Aven is dynamically regulated during *Xenopus* oocyte maturation and is required for oocyte survival

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We have analyzed the expression and function of the cell death and cell cycle regulator Aven in *Xenopus*. Analysis of *Xenopus* Aven expression in oocytes and embryos revealed a band close to the predicted molecular weight of the protein (36 kDa) in addition to two bands of higher molecular weight (46 and 49 kDa), one of which was determined to be due to phosphorylation of the protein. The protein is primarily detected in the cytoplasm of oocytes and is tightly regulated during meiotic and mitotic cell cycles. Progesterone stimulation of oocytes resulted in a rapid loss of Aven expression with the protein levels recovering before germinal vesicle breakdown (GVBD). This loss of Aven is required for the G2–M1 cell cycle transition. Aven morpholino knockdown experiments revealed that early depletion of the protein increases progesterone sensitivity and facilitates GVBD, but prolonged depletion of Aven results in caspase-3 activation and oocyte death by apoptosis. Phosphorylated Aven (46 kDa) was found to bind Bcl-xL in oocytes, but this interaction was lost in apoptotic oocytes. Thus, Aven alters progesterone sensitivity in oocytes and is critical for oocyte survival.

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In order for oocytes to undergo fertilization, meiotic maturation is required, that is the progression from a G2 phase-arrested oocyte to an M phase-arrested oocyte. In *Xenopus laevis* oocytes, progesterone is used to stimulate resumption of the meiotic cell cycle, whereupon adenylyl cyclase activity is inhibited, cAMP levels are decreased and protein kinase A (PKA) activity is suppressed.¹⁻³ This decrease in PKA activity is necessary for synthesis of MOS protein and initiation of the MOS-MEK-MAPK signaling cascade and subsequent germinal vesicle breakdown (GVBD).⁴⁻⁵

Aven is an apoptotic regulator, inhibiting mitochondrial apoptosis by binding to and inhibiting the self-association of pro-apoptotic Apaf-1 and binding to and enhancing anti-apoptotic Bcl-xL activity.⁶⁻¹⁰ Overexpression of human Aven mRNA in *Xenopus* oocytes delays oocyte maturation, whereas overexpression of human and *Drosophila* Aven in *Xenopus* egg extract causes mitotic cell cycle arrest.¹¹⁻¹² Following DNA damage, it was shown that Aven activates ataxia telangiectasia-mutated (ATM) kinase to inhibit G2/M cell cycle progression.¹¹

There is evidence for steroidal regulation of Aven. It has recently been shown that estrogen upregulates Aven expression at both the mRNA and protein level, in rat seminiferous tubules cultured *ex vivo*.¹³ Also, neonatal rat ovaries cultured in the presence of progesterone show decreased Aven gene expression in microarray analysis of the ovarian transcriptome.¹⁴

Aven has been linked to several diseases including cancer, Prader-Willi syndrome and amyotrophic lateral sclerosis. Aven gene expression is associated with poor prognosis in several cancers, including childhood acute lymphoblastic leukemia, acute myeloid leukemia¹⁵ and breast cancer.⁹ Microarray analysis has also identified Aven as being overexpressed in an ovarian carcinoma cell line resistant to the chemotherapeutic agent vincristine,¹⁶ and underexpressed in colon cancer cell lines resistant to methotrexate.¹⁷

Here we present an analysis of endogenous Aven expression during oogenesis, oocyte maturation and early embryonic development in *Xenopus laevis*. We show that progesterone-stimulated maturation leads to an initial loss of Aven protein that is required for progression to GVBD with protein levels recovering thereafter. In addition, morpholino knockdown revealed that loss of Aven protein facilitated a rapid progression to GVBD but ultimately resulted in oocyte death by apoptosis.

