DEADSouth protein localizes to germ plasm and is required for the development of primordial germ cells in *Xenopus laevis*

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**Summary**

DEADSouth mRNA is a component of germ plasm in *Xenopus laevis* and encodes a DDX25 DEAD-box RNA helicase. To determine the intracellular localization of DEADSouth protein, we injected mRNA encoding DEADSouth tagged with mCherry fluorescent protein into fertilized eggs from transgenic *Xenopus* expressing EGFP fused with a mitochondrial targeting signal. The DEADSouth-mCherry fusion protein was localized to the germ plasm, a mitochondria-rich region in primordial germ cells (PGCs). **DEADSouth** overexpression resulted in a reduction of PGC numbers after stage 20. Conversely, **DEADSouth** knockdown using an antisense locked nucleic acid gapmer inhibited movement of the germ plasm from the cortex to the perinuclear region, resulting in inhibition of PGC division at stage 12 and a decrease in PGC numbers at later stages. The knockdown phenotype was rescued by intact **DEADSouth** mRNA, but not mutant mRNA encoding inactive DEADSouth helicase. Surprisingly, it was also rescued by mouse *vasa* homolog and *Xenopus* *vasa*-like gene 1 mRNAs that encode DDX4 RNA helicases. The rescue was dependent on the 3′ untranslated region (3′UTR) of **DEADSouth** mRNA, which was used for PGC-specific expression. The 3′UTR contributed to localization of the injected mRNA to the germ plasm, resulting in effective localization of DEADSouth protein. These results demonstrate that localization of DEADSouth helicase to the germ plasm is required for proper PGC development in *Xenopus laevis*.

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Key words: RNA helicase, Germline, LNA gapmer

**Introduction**

In most animals, germline cells are generated apart from the somatic lineage in early development. In contrast to mice, in which the germline is generated by inductive signals from the surrounding tissues, the *Xenopus* germline is established by inheriting specialized cytoplasm localized in the vegetal cortex of the egg. Such cytoplasm in *Xenopus* is also observed in various animals including zebrafish, nematode and fly, which is termed ‘germ plasm’ in general, the P-body in *Caenorhabditis elegans* and ‘polar plasm’ in *Drosophila melanogaster* (Ikenishi, 1998). It has been demonstrated that germ plasm contains determinants required and sufficient for germline differentiation in *Drosophila* (Illmensee and Mahowald, 1974; Okada et al., 1974) and *Xenopus* (Buehr and Blackler, 1970; Tada et al., 2012).

In early *Xenopus* development, germ plasm at the vegetal cortex of fertilized eggs is divided into about four blastomeres through the first two cleavages, and then distributes to only one side of two daughter cells because it is present in a particular region of the cortex of blastomeres (Whittington and Dixon, 1975). Thus, the number of primordial germ cells (PGCs) harboring germ plasm remains at about four, although the size of PGCs becomes gradually smaller through repeated cell division until the late blastula at stage 9. Early in gastrulation, germ plasm moves from the cortex to the perinuclear region in PGCs and divides equally into two daughter PGCs in later cell divisions. PGCs incorporating germ plasm divide about three times and migrate to the genital ridge (Dziadek and Dixon, 1977). The germ plasm is composed of electron dense granules, many mitochondria, and specific mRNAs and proteins. Although many molecules in germ plasm have been identified and investigated, the mechanisms of germline development remain unknown (Cuykendall and Houston, 2010; King et al., 2005).

**VASA/DDX4** of the DEAD-box RNA helicase family is a component of germ plasm and widely used as a germline marker in various animals because of its germline-specific expression (Gustafson and Wessel, 2010). It has been reported that *vasa* is required for germ cell development. In fly and nematode, loss-of-function of *vasa* results in a defect of oogenesis (Kuznicki et al., 2000; Styhler et al., 1998). **MVH** (mouse *vasa* homolog) is present in the chromatoid body that is observable during spermatogenesis and resembles germ plasm (Toyooka et al., 2010). MVH-null mutants show defects in spermatogenesis (Tanaka et al., 2000). Recently, it was also revealed that *vasa* is involved in cell cycle progression in fly (Pek and Kai, 2011a) and sea urchin (Yajima and Wessel, 2011), and piRNA production in mice (Kuramochi-Miyagawa et al., 2010).
contrast, \textit{XVLG1} (\textit{Xenopus} vasa-like gene 1) is expressed in somatic and germ cells in \textit{Xenopus}, although \textit{XVLG1} is certainly a homolog of \textit{vasa/Ddx4} (Ikenishi and Tanaka, 2000; Komiya et al., 1994). Functional inhibition of \textit{XVLG1} by an antibody results in aberrant morphogenesis of somatic cells and the loss of germ cells, suggesting that \textit{XVLG1} is involved in the differentiation of both somatic and germline cells (Ikenishi and Tanaka, 1997; Ikenishi and Tanaka, 2000).

