

Dnd Is a Critical Specifier of Primordial Germ Cells in the Medaka Fish

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SUMMARY

Primordial germ cell (PGC) specification occurs early in development. PGC specifiers have been identified in *Drosophila*, mouse, and human but remained elusive in most animals. Here we identify the RNA-binding protein Dnd as a critical PGC specifier in the medaka fish (*Oryzias latipes*). Dnd depletion specifically abolished PGCs, and its overexpression boosted PGCs. We established a single-cell culture procedure enabling lineage tracing in vitro. We show that individual blastomeres from cleavage embryos at the 32- and 64-cell stages are capable of PGC production in culture. Importantly, Dnd overexpression increases PGCs via increasing PGC precursors. Strikingly, *dnd* RNA forms prominent particles that segregate asymmetrically. Dnd concentrates in germ plasm and stabilizes germ plasm RNA. Therefore, Dnd is a critical specifier of fish PGCs and utilizes particle partition as a previously unidentified mechanism for asymmetric segregation. These findings offer insights into PGC specification and manipulation in medaka as a lower vertebrate model.

INTRODUCTION

Primordial germ cells (PGCs) are germ stem cells capable of generating precursors of eggs and sperm and thus offer the basis for reproduction and fertility (Ko et al., 2010; Lin, 2007, 2012). PGC specification demarcates the soma-germline separation and thus sets a balance between individual life and species continuity. PGCs form early in development and migrate into the developing gonad (Wylie, 1999). In adult animals, gonadal germ cells undergo meiosis and produce eggs and sperm. PGC development has been extensively studied in several model organisms including *Drosophila*, *Caenorhabditis elegans*, and *Xenopus* (Houston and King, 2000), zebrafish (Lin et al., 1992; Raz, 2003), and mouse (Hayashi et al., 2011; Ohinata et al., 2005; Saitou et al., 2002; Tam and Zhou, 1996). In these model invertebrates and lower vertebrates, PGCs are cell autonomously preformed by maternally supplied germ plasm, a membrane-less organelle composed mainly of RNA-binding proteins and their mRNAs (Houston and King, 2000; Raz, 2003). In mouse, PGCs are epigenetically induced by cell-cell interactions in the absence of germ plasm (Tam and Zhou, 1996), where signaling molecules such as bone morphogenetic factor 4 play a critical role (Ying et al., 2001).

Dozens of genes essential for PGC development are known in the model invertebrates and lower vertebrates (Houston and King, 2000); clearly defined specifiers of the PGC fate has, however, been limited to *Drosophila*. In this or-

ganism, *oskar* acts as the PGC specifier, as it is necessary for PGC formation and more importantly, sufficient for ectopic PGC induction in a dose-dependent manner (Ephrussi and Lehmann, 1992). However, *osk* is restricted to certain insects. An evolutionarily conserved gene, *piwi*, is also necessary for PGC formation and sufficient for tripling the PGC number but incapable of ectopic PGC induction (Megosh et al., 2006). In vertebrates, *piwi* is dispensable for PGC specification but essential for subsequent PGC development, such as spermatogenesis in mouse (Deng and Lin, 2002), germ cell maintenance in zebrafish (Houwing et al., 2007), and PGC migration in medaka (Li et al., 2012). In mice, *blimp1* (encoded by *prdm1*) and *prdm14* are transcriptionally induced by BMP4 in the epiblast at E6.25, which together with *tfap2c* constitute a tripartite genetic network to induce the PGC fate in vivo (Magnusdottir et al., 2013; Ohinata et al., 2005) and in vitro from embryonic stem (ES) cells (Magnusdottir et al., 2013; Nakaki et al., 2013). In human, SOX17 has most recently been identified as a critical specifier of PGCs in ES cells (Irie et al., 2015). Accumulated data from *Drosophila*, mouse, and human suggest that PGC specifiers show remarkable diversity and do not follow the evolutionary history, although many genes involved in subsequent germ cell development are highly conserved across animal phyla.

A vertebrate-specific germ gene, *dead end* (*dnd*), was first identified in zebrafish as a germ plasm component encoding an RNA-binding protein crucial for PGC migration

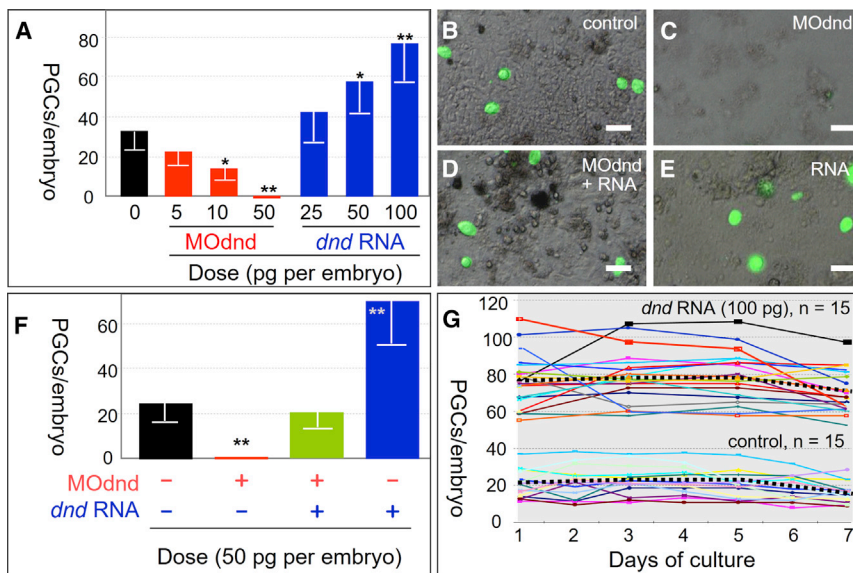


Figure 1. Dnd Dosage Determines the PGC Number

(A) Number of PGCs in embryos. Data are means \pm SD (bars) of 12–21 embryos at stage 20.

(B–E) PGCs from midblastula cells at 2 days of culture. Scale bars, 20 μ m.

