeled polyadenylation components and mutant mGLD-2, D215A, fused with GST. (C) Schematic representation of the gene targeting strategy. For detail, refer to the materials and methods. Restriction enzyme sites indicated are as follows: S, SacI; G, BglII; P, PstI; X, XhoI, Sa; SalI. (D) Southern blot analysis of genomic DNA from wild-type ($mGLD-2^{+/+}$, +/+), heterozygous ($mGLD-2^{+/c}$, +/-) and homozygous ($mGLD-2^{-/-}$, –/–) mice. Genomic DNA was digested by BglII and PstI, separated by agarose gel electrophoresis, and subjected to Southern blot analysis using a $^{32}$P-labeled DNA fragment (S-probe in panel C) as a probe. The wild-type and targeted alleles yielded 4.7- and 5.5-kbp DNA bands, respectively. (E) Northern blot analysis of total RNAs from liver tissues. Two forms (3.5 and 2.3 kbp) of mGLD-2 mRNA, distinguished by the length of the 3’-untranslated region encoded by the same exon of the mGLD-2 gene, were absent in $mGLD-2^{-/-}$ mouse liver. G3PDH, glyceraldehyde-3-phosphate dehydrogenase.
dehydrogenase. (F) Western blot analysis of protein extracts from liver tissues.

Proteins (10 μg) were loaded in each lane and probed with affinity-purified anti-mGLD-2 or anti-β-tubulin antibody. The mGLD-2+/ mouse liver lacked mGLD-2 with the size of 54 kDa. An asterisk indicates the location of a 51-kDa protein nonspecifically immunoreactive with the anti-mGLD-2 antibody.

Figure 2

Polyadenylation activity in mGLD-2+/ mouse liver. (A) Enzyme activity of mGLD-2 in the liver nuclear extract. Nuclear protein (0.5 mg) of mGLD-2+/+ (open circle), mGLD-2+/- (shaded circle), and mGLD-2-/- (closed circle) mice were immunoprecipitated using affinity-purified anti-mGLD-2. Polyadenylation activities of the precipitation were monitored by measuring the incorporation of AMP from [α-32P]ATP into oligo(A)12 primer. (B) Total polyadenylation activity of nuclear and cytoplasmic protein (4 μg) extracted from mGLD-2+/+ (shaded column) and mGLD-2-/- (closed column) mouse liver. Relative activities measured at 6 h are indicated as means ± S.D., where n = 3.

Figure 3
Polyadenylation activity in \( mGLD-2^{-/-} \) mouse oocytes. (A) RT-PCR analysis of total RNAs from metaphase II-stage oocytes. No DNA band was detected in \( mGLD-2^{-/-} \) mice. G3PDH, glyceraldehyde-3-phosphate dehydrogenase. (B) Western blot analysis of protein extracts from metaphase II-stage oocytes. Proteins of oocytes (50 cells per lane) were loaded in each lane and probed with affinity-purified anti-mGLD-2 or anti-\( \beta \)-tubulin antibody. (C) Total polyadenylation activity of protein extracts (50 cells) from metaphase II-stage oocytes of \( mGLD-2^{+/+} \) (shaded column) and \( mGLD-2^{-/-} \) (closed column) mice. Relative activities measured at 6 h are indicated as means ± S.D., where \( n = 4 \). (D) The sizes of poly(A) tails of cyclin B1, Mos, and \( \beta \)-actin, as a negative control. Total RNAs from the GV-, MI- and MII-stage oocytes (50 cells) of \( mGLD-2^{+/+} \) and \( mGLD-2^{-/-} \) mouse were subjected to PCR-based poly(A) test.

**Figure 4**

Cytoplasmic polyadenylation activity of \( mGLD-2^{-/-} \) mouse oocytes. (A) Schematic representation of the synthetic reporter RNAs. EGFP open reading frame was fused to 48 bp of the cyclin B1 3’-UTR. The polyadenylation hexanucleotide (AAUAAA), CPE and poly(A) tail are indicated. (B) Fluorescence images of \( mGLD-2^{+/+} \) and
mGLD-2+/− mouse oocytes 6 h after the injection of the reporter RNAs. TRITC-labeled dextran was coinjected with the RNA to normalize the amount of RNA injected (D). Scale bar = 100 μm. (C) PCR-based poly(A) test of reporter RNAs injected into mGLD-2+/− and mGLD-2−/− mouse oocytes. Oocytes (8 cells) were subjected to the analysis 6 h after the RNA injection. (D) Quantification of cytoplasmic polyadenylation activities of mGLD-2+/− and mGLD-2−/− mouse oocytes. EGFP fluorescence of the oocytes was measured at 2-h intervals, and was divided by that of the TRITC-labeled dextran coinjected with the reporter RNA. The highest intensity of an oocyte after 6-h incubation was set at 100%. Number of mGLD-2+/− oocytes examined: 15 (n = 2) and 8 (n = 1) injected with CPE(−) and CPE(+) RNA, respectively. Number of mGLD-2−/− oocytes examined: 17 (n = 2) and 10 (n = 1) injected with CPE(−) and CPE(+) RNA, respectively. The data indicate the means ± S.D.
Figure 1. Nakanishi et al.

A

mGLD-2

B

CPEB1
CPEB2
CPSF160
PABPC1

C

Wild-type allele

Targeting construct

Targeted allele

D

E

F

β-Tubulin
Figure 2. Nakanishi et al.
Figure 3. Nakanishi et al.

A

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B

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C

Oocytes

Relative activity (%)

D

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bp

600
400
200
Figure 4. Nakanishi et al.

A

CPE (+) EGFP UUUUAAU AAUAAA A₄₅
CPE (-) EGFP UUUGGAU AAUAAA A₄₅

Cyclin B1 3'-UTR

B

EGFP

Dextran

C

CPE ⁺⁻ ⁺⁻ bp

400
300
200
100

D

Relative intensity

0.0 0.2 0.4 0.6 0.8 1.0

Time (h)

0 2 4 6

CPE (+)

CPE (-)