\textbf{DEADSouth is a DEAD-box RNA helicase belonging to DDX25, but not the VASA/DDX4 family, and its transcript is an RNA component of germ plasm in \textit{Xenopus} (MacArthur et al., 2000). \textit{DEADSouth} transcripts were initially detected prior to mitochondrial cloud formation, accumulate in the mitochondrial cloud and then co-localize to the germ plasm during early development. It is also expressed in spermatogonia, spermatocytes and spermatids in testes. In mammals, DDX25 has been identified as a gonadotropin-regulated testicular RNA helicase (GRTH) (Tang et al., 1999). Knockout of the gene results in remarkably diminished sizes of chromatioid bodies in spermatids and incomplete spermatogenesis in mice (Tasi-Morris et al., 2004). It has been suggested that GRTH/DDX25 may be required to maintain chromatioid bodies in spermatogonia (Sato et al., 2010). In this study, we focus on the function of \textit{DEADSouth} in \textit{Xenopus} germline development in relation to VASA/DDX4. By expression of mRNA encoding \textit{DEADSouth} fused with mCherry fluorescent protein, we show that the \textit{DEADSouth} protein is localized to the germ plasm. We show that PGCs decrease in number both in \textit{DEADSouth-overexpressing} and -depleted embryos. The depleted embryos are deficient for translocation of germ plasm from the cortex to the perinuclear region in PGCs before midblastula transition (MBT), causing inhibited PGC division. Surprisingly, such a knockdown phenotype is rescued by expression of MVH and \textit{XVLG1}, but not inactive \textit{DEADSouth} helicases. The rescue is dependent on the 3' untranslated region (3' UTR) of \textit{DEADSouth} mRNA, which is required for localization of the injected mRNA to the germ plasm. Together, we demonstrate that localization of \textit{DEADSouth} protein is required for proper development of \textit{Xenopus} PGCs.

\section*{Materials and Methods}

\textbf{Xenopus embryos}

Adult male and female wild-type \textit{Xenopus laevis} were purchased commercially and maintained at 22°C in circulatory tanks. Female transgenic \textit{Xenopus laevis} expressing EGFP fused with a mitochondrial targeting signal (mito-EGFP \textit{Xenopus}) was also used in this study (Taguchi et al., 2012). Because embryos from the transgenic frog had maternally supplied mitochondria with EGFP, the mitochondria-rich germ plasm was clearly visible until stage 20 under a fluorescence microscope. Embryos were obtained as described previously (Kataoka et al., 2006), allowed to develop at 18°C, and staged according to Nieuwoop and Faber (Nieuwoop and Faber, 1994).

\textbf{Preparation of constructs}

\textit{Mvh} cDNA was a gift from Drs Kuniya Abe (RIKEN BioResource Center) and Toshiaki Noce (Keio University). \textit{DEADSouth} and \textit{XVLG1} CDAs were amplified by PCR using \textit{Xenopus laevis} ovarian CDA as a template, and cloned into pCR2.1 or pCRIII vectors. Then, the ORFs and 3'UTR's were amplified by PCR using PrimeStar polymerase (TaKaRa), and inserted into appropriate regions of pCS2-based vectors such as pCS2-Venus-DEADSouth 3' UTR and pCS2-mCherry-DEADSouth 3' UTR (Kataoka et al., 2006) by In-Fusion technology (Clontech). \textit{pcS2-DEADSouth-AAA-DEADSouth} 3' UTR and pCS2-DEADSouth-AAA-DEADSouth 3' UTR were constructed by inverse PCR with mutated primers and In-Fusion technology. The coding regions of all constructs were confirmed by sequencing. \textit{XVLG1} ORF from pCS2-XVLG1-DS 3' UTR and pCS2-XVLG1-XVLG1 3' UTR were generated by removing the EcoRV fragment of the \textit{XVLG1} ORF from pCS2-XVLG1-DS 3' UTR and pCS2-XVLG1-XVLG1 3' UTR plasmids, respectively.

\section*{Locked nucleic acid (LNA) gapmers}

According to previous reports (Kurreck et al., 2002; Brasch et al., 2002), antisense and mismatched LNA gapmers (19 bases each) were designed. Sequences of antisense and mismatched LNA gapmers were TTAGCGGCCATCGTTTCCCT and TATCCGGCCATCGTTGGAG (modification with LNA underlined), respectively (Nippon EGT). LNA gapmers were dissolved in water at 20 µM. Ninety-two femtomoles of gapmer were injected into each embryo.