**Results**

**Aven expression and post-translational modification in oocytes and embryos.** The predicted molecular weight of *Xenopus* Aven is 34.6 kDa and an analysis of Aven expression in oocytes and embryos revealed a band close to the predicted molecular weight of the protein (36 kDa) in...
addition to two bands of higher molecular weight (46 and 49 kDa) (Figure 1a). The 36 and 46 kDa bands were detected throughout oogenesis and in pre- and post-mid blastula transition (MBT) embryos. The highest molecular weight 49-kDa band was only detected in previtellogenic stage II and III oocytes. Comparing nuclear and cytoplasmic fractions from stage VI oocytes, we demonstrated that Aven is entirely cytoplasmic (Figure 1b). This is in agreement with previous reports that Aven primarily localizes to the cytosol, whereas a small fraction is reported to be nuclear. Studies have shown that the intracellular localization of Aven is highly regulated and Aven contains a highly conserved leucine-rich nuclear export sequence (LR-NES).

The intensity of the 36- and 46-kDa band varied, with levels decreasing as oogenesis proceeded. In pre-MBT embryos, the 46-kDa band in particular fluctuated, this may reflect stage-dependent changes in the protein or cell cycle-dependent changes as embryos synchronously divide (Figure 1c). Post MBT in gastrulation stage embryos, the 46-kDa band was stabilized and predominated.

To verify that the antibody was in fact detecting Aven protein, we used an Aven morpholino oligonucleotide to inhibit Aven protein synthesis. Stage VI oocytes were microinjected 1 h before progesterone-stimulated in vitro maturation. Western blot analysis of GVBD samples showed the knockdown of both the 36-kDa and the 46-kDa bands, indicating that both bands represent the Aven protein (Figure 1d).

The Xenopus AVEN protein sequence has several predicted phosphorylation sites by multiple kinases (Supplementary Figure S1). Treatment of stage II and stage VI oocytes with potato acid phosphatase caused the depletion of the 46-kDa band after a 2-h incubation (Figure 1e). This indicates that 46-kDa band represents a phosphorylated form of the AVEN protein.

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Figure 1 AVEN expression and subcellular localization in oocytes. (a) Representative western blot analysis of Aven expression in stage II–stage VI oocyte extracts. GAPDH was used as a loading control (n = 5). (b) Western blot analysis of stage VI whole oocyte extract, cytoplasmic extract and nuclear extract (n = 3). (c) Aven expression in embryos from single cell to stage 11.5 (n = 3). (d) Aven morpholino results in a loss of Aven expression at GVBD (n = 6). (e) Phosphatase treatment of Stage II and Stage VI oocytes caused depletion of the 46-kDa Aven band, (n = 3)
the expression of both bands peaking at MI and MII, as determined by phospho MAPK levels. Calcium ionophore (A23187) induced metaphase II release, confirmed by pHistone-H3 depletion, resulted in a decrease in Aven expression (Figure 2b). Next, we examined Aven levels throughout the mitotic cell cycle using cycling extracts where oscillations were assessed by following pHistone-H3 expression. Aven protein levels, particularly the 46-kDa band, appear to peak just before the M phases of the cell cycle (Figure 2c).

**Oocytes injected with the Aven morpholino had increased sensitivity to progesterone, with 10-fold less progesterone needed to induce 50% GVBD by 5 h compared with control oocytes (Figure 3e). Strikingly, at a very low concentration of progesterone (5 μg/ml), 37% of oocytes injected with the morpholino showed GVBD compared with only 8% of the control oocytes.**

Aven knockdown led to oocyte death following the onset of GVBD, with 50% of oocytes dying by 6 h post progesterone addition (Figure 3d). This oocyte death was characterized by the classic apoptotic phenotype previously described.  

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**Figure 2** Dynamic changes in Aven expression during oocyte maturation. (a) Western blot analysis of Aven in oocyte extracts before and at the indicated time points (hours) after progesterone (P4) addition. In all, 64% of oocytes reached GVBD by 3.5 h as determined by the appearance of a white spot (92% at 4 h) and GVBD and MII were confirmed by pMAPK and pHistone-H3 expression. Aven expression is rapidly lost on progesterone addition and recovers by GVBD, error bars correspond to S.D., based on four experiments. (b) Loss of Aven expression following calcium ionophore A23187 induced metaphase release, error bars correspond to S.D., (n = 4). (c) Fluctuations in Aven expression in cycling extracts with pHistone-H3 peaks marking the M phase of the cell cycle.
where the oocytes displaced pigment granules to form a large white spot at the animal pole of the oocyte (Figure 3f).