(F) Number of PGCs at day 2 post culture from ≥ 12 embryos. *Significant difference ($p \leq 0.05$); **very significant difference ($p \leq 0.01$) compared with non-injected control.

(G) Time course of PGC numbers from individual blastula embryos in culture.

and survival (Tzung et al., 2015; Weidinger et al., 2003). Mouse *dnd* mutations do not prevent PGC formation (Youngren et al., 2005). The medaka fish (*Oryzias latipes*) is an excellent model for studying vertebrate development (Wittbrodt et al., 2002), stem cells (Centanin et al., 2011; Hong et al., 1996, 1998b), germ cells, and reproductive technologies. This fish has haploid ES cells capable of whole-animal production by semicloning (Yi et al., 2009), male germ stem cells capable of test-tube production (Hong et al., 2004a), and transgenic lines for PGC visualization (Li et al., 2009; Tanaka et al., 2001). Unusually, maternal germ plasm components distribute widely in medaka (Shinomiya et al., 2000; Xu et al., 2009), rather than locally as in zebrafish (Weidinger et al., 2003; Yoon et al., 1997). This suggests the presence of an unknown key factor that determines the PGC fate. Here we identify Dnd as such a critical PGC specifier in medaka. Interestingly, *dnd* RNA uses particle formation and partition as a mechanism for asymmetric segregation and cell fate decision in early developing embryos. These results provide insights into our understanding of PGC formation and manipulation in medaka as a lower vertebrate model.

RESULTS

Dnd Dosage Determines the PGC Number In Vivo

Targeted *dnd* disruption in medaka embryos led to normal survival and development to adulthood (Wang and Hong, 2014). The *dnd* mutant medaka adults are sterile and not suitable for studying its role in PGC development during embryogenesis. Conditional knockout of genes essential

for early development such as PGC formation is not yet available in fish. We thus adopted direct embryo microinjection of *dnd*-targeting morpholino oligos for gene depletion and *dnd* mRNAs for gene overexpression. Two morpholino oligos were used for *dnd* depletion: MOdnd targets the medaka *dnd* mRNA and inhibits its translation, and MOddm is a mutant derivative of MOdnd by introducing four mismatches (Figure S1A). For overexpression, mRNAs *dnd:ch* and *dndΔ1:ch* were synthesized from pCSdnd:chDD and pCSdndΔ1:chDD (Figure S1B); the former encodes a cherry fluorescent protein-tagged wild-type Dnd and the latter a tagged deletion mutant Dnd. The morpholino oligos and mRNAs were microinjected alone or in combination into one-cell embryos of transgenic medaka NgVg expressing GFP specifically in PGCs (Hong et al., 2010; Li et al., 2009).

A medaka embryo at stages 18–22 has ~ 32 PGCs that are recognizable by GFP expression (Figure 1A). Injection of mismatch-containing MOddm had no effect on the PGC number (Figure S2A). Remarkably, *dnd* depletion by injection with 50–100 pg of MOdnd caused the complete absence of PGCs in all ($n = 333$) manipulated embryos (Figures 1A and S2B). When MOdnd was coinjected with *dnd:ch* RNA, the PGC number was rescued (Figure S2C), whereas *dndΔ1:ch* mRNA did not rescue (Figure S2D). Moreover, injection with 50 pg of *dnd:ch* RNA alone was sufficient to increase the PGC number (Figure S2E). Injection of *dnd:ch* RNA at 100 pg considerably boosted the PGC number, so that a significant number of PGCs were located in ectopic sites as well as numerous PGCs in the gonad (Figure S2F). Clearly, altering *dnd* expression by injection of either MOdnd or *dnd:ch* RNA altered the PGC



number, ranging from the complete absence to an increase by ~2-fold (Figure 1A). The effect of MO*dnd* is specific, as it does not affect somatic development by morphological criteria, and *dnd* RNA but not its deletion mutant is capable of phenotypic rescue. Taken together, *dnd* is essential for PGC development and its dosage determines the PGC number in vivo.

Dnd Depletion Does Not Cause PGC Death

To determine whether the absence of PGCs observed from stage 21 onward was due to the absence of PGC formation or loss of established PGCs, we examined NgVg embryos earlier at stage 15 when PGCs are unambiguously visible by the GFP signal. Cell counting in more than 12 embryos revealed that the average PGC number was 22.3, 9.7, 0., and 49.8 in the MO*dnd*-injected control embryos, embryos injected with 20 pg and 50 pg of MO*dnd*, and embryos injected with 50 pg of *dnd:ch* RNA, respectively (Figures S3A–S3E). These results indicate that a PGC decrease and loss in MO*dnd*-injected embryos occurred as early as at stage 15, arguing against PGC loss between stage 15 and subsequent stages. To rule out the possible loss of PGCs at even earlier stages, we examined cell death by acridine orange staining (Li et al., 2009). No difference was observed at the blastula stage between control (Figure S3F) and MO*dnd*-injected embryos (Figures S3G and S3H), indicating that MO*dnd* depletion does not cause cell death. Collectively, the absence of PGC caused by *dnd* knockdown is not due to PGC loss by death prior to observation.

Dnd Dosage Determines the PGC Number In Vitro

In medaka, PGCs can be formed in culture from dissociated single cells of gastrulae (Li et al., 2009) and midblastula embryos (Li et al., 2014). We made use of this cell culture system to analyze the effect of *dnd* dosage on the PGC number in vitro. To this end, NgVg embryos were injected at the one-cell stage with MO*dnd* and/or *dnd:ch* mRNA and dissociated into single cells at the midblastula stage. The dissociated cells were seeded into 96-well plates, one embryo per well, and the cell fate decision was observed at day 2 post culture (dpc). PGC formation was seen in all embryos except for those *dnd*-depleted embryos injected with MO*dnd* (Figures 1B–1E). Cell counting revealed that the PGC number was dependent on Dnd dosage, as Dnd overexpression via injecting 50 pg of *dnd:ch* RNA increased the number of PGCs from 24 to 69, whereas *dnd* knockdown via injecting 50 pg of MO*dnd* led to the absolute absence of PGCs (Figure 1F). We observed a time course change in the PGC number per embryo in culture. In 12 control embryos, the PGC number per embryo ranged from 10 to 38 at 1 dpc and changed during subsequent days of culture (Figure 1G), revealing the individual variability in the PGC number per embryo and dynamics of PGC proliferation

and survival. Injection of *dnd* RNA significantly increased the PGC number ranging from 55 to 109 ($n = 15$) at 1 dpc, without altering the variability and dynamics (Figure 1G). Thus, Dnd dosage controls the PGC number in vitro without influencing PGC proliferation and/or survival.