\section*{Preparation and microinjection of mRNA}

Template plasmids were linearized by digestion with \textit{Xhol}, and used as templates for \textit{in vitro} mRNA synthesis with a mMESSAGE mMachine SP6 kit (Ambion). mRNAs of \textit{XVLG1}-DS 3' UTR and \textit{XVLG1A}-XVLG1 3' UTR were labeled using a Label IT Cy3 labeling kit (Mirus). mRNAs and LNA gapmers were microinjected into the cortical region at the vegetal pole of a fertilized egg with a Nonojekt II microjector (Drummond Scientific Company). The amount of each injected mRNA per egg was determined empirically as follows. Four-hundred and sixty picograms of \textit{Venus-DEADSouth} 3' UTR (v-DS), \textit{DS-mCherry-DS} 3' UTR, \textit{DS-mCherry-XVLG1} 3' UTR or \textit{mCherry-DS} 3' UTR; 46 pg (for overexpression) or 92 pg (for rescue) of DS-full, 46 pg of DS-stop; 92 pg of \textit{DS-AAA}, \textit{Mvh-DS} 3' UTR or \textit{DS-XVLG1} 3' UTR; 276 pg of \textit{VXLG1-DS} 3' UTR or \textit{XVLG1-XVLG1} 3' UTR; 3.7 ng of \textit{Cy3-XVLG1A-DS} 3' UTR or \textit{Cy3-XVLG1A-XVLG1} 3' UTR were injected. Injection at these doses did not affect normal development.

\section*{PGC isolation and observation}

Mito-EGFP \textit{Xenopus} or v-DS-injected embryos at the indicated stages were dissected in 70% Dulbecco's phosphate-buffered saline without Ca²⁺ and Mg²⁺ (PBS⁻) and incubated for 1 hour. Then, PGCs were collected manually with a micropipette and observed under an MZ16f fluorescence stereomicroscope (Leica) equipped with a DS-5Mc digital camera (Nikon).

The diameter of PGCs was measured in images with a scale. To visualize mCherry protein, isolated PGCs were fixed in 2% paraformaldehyde/0.1 M MOPS (pH 7.5)/0.5 M NaCl at 4°C for at least 2 days, stained with 2 µg/mL Hoechst 33342 (Sigma) in PBS⁻, washed with PBS⁻ and mounted in 50% glycerol/PBS⁻. Cells were observed under a BX60 fluorescence microscope (Olympus) equipped with a Nuance Fx digital camera (Caliper PerkinElmer).

\section*{Quantitative real-time RT-PCR}

Total RNA was extracted from 10 embryos using TRIzol (Invitrogen). One microgram of total RNA was subjected to cDNA synthesis using Ready-To-Go You-Prime First-Strand Beads with random hexamers (GE healthcare Life Science), according to the manufacturer’s protocol. mRNA quantification was performed by a PRISM7000 Sequence Detection System (Applied Biosystems) using SYBR Premix Ex Taq (TaKaRa). Primer sequences and annealing temperatures were as follows. \textit{DEADSouth} forward ATGGGCTTCAACAGAATGCT and reverse ATGGGCTTCAACAGACCTTC and reverse TCCACAGGCTCACTCAGATAC, 60°C; \textit{Xvlg1} forward GAGAGGAAATTGTTGCTGGATATGC and reverse GAGAGGAAATTGTTGCTGGATATGC, 66°C. Standard curves were generated for each gene by serial dilution of cDNA from unjected stage 8 embryos. \textit{Ef1a} gene expression was used as an internal control. Each experiment was repeated three times and RNA quantification was performed twice for each experiment.

\section*{Immunostaining}

Embryos were fixed in 4% paraformaldehyde/PBS⁻ at 4°C overnight, washed in PBS⁻ (0.1% Triton X-100 in PBS⁻), and then incubated in blocking buffer (10% goat serum in PBS⁻) at room temperature for 2 hours. Samples were treated with blocking buffer containing mouse anti-Xdaxl (1:1000 dilution) (Mita and Yamashita, 2000) and rabbit anti-capase-3 (1:1000 dilution) (Abcam) antibodies at 4°C for 3 days, washed in PBS⁻, incubated in blocking buffer containing Alexa Fluor 488-conjugated anti-mouse IgG (H+L) (1:1000 dilution) (Molecular Probes) and Cy3-conjugated anti-rabbit IgG (1:1000 dilution) (Rockland) at 4°C overnight and then washed in PBS⁻. Specimens were dissociated manually in PBS⁻ and then observed under an MZ16f fluorescence stereomicroscope (Leica).

\section*{Statistical analyses}

The Student’s t-test was carried out for all statistical analyses. One-tailed P-values were used for significance.