**Aven is required for oocyte survival**

**Aven promotes oocyte survival by interacting with Bcl-xL and preventing apoptosis.** To assess whether the oocyte death induced by Aven protein depletion is due to apoptosis, morpholino-injected oocytes were examined by western blot for the presence of active caspase-3 (Figure 4a). Aven morpholino-injected oocytes showed activated caspase-3 by 5 h after progesterone addition,
whereas no caspase-3 activation was detected in uninjected or control morpholino-injected oocytes. This supports activation of an apoptotic pathway in Aven-depleted oocytes, as indicated by the visual morphology (Figure 4b).

The interaction between Aven and Bcl-xL is important in promoting cell survival, therefore, we compared Aven–Bcl-xL interaction in normal and DNA damage-induced apoptotic oocytes. Progesterone-stimulated and γ-irradiated oocytes underwent apoptosis at a time equivalent to MII as indicated by caspase-3 activation (Figure 4c, input MII*). Despite the decreased Aven expression in apoptotic oocytes, it is possible to concentrate the remaining 46-kDa form of the protein by immunoprecipitation and Bcl-xL co-immunoprecipitates with this (Figure 4c, Aven IP). Similar levels of Bcl-xL from both normal and apoptotic oocyte samples were immunoprecipitated, however significantly less Aven protein was associated with Bcl-xL in the apoptotic oocytes (Figure 4c, Bcl-xL IP). This loss of Aven interaction with Bcl-xL in apoptotic oocytes points to the significance of this interaction at GVBD and MII for oocyte survival.

![Figure 4](image-url)

**Figure 4** Cell death activation in Aven-depleted oocytes. (a) Western blot analysis for active caspase-3 expression in uninjected, control MO-injected and Aven MO-injected, progesterone-treated oocytes (100–120 oocytes per experiment). Active caspase-3 was only detected in Aven-depleted oocytes 5 h after progesterone treatment. Shown is a representative experiment from n = 7. (b) Representative images (light and fluorescence) of Aven MO-injected oocytes undergoing precocious GVBD and subsequent apoptosis at the indicated times after progesterone addition. (c) A comparison of Aven–Bcl-xL interaction in control and apoptotic oocytes. Aven and Bcl-xL immunoprecipitates were analyzed by western blot at GV, GVBD, MII and MII*; an MII equivalent time point in irradiated oocytes (n = 3). Normal rabbit IgG antibody was used in control immunoprecipitations. Bcl-xL is bound to Aven in immunoprecipitated MII-arrested egg extract, whereas in DNA damage-induced apoptotic extracts, little Aven is associated with Bcl-xL.
Discussion

To date, the only reported endogenous expression of Aven indicates a protein with a higher molecular weight than predicted; analysis of human Aven expression in HeLa cells identified a protein with an apparent molecular weight of 55 kDa, even though the predicted molecular weight of the human protein is 38.6 kDa. A recombinant HA-tagged Drosophila Aven yielded a protein of ~36 kDa, excluding the HA tag. The detection of a 36-kDa band in Xenopus oocytes and embryos is the first report of endogenous Aven close to its predicted molecular weight. The bands of higher apparent molecular weight detected in Xenopus may represent a similar form to that reported in humans and in all likelihood these represent post-translationally modified forms of Aven. Recombinant Aven expression in a Chinese hamster ovary (CHO) cell line resulted in four closely separated bands being detected by western blot, providing further evidence that Aven is post-translationally modified.

Human Aven is a known substrate for the ATM kinase at two SQ sites and evidence of additional basal phosphorylation at Ser and Thr sites also exists. Treatment of stage II and stage VI oocytes with potato acid phosphatase caused the depletion of the 46-kDa band after a 2-h incubation (Figure 1e), indicating that this band represents a phosphorylated form of the Aven protein. The 10-kDa molecular weight shift suggests hyperphosphorylation of the protein at multiple sites. Analysis of the protein sequence reveals several different kinase phosphomotifs, including potential CK2, PKA and an ATM site, (Supplementary Figure S1a).

The phosphomotifs identified are putative substrates for kinases that are known regulators of the cell cycle and/or oocyte maturation. PKA, in particular, is a negative regulator of the G2/M1 transition, and is rapidly depleted in response to progesterone stimulation. It has long been postulated that this decrease in PKA activity results in decreased phosphorylation of a presumptive maturation-inhibiting phosphoprotein. Aven would be a candidate for such a protein if phosphorylation by PKA stabilized the protein preventing GVBD.