Dnd Dosage Determines the pPGC Number In Vitro

The first embryonic cells that will produce exclusively germ cells by clonal mitotic divisions are PGCs, and the PGC precursors, which are often morphologically indistinguishable from the surrounding somatic cells, are called presumptive PGCs (pPGCs) (Nieuwkoop and Sutasurya, 1979). At the final stage of fate decision, a pPGC divides to produce one PGC and one somatic cell. Hence, the pPGC number is equal to the initial PGC number (Figure 2A). We developed a procedure for lineage tracing of PGC development in single-cell culture (Figure 2B). Medaka PGCs can be formed in culture from gastrulae (Li et al., 2009) and midblastulae (Li et al., 2014). Cleavage blastomeres from 32-cell embryos are capable of cultivation (Li et al., 2011). We asked whether cleavage blastomeres were capable of PGC formation in vitro. To this end, blastomeres were dissociated from NgVg embryos at the 32- and 64-cell stage and individually seeded into 96-well plates, and PGCs were identified by phenotype and GFP expression. PGCs were found in culture from blastomeres isolated from embryos at the 32-cell stage ($n = 24$; Figures S4A and S4B) and 64-cell stage ($n = 33$; Figures S4C and S4D). Thus, dissociated cleavage blastomeres can form PGC in culture.

We then tested the ability of cleavage blastomeres to form PGCs in single-cell culture. A proportion of blastomeres individually cultured in 96-well plates produced PGCs. These PGCs were first visible at 14 hr of culture by a larger size and faint GFP expression (Figures 2C and C'), which became more distinct with a brighter GFP signal at 48 hr (Figures 2D and D'). We made use of this single-cell culture system to lineage trace pPGCs in vitro. A total of 378 blastomeres from 64-cell NgVg embryos were seeded for single-cell culture; 331 survived PGC observations at day 2 of culture (Table S1), 42 of which were pPGCs (Figure 2E). This suggests that 12.7% of 64-cell blastomeres are pPGC, equivalent to 8.1 pPGCs per embryo at this stage (Figure 2F). These 42 pPGCs produced a total of 181 PGCs, corresponding to 4.3 PGCs per pPGC (Figure 2G). By extrapolation from these values, a medaka is able to produce 37.3 PGCs at 3 dpf (Table S1), comparable with 35.3 for developing embryos (Figure 1A). Hence, ~8 pPGCs are committed in a medaka embryo already at the 64-cell cleavage stage, which demonstrates PGC preformation in this organism.

We analyzed *dnd*-dependent PGC development in single-cell culture. Since *dnd* knockdown causes the complete

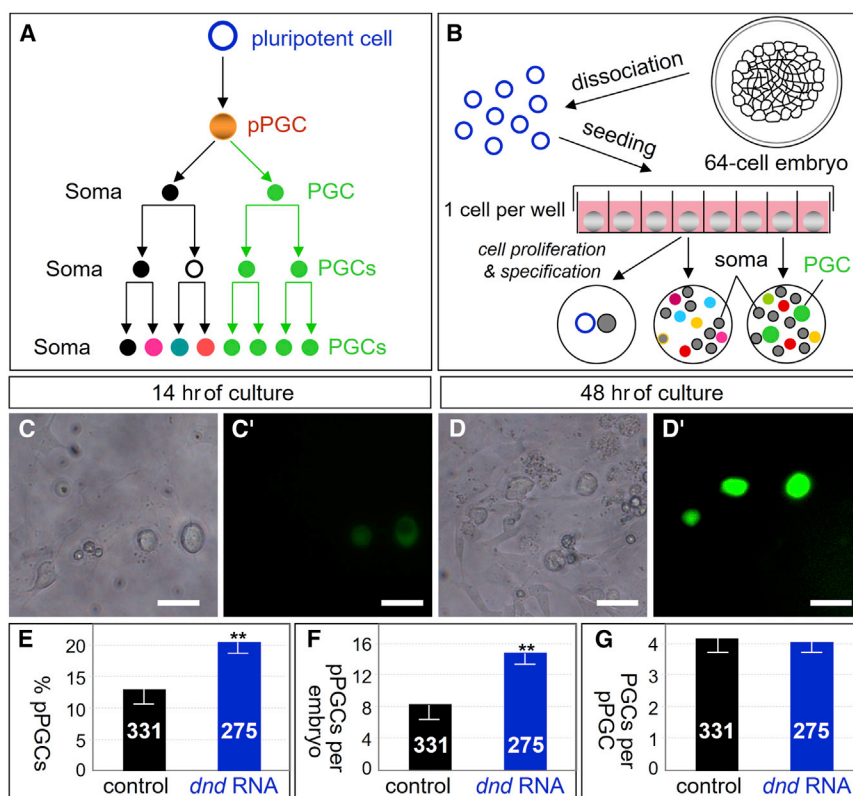


Figure 2. Lineage Tracing of PGC Formation

(A) Conceptual PGC formation. (B) Strategy of single blastomere culture. (C–D') 64-cell embryo blastomeres in single cell culture, showing PGCs (green) at 14 and 48 hr post culture. Scale bars, 50 μ m. (E) Percentage of pPGCs in culture. (F) Number of pPGCs per embryo in culture. (G) Number of PGCs per pPGC in culture. Numbers of blastomeres analyzed are shown in the columns. Data are means \pm SD (error bars) from ≥ 7 independent experiments. **Very significant difference ($p \leq 0.01$) compared with control.