\section*{Results}

\textbf{Localization of \textit{DEADSouth} protein in \textit{Xenopus} PGCs}

Although \textit{DEADSouth} mRNA is localized to the germ plasm during oogenesis and early development (MacArthur et al., 2000), localization of the protein is unknown. Because an...
antibody against DEADSouth protein was not available, we expressed DEADSouth protein tagged with a fluorescent protein. To reveal the relative location of DEADSouth protein in germ plasm, we also used mito-EGFP *Xenopus* (Taguchi et al., 2012). Germplasm is a PGC-specific organelle enriched with mitochondria that are a useful marker of germ plasm (Venkatarama et al., 2010; Elinson et al., 2011). We prepared mRNA encoding DEADSouth protein and mCherry fluorescent protein, followed by the DEADSouth 3’ UTR (DS-mCherry-DS 3’ UTR), and injected it into fertilized eggs obtained by crossing a mito-EGFP female with a wild-type male. At stage 12, the germ plasm was observed in the perinuclear region of PGCs (Fig. 1A–E). The mCherry signal clearly overlapped with the germ plasm and appeared to be observable in the nuclei of PGCs. This observation suggested that DEADSouth protein played a role in the nucleus and the germ plasm, but required further analysis. As a control, mCherry signal from mCherry-DS 3’ UTR was observed throughout PGCs, but not localized to the germ plasm (Fig. 1F–J). This result indicates that DEADSouth protein is localized to the germ plasm.

**Overexpression effects of the DEADSouth gene**

To elucidate the function of DEADSouth in PGC development, we synthesized full-length DEADSouth mRNA (DS-full) *in vitro* and injected it into the vegetal pole of fertilized eggs, together with Venus-DEADSouth 3’ UTR mRNA (v-DS), to monitor PGCs in a living embryo (Kataoka et al., 2006). At stage 32, we externally observed that PGCs were significantly reduced in number (Fig. 2A–D). The number of PGCs per embryo injected with water (control) or DS-full were 17.6±2.3 and 10.7±1.2 (average ± s.d., P<0.001), respectively. No effect was observed in embryos injected with mRNA encoding nonsense DEADSouth (DS-stop), in which a stop codon was inserted downstream of the initial methionine of the DEADSouth ORF. This result indicated that the loss of PGCs was due to overexpression of DEADSouth protein.

To determine when the effect of DEADSouth gene overexpression appeared, we used mito-EGFP transgenic *Xenopus*. Embryos are more suitable for monitoring PGCs at an early stage, compared with the use of v-DS as a PGC tracer, because of no delayed EGFP expression. We dissociated embryos injected with DS-full and checked the PGCs in detail. At stages 12 and 20, no difference in PGC numbers between DS-full and the control (uninjected) was detected (Fig. 2E). DEADSouth-overexpressing embryos also showed no difference in PGC size and intracellular localization of germ plasm, compared with that of the control at stages 12, 20 and 32 (supplementary material Figs S1, S2). To determine whether the loss of PGCs resulted from apoptosis, we performed immunostaining for Xdazl and caspase-3 to detect apoptotic PGCs in DEADSouth-overexpressing embryos at stage 28 (Fig. 2F). As expected, 17% of PGCs (10 caspase-3-positives out of 60 Xdazl-positives) were apoptotic, compared with 4.8% of PGCs in control embryos (3 out of 63). This result indicated that the loss of PGCs in DEADSouth-overexpressing embryos, at least partially, resulted from apoptosis.

**Knockdown of DEADSouth with an antisense LNA gapmer**

To reveal the function of DEADSouth, we performed knockdown experiments using an antisense LNA gapmer. Because an antibody against DEADSouth is unavailable, it is difficult to evaluate the extent of knockdown after translational inhibition using morpholino oligos. Therefore, we chose an LNA gapmer as a knockdown reagent because it caused degradation of the target mRNA that was quantified easily by PCR. As shown in Fig. 3A, we designed an antisense LNA gapmer targeting around the start codon of DEADSouth mRNA and a mismatch gapmer as a control that was not expected to act on the target.

First, we evaluated the extent of DEADSouth mRNA degradation in embryos injected with the LNA gapmer by quantitative RT-PCR. The mRNA level of Xpat, which is expressed in a PGC-specific manner (Hudson and Woodland, 1998), was also examined as a control. Because PGC number and the amount of germ plasm depended on the embryos and stages, the amount of DEADSouth mRNA was normalized to that of EF1z mRNA and compared between embryos injected with antisense or mismatch LNA gapmers (Fig. 3B). At stages 8 and 32, DEADSouth mRNA levels in antisense LNA-injected embryos were reduced significantly compared with those in mismatch LNA gapmer-injected embryos (P<10⁻⁹ at stage 8). In contrast, Xpat mRNA levels in both embryos were similar at stages 8 and 32. These mRNA levels in mismatch LNA gapmer-injected embryos were similar to those in uninjected embryos (data not shown). These data indicated that injection of the LNA gapmer resulted in degradation of DEADSouth mRNA in a sequence-specific manner. Unfortunately, stage-dependent changes of the mRNA level could not be examined because the total amount of mRNA in the embryo changed during development.