The 49-kDa band in stage II extracts was resistant to phosphatase treatment, suggesting that either different post-translational modifications are altering the proteins apparent molecular weight or there are additional isoforms of the protein present. Studies on human and mouse Aven have indeed considered the possibility of different isoforms being present. Northern blot analysis detected a single 1.7-kb Aven transcript in all human adult tissues tested, with high expression in heart, skeletal muscle, kidney, liver, pancreas and testis and a number of cell lines. Analysis of the mouse Aven gene revealed two isoforms, a short (m4Aven-S) and a longer (m4Aven-L) transcript (corresponding to the Aven gene found in other species) encoding proteins of 216 and 342 amino acids, respectively. Thus, there is no evidence from other species studied that the 49–55 kDa protein identified represents a different isoform of the protein. The Aven sequence shows putative O and N glycosylation sites, and it will be interesting to investigate whether such post-translational modifications are responsible for the highest molecular weight forms of the protein.

The expression of Aven as two bands of 36 and 46 kDa is evident at different stages during meiotic maturation. Downregulation of the protein is evident at two points; immediately following progesterone stimulation and a less significant reduction in protein levels between MI and MII. It is of note that there is no evidence of a classical destruction box in the Aven sequence. However, there are several different mechanisms by which proteins might be rapidly degraded, including ubiquitin-mediated proteasomal degradation and proteolytic cleavage. Our preliminary analysis using a proteasome inhibitor suggests proteasomal degradation does not have a role in Aven degradation. The CK2-regulated caspase 8/9 site identified in human Aven is not conserved in Xenopus, ruling out cleavage by these caspasas. Aven cleavage by Cathepsin D is important for the anti-apoptotic potential of the protein and the Cathepsin D cleavage sites are conserved across species, including Xenopus. Using an antibody directed against the C-terminal of the Aven protein, we have been able to detect a band with an apparent molecular weight of 28 kDa, this could represent the predicted Cathepsin D Aven fragment (Supplementary Figure S1b). This fragment is not degraded on progesterone treatment and any generation of a fragment by Cathepsin D activity is more likely related to oocyte survival rather than the required loss of the higher molecular weight forms for G2–MI progression following progesterone treatment.

During meiosis, Aven is consistently associated with BCL-XL and specifically the 46-kDa phosphorylated form of the protein. This suggests that the BCL-XL interaction is dependent on particular post-translational modification(s) of the protein and only a subpopulation of the AVEN interacts with BCL-XL.

In silico analysis of the Aven promoter region, using Matbase from Genomaxit (http://www.genomaxit.de), enabled the identification of a progesterone receptor-binding site and potential CDE/CHR cell-dependent element, which is known to regulate cell cycle-dependent transcription (Supplementary Figure S1c). The role of progesterone in regulating Aven warrants further study and while cell cycle regulation may not be relevant during oocyte maturation, it may have a role in the mitotic cell cycle regulation of Aven observed within the embryo.

Modulation of Aven expression initially impacts on G2–MI progression and ultimately has consequences for oocyte survival. The mechanism by which the progesterone induced cytoplasmic signaling pathway that leads to oocyte maturation impacts on Aven expression remains to be elucidated. It is known that Aven can prevent M-phase entry following DNA damage through its ability to activate ATM and overexpression of the protein inhibits G2/M in oocytes. Our analysis of endogenous Aven expression shows how its loss is required for this cell cycle transition to occur, suggesting that Aven has the capacity to interfere with cell cycle progression independent of a DNA damage signal. Recovery of Aven protein levels is required for oocyte viability, highlighting Aven’s importance as an apoptotic inhibitor in the maturing oocyte. Aven clearly has a dual role in regulating both cell cycle progression and oocyte survival and whether these activities are integrated or are underpinned by completely separate biochemical activities requires further analysis.
Materials and Methods

**Xenopus oocyte isolation and embryo culture.** All animal experiments were conducted under license from the Department of Health and Children (B100/2003) and according to protocols approved by the University College Dublin Animal Research Ethics Committee (AREC-P-10-68).