absence of PGCs in vivo and in vitro (Figures 1A, 1F, and S1), we analyzed how *dnd* overexpression enhanced the PGC number. To this end, 64-cell embryos injected at the one-cell stage with 50 pg of *dnd:ch* RNA were used for single-cell culture. A total of 308 blastomeres from *dnd:ch* RNA-injected 64-cell embryos were seeded, 275 survived observation at day 2 of culture (Table S1), 20.7% ($n = 57$) of which were pPGCs (Figure 2E). This corresponds to 13.3 pPGCs per embryo (Figure 2F). Therefore, *dnd* RNA injection at 50 pg increases the pPGCs number by up to 64%. On average, a pPGC from *dnd:ch* RNA-injected embryos produced 4.2 PGCs (Figure 2G), which is fully comparable with that of pPGCs from control embryos. Consequently, the PGC number per embryo increased by 53% to 57.1 upon *dnd:ch* RNA injection. Taken together, Dnd overexpression increases PGCs through increasing pPGCs without increasing PGC survival and/or proliferation.

dnd RNA Forms Particles for Asymmetric Segregation

The medaka *dnd* RNA is specific to germ cells (Liu et al., 2009). We analyzed its temporospatial expression during early embryogenesis. In a majority (91%; $n = 145$) of cleavage embryos, *dnd* RNA existed predominantly in prominent cytoplasmic particles (Figures 3A–3C), which has never been seen for mRNAs of other germ genes such as *boule*, *dazl*, and *vasa* (Xu et al., 2009). As early as in

the one-cell embryo, there were ~ 40 particles of varying shapes and sizes ranging from 1 μ m to 30 μ m, which were not localized to a particular region of the embryo (Figure 3A). In 2- to 16-cell embryos, cells were different in the absence and presence of particles, suggesting irregular asymmetry of particle distribution between daughter cells (Figure 3A). In midblastula embryos comprising $\sim 2,000$ cells, *dnd* RNA was seen in 6–11 prominent particles (Figure 3B). A closer inspection at larger magnification revealed that each prominent particle was frequently associated with 3–7 nuclei (Figure 3C). Until stage 13, *dnd* RNA eventually became dispersed and concentrated in the perinuclear region of PGCs (Figures 3D and 3E). We furthered our observation at better sensitivity and resolution by fluorescence in situ hybridization (FISH). This revealed 6–11 big particles 10–20 μ m in size, ~ 5 medium-sized (6–9 μ m) particles, and ~ 30 small particles ≤ 5 μ m in medaka embryos at the early blastula stage (Figures 4A–4C). At this stage, *dnd* RNA particles were seen in all three to four cell layers on cryosections (Figures 4D–4G'). The *vasa* RNA did not form particles and displayed wide distribution among deep cells (Figures 4G and 4G'). Taken together, maternal *dnd* mRNA exists predominantly in cytoplasmic particles and segregates irregularly into a randomly positioned subset of blastomeres before its perinuclear localization in PGCs.

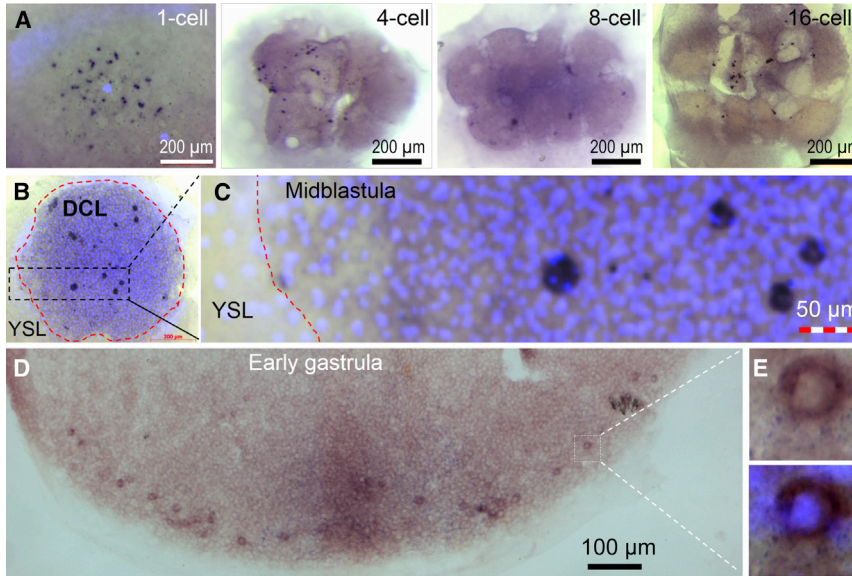


Figure 3. *dnd* RNA Particles

(A) Cleavage embryos at 1- to 16-cell stages showing *dnd* RNA particles and their intercellular distribution.

(B) Midblastula embryo showing prominent *dnd* RNA particles in the deep cell layer (DCL) but absent in the extraembryonic yolk syncytial layer (YSL).

(C) Large magnification of the area framed in (B), highlighting association of prominent particles with multiple nuclei (blue).

(D) Early gastrula embryo squash showing PGCs scattered at the submarginal region.

(E) Large magnification of the boxed area in (D), highlighting perinuclear localization of *dnd* RNA within a PGC.