Second, we examined the effects of the LNA gapmer on PGC development by coinjection with v-DS as a PGC tracer. External observation at stage 32 showed that PGCs in antisense gapmer-injected embryos decreased remarkably in number compared with those in mismatch gapmer-injected embryos (antisense, 6.3±1.7; mismatch, 17.3±2.1). In addition, the reduction of PGC number in antisense gapmer-injected embryos was rescued by coinjection with DS-full mRNA that was ineffective for antisense gapmer targeting because of codon substitutions in the target region (Fig. 3). Although this rescue was significant (P<10⁻⁵ between antisense and antisense+DS-full), it appeared to be
In contrast, this reduction was not rescued by co-injection with DS-AAA mRNA encoding a putatively inactive helicase of DEADSouth, in which the amino acid sequence SAT was substituted with AAA in motif III (Pause and Sonenberg, 1992) (Fig. 4). These results clearly show that DEADSouth RNA helicase is required for proper PGC development.

We also dissociated DEADSouth-knockdown embryos at stages 12, 20 and 32, and examined PGC number per embryo and size. For stage 12 and 20 embryos, mito-EGFP transgenic Xenopus were used because of easier PGC tracing. PGCs from knockdown embryos at stages 12, 20 and 32 showed less increasing numbers (6.6±1.8/embryo at stage 12 to 9.3±1.7/embryo at stage 32) in contrast to control embryos (mismatch LNA injected) (10.3±2.5/embryo at stage 12 to 21.9±3.4/embryo at stage 32). At stage 12, knockdown embryos already showed that the number of PGCs was significantly decreased compared with that of the control (P<0.05 between mismatch and antisense+DS-full). In contrast, this reduction was not rescued by co-injection with DS-AAA mRNA encoding a putatively inactive helicase of DEADSouth, in which the amino acid sequence SAT was substituted with AAA in motif III (Pause and Sonenberg, 1992) (Fig. 4). These results clearly show that DEADSouth RNA helicase is required for proper PGC development.

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The decrease in number and increase in size of PGCs in DEADSouth-knockdown embryos appeared to be caused by inhibition of PGC division. In addition, we found a defect in the subcellular localization of germ plasm in DEADSouth-knockdown PGCs (Fig. 5B). In normal development, the germ plasm is present in a particular region of the cortex in PGCs...
before MBT. Then, it moves to a perinuclear region and is divided equally into two daughter PGCs after MBT. The germ plasm in PGCs from embryos injected with the mismatch LNA gapmer, antisense LNA gapmer, antisense LNA gapmer plus DS-full and antisense LNA gapmer plus mutant DEADSouth-DS 3’ UTR (DS-AAA), in addition to v-DS as a PGC-tracer, respectively. Scale bar: 1 mm. (A–D’) High magnification of the areas indicated in (A–D), respectively. (E) The number of PGCs per embryo at stage 32 injected with the indicated LNA gapmer(s) and mRNAs. PGC numbers were determined by externally counting from both sides. (F) The number of PGCs per embryo at the indicated stages with mismatch or antisense LNA gapmers. The injected mito-EGFP embryos were dissociated, and EGFP-positive and large cells were counted as PGCs. N and n indicate the number of experiments and total embryos examined, respectively. P-values were calculated by the one-tailed t-test. Error bars indicate s.d.

**Fig. 4.** The number of PGCs is decreased by knockdown of the **DEADSouth** gene. (A–D) Representative examples of stage 32 embryos injected with the mismatch LNA gapmer, antisense LNA gapmer, antisense LNA gapmer plus DS-full and antisense LNA gapmer plus mutant DEADSouth-DS 3’ UTR (DS-AAA), in addition to v-DS as a PGC-tracer, respectively. Scale bar: 1 mm. (A’–D’) High magnification of the areas indicated in (A–D), respectively. (E) The number of PGCs per embryo at stage 32 injected with the indicated LNA gapmer(s) and mRNAs. PGC numbers were determined by externally counting from both sides. (F) The number of PGCs per embryo at the indicated stages with mismatch or antisense LNA gapmers. The injected mito-EGFP embryos were dissociated, and EGFP-positive and large cells were counted as PGCs. N and n indicate the number of experiments and total embryos examined, respectively. P-values were calculated by the one-tailed t-test. Error bars indicate s.d.

**Fig. 5.** Knockdown of the **DEADSouth** gene affects PGC division and translocation to germ plasm. (A) Size distribution of PGCs at stages 12, 20 and 32 from embryos injected with mismatch or antisense LNA gapmers. All mito-EGFP- (at stages 12 and 20) and v-DS-labeled (at stage 32) PGCs from the injected embryos were isolated to measure their diameter. Total PGC numbers after each injection are shown as 100%. ‘n’ indicates total PGC numbers. Arrows indicate average diameters from the indicated experiments. (B) Localization of the germ plasm in PGCs isolated from mito-EGFP embryos at stages 7 and 12, which were injected with mismatch or antisense LNA gapmers. Scale bar: 100 μm. Note that at stage 12, PGCs injected with the antisense LNA gapmer are relatively large and contain the germ plasm beneath the cell membrane, compared with perinuclear localization of the germ plasm in PGCs with the mismatch LNA gapmer. (C) Ratio of PGCs with three localization patterns of germ plasm from stage 12 embryos injected with mismatch or antisense gapmers, respectively. According to the localization patterns, 120 and 124 PGCs with mismatch or antisense gapmers, respectively, were classified into three groups: cortex, perinucleus and intermediate (cortex-perinucleus) shown at the bottom of the panel.
Functional substitution of *vasa* for *DEADSouth*