Stage II–VI oocytes for oogenesis experiments were obtained via collagenase defolliculation (2 mg/ml, Sigma, St. Louis, MO, USA). Stage VI oocytes were obtained by manual defolliculation, matured with progesterone (Sigma, P7556) at the indicated concentrations, observed for GVBD and nuclear extracts were made as described. For phosphatase treatment, St II and St VI oocyte extracts were incubated with potato acid phosphatase at 0.5 μg/ml extract. Calcium ionophore A23187 (Sigma, C7522) was added to OR2 culture media at a concentration of 2 μM to activate metaphase II-arrested oocytes. In vitro fertilization was carried out as described. Embryos were staged, collected and snap-frozen in liquid nitrogen at the indicated stages. Cycling extracts were prepared from unfertilized eggs activated using calcium ionophore A23187 (Sigma, C7522), as described in Murray. Samples were collected at 10-min intervals and snap-frozen in liquid nitrogen.

**γ-Irradiation.** Irradiation of oocytes and embryos was performed using a Gamma Cell Cs137 source. Irradiated samples (33 Gy) were subsequently treated with progesterone to induce maturation.

**Aven knockdown.** Fluorescein-tagged Aven morpholino (5'-GGCCC CGCCTCAATGTTGCTTACCT-3') and control morpholino (5'-CCTCTACTT CAGTTACAATTTTATA-3') were obtained from Gene Tools, LLC (Philomath, OR, USA) (http://www.gene-tools.com). The morpholinos were microinjected into stage VI oocytes, at a final concentration of 5 μM, 1 h before progesterone treatment. To confirm microinjection, live oocytes were observed for fluorescence using a Carl Zeiss Ltd. (Cambridge, UK) stereo Lumar V12 with FITC filter.

**Immunoblotting, immunoprecipitation and antibodies.** Protein samples were separated according to their apparent molecular mass on NuPAGE 4–12% Bis-Tris Gels (Invitrogen, Carlsbad, CA, USA), and transferred to a PVDF membrane. After blocking in 5% bovine serum albumin (BSA) in PBS–0.1% Tween-20, membranes were incubated with the primary antibody, followed by incubation with horseradish peroxidase-conjugated secondary antibodies. Proteins were visualized using Enhanced Chemiluminescence Western Blotting substrate (Thermo Scientific, Rockford, IL, USA; #23106). Densitometry was performed using the ImageJ program (http://rsweb.nih.gov/ij/).

Endogenous Aven was immunoprecipitated by incubating 40 μl of oocyte extract with 20 μl of Aven antibody or control normal rabbit IgG antibody overnight at 4 °C, then incubating with protein A/G agarose beads (Thermo Scientific, 20421) for 2 h at room temperature. Washes were performed using IP buffer (25 mM Tris, 150 mM NaCl; pH 7.2).

Anti-AVEN antibodies (Cell Signaling Technology Inc., Danvers, MA, USA; #2300; Abcam, Cambridge, UK, ab77014), anti-phospho-histone H3 (Ser10) antibody (Merck Millipore, Bitterica, MA, USA, cat. #06-570), anti-GAPDH antibody (Cell Signaling, #3200), anti-p-MAPK (Cell Signaling, #9101), anti-Caspase-3 (BD Biosciences, Oxford, UK, #559565), anti-Bcl-x(L) (Santa Cruz Biotechnology, Santa Cruz, CA, USA, sc-77014), normal Rabbit IgG (Cell Signaling, #2729).

Two antibodies raised against an N- and C-terminal epitope in human Aven were compared for their ability to detect Xenopus Aven (Supplementary Figure S1b). The N-terminal antibody detected bands of 36 and 46 kDa, both of which were knocked down following injection of Aven morpholino to oocytes (Figure 1d). This antibody was used throughout the study. The C-terminal antibody also detected the 46-kDa band and in addition a 28-kDa band, which may be the previously reported cathepsin D-cleaved C-terminal proteolytic fragment.

**Statistical analysis.** Statistical analyses were carried out using the GraphPad Prism version 6.0 software package for Windows. Differences between groups were considered significant when P-values were less than 0.05.

**Conflict of Interest.** The authors declare no conflict of interest.

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