Dnd Protein Is a Component of Germ Plasm

Many germ cell markers are localized in germ plasm, a membrane-less organelle that is closely associated with mitochondria (Houston et al., 1998; Kloc et al., 1998; Knaut et al., 2000). Our finding that Dnd determines the PGC fate provoked us to examine its intracellular localization. Fusion proteins PF (puromycin acetyltransferase:GFP) and Vas:GFP were exclusively cytoplasmic (Figures 5A and 5B), whereas Dnd:Ch was both nuclear and cytoplasmic in ES cells (Figures 5A and 5B) and early embryos (Figures 5C and 5D). We performed correlative light and electron microscopy to precisely determine the intracellular localization of Dnd protein. Embryos were microinjected with the *dnd:ch* RNA, single blastomeres were dissociated at the blastula stage and fixed by high-pressure freezing to preserve cellular architecture. After freezer substitution, ultrathin sections were transferred onto a copper grid for light microscopy (LM) and electron microscopy (EM). Dnd:Ch was seen mostly in cytoplasmic particles of 0.5–5 μm in size under LM, which appeared as dense bodies under EM (Figures 5E and 5F). At a large magnification, these dense bodies lacked a membrane, possessed granulo-fibrillar structures, and were surrounded by many mitochondria (Figure 5G). These features are diagnostic of germ plasm that has been described in embryos of diverse organisms including *Xenopus* (Houston et al., 1998) and zebrafish (Knaut et al., 2000). Thus, Dnd protein is a component of germ plasm present in certain blastomeres prior to PGC formation.

Dnd Stabilizes Germ Plasm RNA

In diverse animals, early development is controlled by maternal gene products in the absence of zygotic transcription. miRNAs play an essential role in maternal RNA clear-

ance (Schier, 2007). Specifically, zebrafish MiR-430 promotes degradation of maternal mRNAs (Giraldez et al., 2006), and this degradation is suppressed by Dnd in cultured human germ cells and zebrafish PGCs through inhibiting miRNAs to target mRNAs (Kedde et al., 2007). Dnd interacts with its target RNAs via the DND-binding site. This binding protects the target RNAs from degradation by the MiR430 pathway (Giraldez et al., 2006). The binding sites for Dnd and MiR430 are present in the medaka *dnd* (Figure 6A; Figure S5A) and the zebrafish *nos3* (initially *nos1*) (Kedde et al., 2007; Figure S5B). To determine the effect of Dnd binding on target RNAs in medaka, we exploited reporter assays by microinjection of *gfpMiR430* and *gfpNOS* reporter RNAs (Figures 6A–6C). The *gfpMiR430* RNA contains three copies of the MiR-430 target site (Giraldez et al., 2006). Upon coinjection into medaka embryos at the one-cell stage, GFP expression from injected *gfpMiR430* RNA was monitored at stages 16 and 30. A preferential loss of GFP expression in many somatic cells was observed upon coinjection with *dnd:ch* but not *dnd Δ 1:ch* RNA, whereas coinjection of *MOdnd* reduced GFP expression to a barely detectable level (Figure 6B). A similar and more evident result was obtained by injection of *gfpNOS* reporter RNA (Figure 6C), which encodes GFP and contains the zebrafish *nos3* 3'-UTR capable of localization in PGCs (Koprunner et al., 2001). Thus, medaka Dnd regulates the level of germ plasm mRNA translation.

DISCUSSION

In this study we provide several lines of evidence that Dnd serves as a critical PGC specifier in medaka. First, *dnd*

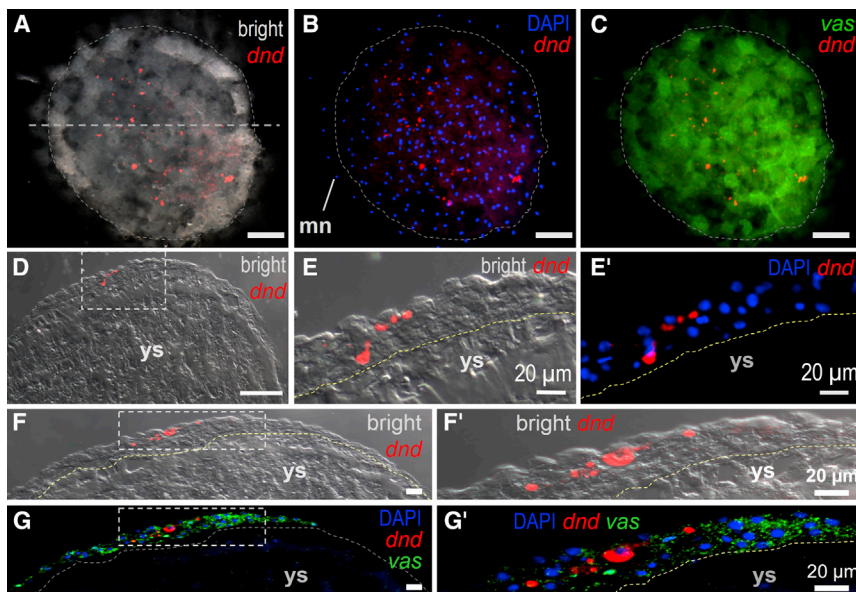


Figure 4. Asymmetric Segregation of *dnd* RNA Particles

(A–C) Early blastula with one layer of marginal cell nuclei (mn).

(D) Cryosection through the broken line in (A).

(E and E') Large magnification of the boxed area in (D), showing *dnd* RNA in all cell layers but absent in the yolk sac (ys).

(F and G) Early blastula cryosection.

(F' and G') Large magnification of the boxed areas in (F) and (G), highlighting fewer *dnd*-positive cells than *vas*-positive cells.

Scale bars, 100 μ m unless indicated.

knockdown is capable of specifically and completely abolishing PGC formation and its overexpression is sufficiently capable of inducing ectopic PGCs in embryos and significantly increasing the PGC number. Second, we reveal that *dnd* overexpression boosts PGCs via increasing the number of pPGCs without increasing PGC proliferation and survival in vitro. Third, we show that *dnd* RNA forms prominent particles that irregularly segregate, and thus limits the number of cells for PGC fate decision. Fourth, correlative microscopy reveals that Dnd protein is a component of germ plasm. Finally, we present evidence that Dnd stabilizes germ plasm RNAs essential for PGC development. Our findings pinpoint the mechanism by which Dnd exerts its functions: Dnd specifies the PGC fate through specifying pPGCs and thus determines the initial PGC number. *dnd* is a vertebrate-specific gene that is conserved from fish to mammals in germ cell-specific expression and requirement (Weidinger et al., 2003). In zebrafish and mouse, *dnd* knockdown or mutation does not prevent PGC formation. Knockdown of zebrafish *dnd* and several other germ genes leads to defects in PGC migration and/or survival but does not entirely prevent PGC formation, and their overexpression by RNA injection is unable to markedly induce additional PGCs (Knaut et al., 2000; Koprunner et al., 2001; Weidinger et al., 2003). In medaka, *vasa* knockdown severely affects PGC migration and *vasa* overexpression does not increase the PGC number (Li et al., 2009). Together with the fact that zebrafish null mutant for *nos3* normally specifies PGCs and is defective in maintaining oocyte production (Draper et al., 2007), all the germ genes functionally analyzed so far in fish embryos act primarily on subsequent steps of PGC develop-