In *Drosophila*, the *vasa* gene encodes a DEAD-box RNA helicase that is required for germ cell development (Styhler et al., 1998). *vasa* homologs have been identified in various animals and are expressed specifically in their germline. Also, in *Xenopus*, *vasa*-like gene 1 (*XVLG1*) was identified as a *vasa* homolog (Komiya et al., 1994). However, *XVLG1* is not expressed in a germline-specific manner (Ikenishi and Tanaka, 2000). Instead, *DEADSouth* was identified as another germline-specific DEAD-box RNA helicase (MacArthur et al., 2000). Therefore, we examined whether *vasa* genes could functionally substitute for the *DEADSouth* gene in *Xenopus*. We prepared two constructs, in which the ORF of *Mvh* or *XVLG1* was fused with *DEADSouth* 3′ UTR for PGC-specific expression. Fertilized eggs were injected with the mRNA of *Mvh-DS 3′ UTR* or *XVLG1-DS 3′ UTR* in addition to the antisense LNA gapmer and *v-DS* as a PGC tracer, and allowed to develop until stage 32 to externally observe PGCs (Fig. 6). Compared with mismatch gapmer-injected embryos (PGC number, 16.8±2.5/embryo), antisense gapmer-injected embryos showed significantly reduced PGC numbers (6.4±2.3/embryo) (Fig. 6A). Surprisingly, this reduction was restored by co-injection with *Mvh-DS 3′ UTR* (13.0±4.4/embryo, *P* < 0.001). Co-injection with *XVLG1-DS 3′ UTR* showed similar results. PGC numbers in embryos injected with the mismatch gapmer, antisense gapmer, and antisense gapmer plus *XVLG1-DS 3′ UTR* were 17.1±1.6, 7.6±1.1 and 12.4±0.9, respectively (Fig. 6B). Although the rescue by *XVLG1-DS 3′ UTR* was partial, it appeared to be similar to that by *DS-full* (12.7±1.6/embryo). These results indicated that the function of *DEADSouth* in PGC development can be substituted, at least in part, by another RNA helicase such as VASA.

These constructs contained the 3′ UTR of *DEADSouth* mRNA to ensure PGC-specific expression. Therefore, we evaluated the effect of the 3′ UTR. Two constructs, in which *DEADSouth* or *XVLG1* ORFs were followed by the *XVLG1* 3′ UTR, were generated and subjected to assays to assess their rescue from *DEADSouth* knockdown. Surprisingly, both *DEADSouth* and *XVLG1* ORFs fused with the *DEADSouth* 3′ UTR rescued from *DEADSouth* knockdown, but could not when they were fused with the *XVLG1* 3′ UTR (*DS-XVLG1 3′ UTR*, 8.2±1.3/embryo; *XVLG1-XVLG1 3′ UTR*, 7.7±1.3/embryo). This observation indicates that the rescue depends on the *DEADSouth* 3′ UTR, because there is no distinct difference between the translational activities of the two mRNAs (data not shown).

Localization of *DEADSouth* mRNA and protein to germ plasm via the 3′ UTR

To elucidate whether *DEADSouth* mRNA is localized to germ plasm via its 3′ UTR, we examined the behavior of fluorescent-labeled
mRNA with the DEADSouth 3′UTR in PGCs. Nonsense mRNA encoding a partially deleted XLG1 ORF followed by the DEADSouth 3′UTR (XLG1A-DS 3′UTR) or XLG1 3′UTR (XLG1A-XLG1 3′UTR) was labeled with Cy3 fluorescent dye in vitro and injected at the vegetal pole of fertilized eggs from mitochondria supplied maternally (Taguchi et al., 2012). Together with a PGC visualization technique suitable for observation from about stage 12 onward (Kataoka et al., 2006), it was possible to trace PGCs in a living embryo during early development. Thus, we investigated the function of DEADSouth in the context of PGC development.

First, we investigated the intracellular localization of DEADSouth protein by expression of DEADSouth tagged with mCherry fluorescent protein. DEADSouth-mCherry fusion protein was detected in the germ plasm of PGCs at stage 12 (Fig. 1A–E). Unfortunately, we could not detect the localization at earlier stages because of low expression levels of the fusion protein. In zebrafish, vasa mRNA, but not its protein, is localized to the germ plasm before MBT and the protein is localized to the germ plasm at later stages (Knaut et al., 2000). As a control, mCherry protein was not localized to the germ plasm and distributed throughout PGCs, although mRNA was injected into the cortex of the vegetal pole where the germ plasm was present (Fig. 1F–J). These results suggest that DEADSouth protein includes a signal for localization or anchorage to the germ plasm. DEADSouth-mCherry fusion protein was also detected in the nuclei of PGCs. Considering that germ plasm contains determinants for PGC differentiation in Drosophila (Ikenmeyer and Mahowald, 1974; Okada et al., 1974) and Xenopus (Buehr and Blackler, 1970; Tada et al., 2012), germ plasm migration to the nucleus at this stage suggests that some transport from the germ plasm to the nucleus triggers PGC differentiation. Because DEADSouth is an RNA helicase, it is feasible that DEADSouth is involved in the transport of RNAs from the germ plasm into the nucleus.