ment, but not the first step, namely PGC fate decision. An exception to this is the zebrafish *buc* (Bontems et al., 2009). *buc* overexpression is able to increase the PGC number by up to ~50%. *buc* cannot be a PGC specifier, as it is highly expressed in early embryos and its RNA distributes widely in all blastomeres rather than co-segregating with pPGCs. Our finding that *dnd* acts as a PGC specifier in medaka reveals a previously unidentified role for *dnd* in the initial step of PGC development, namely PGC fate decision rather than subsequent proliferation and survival, because the number of PGCs produced by each pPGC remains unchanged between normal and *dnd* RNA-injected embryos. It remains to be elucidated whether *dnd* acts as a PGC specifier also in other organisms.

Zebrafish makes use of the preformation mode for PGC specification (Raz, 2003). Two previous studies suggest that PGC formation occurs cell autonomously also in medaka (Herpin et al., 2007; Li et al., 2014). Our work here corroborates and extends these studies by providing convincing evidence that, as early as at cleavage stages, the medaka embryo contains pPGCs capable of PGC formation in single-cell culture. Therefore, PGC preformation operates in medaka as in zebrafish and *Drosophila*.

A striking finding from this study is how medaka segregates *dnd* as a cell fate specifier in the absence of a cellular mechanism for asymmetric segregation. In all organisms with PGC preformation, germ plasm components are either prelocalized to the posterior pole of the oocyte as in *Drosophila*, or localized and asymmetrically segregated in early embryos as in *C. elegans*, *Xenopus*, and zebrafish (Raz, 2003; Wylie, 1999). Our results in this study together with accumulated data reveal the presence of two distinct

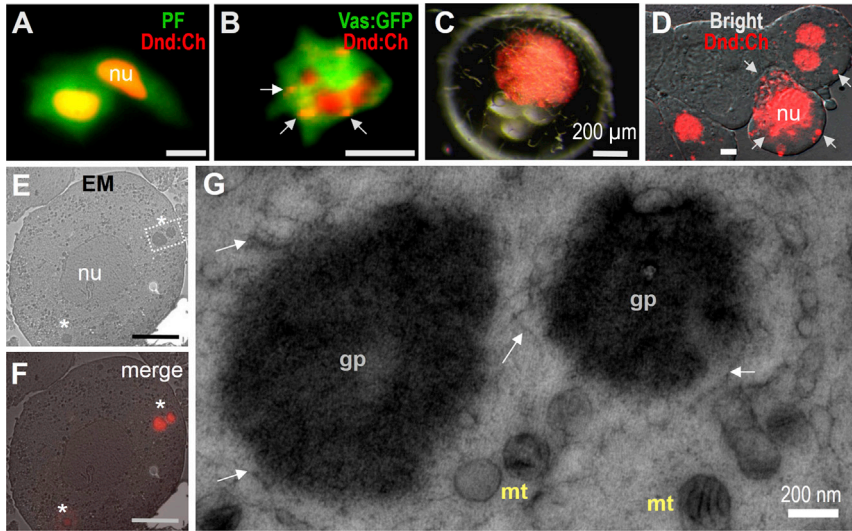


Figure 5. Intracellular Dnd Localization

(A and B) ES cells. (C) Midblastula injected with *dnd:ch* RNA showing Dnd:Ch in many cells. (D) Cells dissociated from the midblastula shown in (C), highlighting Dnd:Ch in the nucleus (nu) and cytoplasmic particles (arrows). (E–G) Correlative microscopy, (E) EM micrograph showing cellular structure and dense bodies (asterisks). (F) Correlative micrographs between fluorescent LM and EM. (G) Large magnification of the area framed in (E), highlighting the fine structure of germ plasm (gp) as dense bodies in association with mitochondria (mt). Germ plasm lacks a membrane and has many granulofibrillar structures (arrows). Scale bars, 10 μ m unless indicated.

modes for asymmetric segregation of germ plasm RNAs in lower vertebrates as exemplified by zebrafish and medaka (Figure 7). In the zebrafish mode, RNAs of *dnd*, *nanos*, and *vasa* are co-localized in four cells that are regularly positioned and committed to the PGC fate (Raz, 2003; Wei-

dingler et al., 2003). In the medaka mode, RNAs of all germ genes studied so far, including *boule*, *dazl*, and *vasa* (Li et al., 2009; Shinomiya et al., 2000; Xu et al., 2009) as well as *piwi* (Li et al., 2012), lack localization but exhibit wide distribution, which apparently suggests the absence