In mammals, GRTH/DDX25 is a homolog of DEADSouth, which is expressed exclusively in testes (Sheng et al., 2003) and essential for completion of spermatogenesis (Tsai-Morris et al., 2004). DDX25 protein is a shuttle protein associated with RNA and localized to chromatid bodies in spermatogonia, which have a morphology similar to that of germ plasm (Sheng et al., 2006). Amino acids regions 61–74 and 101–114 of mouse DDX25 protein have been identified as a nuclear export signal via CRM1 protein and a nuclear localization signal, respectively. We found some similarity in these regions between GRTH/DDX25 and DEADSouth, suggesting that DEADSouth functions as a shuttle protein between the germ plasm and nucleus. However, detailed studies using mutated constructs are required to evaluate whether these regions of DEADSouth play important roles in intracellular transport in PGCs. Translocation of DEADSouth from the germ plasm to the nucleus may be also regulated by phosphorylation (Sheng et al., 2006). Localization of endogenous DEADSouth protein also remains unknown.

To determine the possible function of DEADSouth in PGC development, we performed overexpression experiments in which in vitro synthesized DEADSouth mRNA including the 3′UTR was injected into fertilized eggs. Injected embryos

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**Fig. 7.** DEADSouth 3′UTR localizes the mRNA to the germ plasm.

Localization of Cy3-labeled XLG1A-DS 3′UTR mRNA (A–C) and Cy3-labeled XLG1A-XLG1 3′UTR mRNA (D–F) in PGCs at stage 7. PGCs were isolated from mito-EGFP embryos injected with these mRNAs and observed for Cy3 signals (mRNA, in red) and germ plasm (mitochondria, in green). Note that the Cy3 signal is superimposed on the germ plasm in (A–C). Scale bar: 10 μm.
showed normal PGC numbers and sizes at stages 12 and 20 (Fig. 2E; supplementary material Fig. S1). Translocation of germ plasm from the cortex to the perinuclear region appeared to occur normally before MBT (supplementary material Fig. S2). At stage 32, DEADSouth-overexpressing embryos showed a loss of PGCs, but the PGCs were normal in size and localization of the germ plasm. No PGCs were detected within the endodermal mass of DEADSouth-overexpressing embryos, indicating that the loss of PGCs was not due to a failure to migrate. Abnormal PGC migration is often observed at the tailbud stage in knockdown embryos for germ plasm-specific genes such as Xdazl, XDead end and XGRIPI.1 (Houston and King, 2000; Horvay et al., 2006; Tarbashevich et al., 2007). Recently, it has been shown that loss of PGCs in Nanos1-depleted embryos is due to apoptosis (Lai et al., 2012). Our results also suggest that loss of PGCs in DEADSouth-overexpressing embryos is, at least partially, due to apoptosis (Fig. 2F). Early Xenopus embryos might have a mechanism by which aberrant PGCs are eliminated by apoptosis.

However, PGC abnormalities were observed at stage 12 after depleting DEADSouth mRNA by injection of an antisense LNA gapmer. DEADSouth knockdown resulted in a decrease in number and increase in size of PGCs (Fig. 4; Fig. 5A), but no significant decrease in the total amount of germ plasm as indicated by the Xpat mRNA level (Fig. 3B). The diameter of DEADSouth-knockdown PGCs at stage 12 was about 1.5-fold longer than that of control PGCs, indicating that the volume of DEADSouth-knockdown PGCs was about twice as large as that of the control (Fig. 5A). These results indicate that knockdown of DEADSouth causes inhibition of PGC division. In addition, the knockdown phenotype was rescued by expression of normal DEADSouth, but not mutated DEADSouth encoding a putative inactive RNA helicase, suggesting that helicase activity is involved in PGC division.