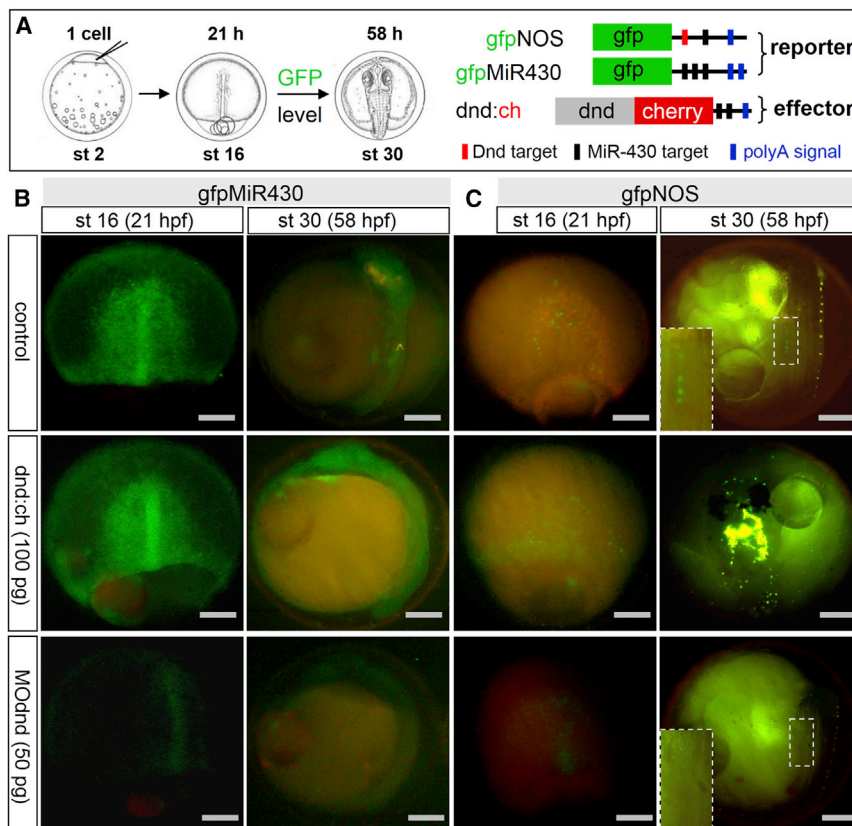


Figure 6. Dnd Protects Germ Plasm RNAs from Degradation

(A) Strategy of a reporter assay. Embryos at the one-cell stage were injected with RNAs from the reporter and effector constructs illustrated and monitored for GFP expression at stages indicated. (B) Dnd stabilizes *gfpMir430* RNA. (C) Dnd stabilizes *gfpNOS* RNA. Scale bars, 200 μ m.

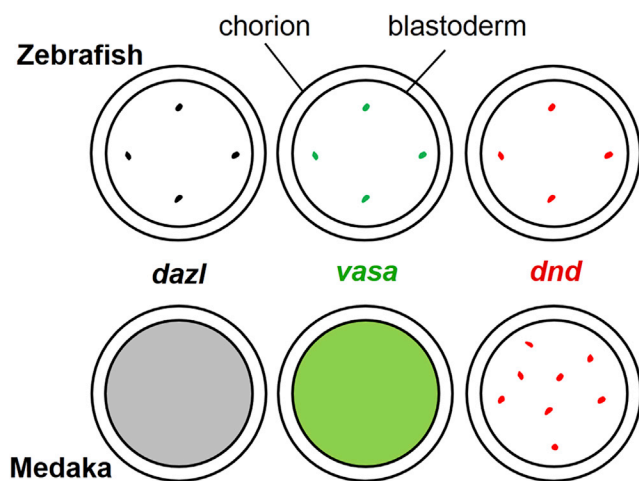


Figure 7. Modes of Germ Plasm RNA Segregation in Fish

Shown here are two distinct modes that regulate the intercellular distribution of germ plasm RNAs such as *dazl* (black), *vasa* (green), and *dnd* (red) RNAs in the embryo until the blastula stage. In the zebrafish mode, germ plasm RNAs are co-localized and asymmetrically segregated into four dots. In the medaka mode, *dazl* and *vasa* are widely distributed, whereas *dnd* is randomly segregated as particles into a subset of cells for the PGC fate decision.

of a cellular mechanism for prelocalization or localized segregation of maternal germ plasm components in medaka. This study has established that *dnd* is a unique exception to wide distribution. In the absence of such a cellular mechanism, the *dnd* RNA makes use of particle formation and partition as a mechanism for asymmetric segregation into pPGCs to specify the PGC fate.

A major challenge in analyzing cell fate decision is the absence of markers to trace steps before bona fide fate decision. An ideal situation is to put a cell under defined conditions and monitor its changes into a state diagnostic of a particular cell. In this study, we have established a single-cell culture system and a procedure to lineage trace PGC formation. This enables us to continuously monitor PGC development by precisely counting the number of potential pPGCs per 64-cell embryo as early as at 4 hr post fertilization, long before 13 hpf when PGCs become visible. We anticipate that this in vitro lineage-tracing system will also be applicable to other cell lineages and other organisms.

An increase in the PGC number caused by *dnd* RNA injection does not exceed 3-fold, even if a high level of Dnd expression occurs in many cells. It is likely that an additional factor essential for PGC specification exists in medaka, which shows expression and PGC-specifying activity similar to *dnd* but *dnd*-independent asymmetric segregation. Alternatively, PGC-competent cells are limited in the medaka embryo for the ultimate fate decision. It is also possible that only a limited number of PGCs are

permissive for embryonic development and survival until observation for PGCs. This possibility is small, because *dnd* RNA-injected embryos are not different from controls in survival rate and cell death as well as survival rate of their blastomeres in single-cell culture. A similar situation has also been recorded for the *Drosophila* PGC specifier *oskar* in vivo (Ephrussi and Lehmann, 1992) and the human PGC specifier Sox17 in vitro (Irie et al., 2015).

During early development of diverse animals, miRNAs play an essential role in maternal-zygotic transcription and PGC development (Giraldez et al., 2006; Schier, 2007). Through inhibiting the accessibility of miRNAs including MiR-430 to target mRNAs, Dnd protects maternal RNAs from clearance in cultured human germ cells and PGCs of zebrafish (Kedde et al., 2007) and medaka (Tani et al., 2009). In this study, by reporter assays we reveal that Dnd counteracts with miRNA-430 to regulate maternal RNA turnover and translation for medaka PGC formation. This suggests that Dnd protects germ plasm RNAs from miRNA-mediated degradation to ensure PGC specification.

Our identification of Dnd as a critical PGC specifier in medaka provides insights into cell fate decisions during germline-soma separation in medaka as a lower vertebrate model. Altering Dnd activity could be a strategy for analyzing and manipulating germ cell formation. Our finding that *dnd* RNA asymmetrically segregates via particle partition reveals the presence of a previously unidentified mechanism that controls asymmetric segregation of macromolecules in early development.

EXPERIMENTAL PROCEDURES

Fish

Work with fish was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Advisory Committee for Laboratory Animal Research in Singapore and approved by this committee (permit number 27/09). Medaka was maintained under an artificial photoperiod of 14-hr light and 10-hr dark at 26°C (Hong et al., 1996, 2010, 2012). Medaka strains (HB32C and af) and transgenic lines (Ng and Vg) were described previously (Hong et al., 2010). Ng and Vg express GFP specifically in germ cells from the medaka *nanos* and *vasa* promoters, respectively (Li et al., 2009). Vg embryos were produced by crossing homozygous Vg males to non-transgenic females. NgVg were produced by crossing homozygous Ng males to homozygous Vg females. Embryos were reared and staged as described (Li et al., 2009).

Morpholino Oligos

Two antisense morpholino oligos (Figure S1A), MOdnd (CAC CAC CTT GCT CTG ATT GTC CAT C) and MOddm containing four mismatches (CAC CAC CTT GCT tTG gTT aTC CAT c, mismatches underlined), were synthesized (Gene Tool) and dissolved in water at 10 mg/ml as stock for microinjection as described (Li et al., 2009).



Plasmids

Plasmids pGEMdnd and pCSdnd:chNS (Liu et al., 2009), pGEMvas and pCVvas:gfpNS (Li et al., 2009), and pCVpf expressing a fusion between puromycin acetyltransferase and GFP (Zhao et al., 2012) have been described. pCSdnd:chDD was made by replacing the 3' UTR of the medaka *nanos3* gene (NS) with the medaka *dnd* 3' UTR (DD). pCSdndΔ1:chDD was constructed by replacing the full-length *dnd* open reading frame (ORF) with a partial *dnd* ORF for the 127 N-terminal amino acids in pCSdnd:chDD. In pCSdnd:chNS, pCSdnd:chDD, and pCSdndΔ1:chDD, four mismatch nucleotides were introduced to the target sequence of MOdnd around the ATG codon without changing the amino acid sequence (Figure S1A). Consequently, the synthetic mRNAs from these constructs were targeted by MOddm but not MOdnd. Plasmid pH2Bgf contains the fusion between the cDNA encoding the bovine histone H2B and EGFP. The fusion gene was PCR-cloned using primers (CGG AGA TCT ATG CCA GAG CCA GCG A and CCG CTC GAG TTA CTT GTA CAG CTC GTC CAT GC; the restriction sites are underlined) from pBOS-H2Bgf (BD Biosciences) and inserted between BamHI and XhoI to replace the cherry cDNA in pCScherry. The 3' UTRs of the medaka *dnd*, *dazl*, and *nanos3* were used to localize synthetic RNAs in germ cells. Structures of these plasmids are illustrated in Figure S1B.

RNA Synthesis and In Situ Hybridization

Sense and antisense *dnd* and *vasa* riboprobes were synthesized from linearized plasmids pGEMdnd (Liu et al., 2009) and pGEMvas (Li et al., 2009). Capped *dnd* and *vasa* mRNA were synthesized by using the mMessage Machine kit (Ambion) from linearized pCSdnd:chDD (Liu et al., 2009) and pCSvas:gfp and stored at -80°C until use. Chromogenic and two-color FISH and nuclear staining with DAPI were performed as described (Liu et al., 2009; Xu et al., 2009).

Microinjection

Medaka embryos were injected at the one-cell stage (Hong et al., 2004b; Li et al., 2009). MOddm and MOdnd (10–5,000 ng/ μl), synthetic mRNAs (10–100 ng/ μl), and plasmid DNA (50 ng/ μl) were injected alone or in combination.

Lineage Tracing in Single-Cell Culture

Embryo preparation, cell isolation, and culture in gelatin-coated plastic ware were done essentially as described (Hong et al., 1996, 1998a, 1998b, 2010; Yi et al., 2009). For single-embryo culture, disperse cells were collected from a single embryo and seeded into a single well of a 96-well plate containing 150 μl of ES cell culture medium ESM2 (Hong et al., 1996), one embryo per well. For single-cell culture, blastomeres were dissociated from embryos at the 32- and 64-cell stages and seeded by pipetting into 96-well plates, one cell per well. PGCs were monitored by GFP expression from 12 hr post culture onward.

Cell Death Detection

Cell death in developing embryos were detected by vital dye acridine orange (AO, Sigma) as described in (Mei et al., 2008). Briefly, embryos were dechorionated by proteinase K (Sigma) and hatching enzyme treatment as described (Yi et al., 2009), then embryos

were incubated in 5 $\mu\text{g}/\text{ml}$ AO for 15–20 min in BSS-1%PEG. After three times washes for 5 min each in BSS-1%PEG, the embryos were then observed under a fluorescent stereo microscopy.

Microscopy

Standard LM for observation and photography was done on a Leica MZFIII stereo microscope, and Zeiss Axiovert invert and Axiovert upright microscopes with a Zeiss AxioCam M5Rc digital camera (Zeiss Corp) as described previously (Hong et al., 2010; Xu et al., 2009; Yi et al., 2009). Preparation of ultrathin cryosections from blastomeres of medaka blastula embryos and correlative light EM were done as described (Yuan et al., 2014).

Statistics

Statistical analyses were calculated by using GraphPad Prism v4.0. Consolidated data were presented as means \pm SD and p values were calculated using the non-parametric Student t test.

SUPPLEMENTAL INFORMATION

Supplemental Information includes five figures and one table and can be found with this article online at <http://dx.doi.org/10.1016/j.stemcr.2016.01.002>.

AUTHOR CONTRIBUTIONS

N.H., H.Z., J.S., and Y.H. designed the research. N.H., M.L., Y.Y., T.W., M.Y., and H.X. performed the experiments. N.H., J.S., and Y.H. wrote the paper.

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