Because we did not observe an abnormal cleavage pattern around the vegetal pole or defects in the size and number of PGCs in DEADSouth-knockdown embryos until stage 7 (data not shown), we concluded that PGC division was inhibited between stages 7 and 12. Interestingly, at that time, germ plasm moves from the cortex to perinucleus in PGCs followed by PGC specification (Whittington and Dixon, 1975; Venkatarama et al., 2010). We also found that aberrant translocation of germ plasm occurred in DEADSouth-knockdown embryos at stage 12, but not at stage 7 (Fig. 5B). In particular, germ plasm appeared to be expanded in the cortex of DEADSouth-knockdown PGCs at stage 12. Such expanded germ plasm was observable more often in larger PGCs. The ratio of PGCs with germ plasm at an aberrant position was higher in DEADSouth-knockdown embryos than that in control embryos (Fig. 5C). In DEADSouth-knockdown embryos, PGCs with germ plasm at the intermediate position between the cortex and perinucleus were also observable. Because translocation of germ plasm from the cortex to perinuclear region occurs prior to symmetric PGC division, we concluded that DEADSouth knockdown causes aberrant translocation of germ plasm, resulting in inhibited or delayed PGC division, which suggests a link between germ plasm translocation and PGC division. At later stages (stages 20 and 32), the size of PGCs in DEADSouth-knockdown embryos appeared to be closer to that of the control. Their germ plasm was also present at the normal position (data not shown). Such a phenotype may have been due to incomplete knockdown. Taken together, DEADSouth RNA helicase is required for proper PGC development. In particular, it is essential for germ plasm translocation required for symmetric division of PGCs after MBT. Moreover, translocation of germ plasm may be controlled via translational regulation by DEADSouth RNA helicase, which is similar to eIF4A (MacArthur et al., 2000).

GRTH/DDX25 protein is present in the chromatoid bodies of spermatids. In mice, GRTH/DDxs25-null mutation causes complete arrest of spermiogenesis and remarkably diminishes chromatoid bodies in round spermatids at the steps before arrest (Tsai-Morris et al., 2004). In addition, protein components of chromatoid bodies, such as MVH and MIWI, are completely excluded from chromatoid bodies in the knockout mouse (Sato et al., 2010). Together with its function as a shuttle protein, these findings suggest that GRTH/DDX25 is essential to govern the structure of chromatoid bodies for storage and processing of mRNAs. Similarly, DEADSouth protein may be essential to maintain the structure of germ plasm in Xenopus, which is supported by the observation that expanded germ plasm was present in DEADSouth-depleted PGCs.

Defects in DEADSouth-knockdown embryos were rescued by expression of XLG1 or MVH RNA helicase belonging to the VASA/DDX4 family, which is different from the DDX25 family, possibly because of functional redundancy among DEAD-box RNA helicases. In mice, although Mvh is expressed both in the ovary and testis, no defect is detected in the oocytes of Mvh-knockout mice, suggesting that another helicase, such as PL10/DDX3, expressed in the ovary has functional redundancy with MVH (Tanaka et al., 2000). Recently, it was reported that VASA/DDX4 plays a role in cell cycle progression and germline development in Drosophila (Pek and Kai, 2011a) and sea urchin (Yajima and Wessel, 2011). In Drosophila, VASA appears to be involved in regulation of mitotic chromosome condensation/segregation in germline cells in a translation-independent manner. Furthermore, in human and Drosophila somatic cells, mitotic chromosome segregation is regulated similarly by Belle/DDX3 RNA helicase, instead of VASA/DDX4 (Pek and Kai, 2011b), suggesting functional redundancy between DDX3 and DDX4 helicases in somatic and germline cells, respectively. In Xenopus, XLG1, a VASA homolog, appears to be involved in general cell specification and proliferation, rather than only in germline development, based on the expression pattern and perturbation of protein function (Ikenishi and Tanaka, 1997; Ikenishi and Tanaka, 2000). Our results suggest that XLG1 and DEADSouth share VASA functions, including general and germline-specific functions, respectively, in early Xenopus development. To clarify such a functional redundancy and sharing between DEADSouth and VASA/DDX4, rescue experiments from vasa deficiency by the DEADSouth gene would be required. In addition, it will be necessary to determine the function of Centroid that is another DEAD-box RNA helicase gene expressed in germ plasm (Kloc and Chan, 2007).

Surprisingly, rescue from defects in DEADSouth-knockdown embryos depended on the 3’UTR of DEADSouth (Fig. 6B). mRNA with the 3’UTR of DEADSouth was localized to the germ plasm, but mRNA with the XLG1 3’UTR was not localized to the germ plasm (Fig. 7). This finding indicates that the 3’UTR of DEADSouth contains a signal to anchor and/or localize the mRNA. Because DEADSouth protein also appears to have signal for localization to the germ plasm, the DEADSouth 3’UTR supports effective localization of the DEADSouth protein. We have previously demonstrated that the DEADSouth 3’UTR
contains information for PGC-specific protein expression, probably via microRNA (Kataoka et al., 2006). It is very interesting to elucidate the relationship between PGC-specific expression and localization of mRNA in the context of transcriptional regulation.

Finally, we demonstrated the utility of an antisense LNA gamper with micro-EGFP transgenic embryos to study the functions of germ plasm-specific genes. In addition to previous methods, such as the host-transfer technique (Heasman et al., 1991) and PGC labeling with EGFP (Kataoka et al., 2006), these tools enable us to analyze the development of PGCs in Xenopus.

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Competing Interests

The authors have no competing interests to declare.

References


