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Zinc sparks are triggered by fertilization and facilitate cell cycle resumption in mammalian eggs

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Abstract

In last few hours of maturation, the mouse oocyte takes up over twenty billion zinc atoms and arrests after the first meiotic division, until fertilization or pharmacological intervention stimulates cell cycle progression towards a new embryo. Using chemical and physical probes, we show that fertilization of the mature, zinc-enriched egg triggers the ejection of zinc into the extracellular milieu in a series of coordinated events termed zinc sparks. These events immediately follow the well-established series of calcium oscillations within the activated egg and are evolutionarily conserved in several mammalian species, including rodents and non-human primates. Functionally, the zinc sparks mediate a decrease in intracellular zinc content that is necessary for continued cell cycle progression, as increasing zinc levels within the activated egg results in the reestablishment of cell cycle arrest at metaphase. The mammalian egg thus uses a zinc-dependent switch mechanism to toggle between metaphase arrest and resumption of the meiotic cell cycle at the initiation of embryonic development.

Introduction

The biological functions of metal ions have traditionally been thought to be limited to structural and catalytic roles within proteins. The alkaline earth metal calcium is one metal in biology for which there are well-established signaling functions. Periodic elevations in the intracellular concentration of free calcium ions, also known as calcium transients or oscillations, are readily detected using fluorescent probes and are known to drive a number of biological processes (reviewed in ref. 1). This is perhaps best illustrated in the egg, where

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repetitive calcium transients are among the earliest observable events after fertilization². Several parameters of these calcium transients, including total number, frequency, and amplitude influence which downstream developmental events are initiated³⁻⁵. These cellular processes, which include cortical granule (CG) exocytosis^{6, 7} and cell cycle progression^{8, 9}, can be initiated in the absence of sperm by stimulatory agents that induce parthenogenesis¹⁰⁻¹³. Interestingly, the normal pattern of calcium oscillations is disrupted in eggs matured under conditions which limit the availability of the transition metal zinc¹⁴, suggesting that the physiologies of these metals are somehow connected in the egg.

In a departure from its well-established role as an enzymatic cofactor or structure-stabilizing agent, fluctuations in the total concentration of intracellular zinc have recently been shown to contribute to the proper cell cycle regulation in maturing oocytes. Intracellular zinc levels increase by more than fifty percent and over 10^{10} atoms per cell are accrued during the final stage of oocyte maturation, also known as meiotic maturation¹⁴. This significant cellular metal accumulation event occurs over a remarkably short time interval and is a physiological imperative, as insufficient accumulation of zinc leads to a premature meiotic arrest at telophase I instead of metaphase II¹⁴. This zinc-dependent meiotic checkpoint arises, in part, because zinc-insufficient eggs fail to reestablish maturation promoting factor (MPF) activity¹⁵, which is necessary for eggs to set up and maintain meiotic arrest at metaphase II. These results revealed a previously unrecognized role for zinc in cell cycle switching events: namely, the regulation of meiotic progression in the egg.

Given this role for zinc in the maturing oocyte, we explored whether this function extended to cell cycle regulation in the fertilized egg. We first investigated the dynamics of zinc and calcium within the physiological context of fertilization using a variety of chemical probes that can report changes in the concentration of labile and loosely bound zinc ions and discovered that the calcium oscillations induced a rapid ejection of zinc through an event termed the zinc spark. We show that zinc sparks are evolutionarily conserved in three mammalian species, and that these fluxes in intracellular zinc availability mediated cell cycle resumption. Together with our previous reports on zinc in the maturing oocyte, we describe herein a new zinc-dependent mechanism for cell cycle regulation in the mammalian egg.

Results and Discussion

Release of zinc at egg activation is a conserved event in mammalian species

Extracellular zinc events were monitored during fertilization of mouse eggs using a membrane impermeant derivative of the zinc fluorophore FluoZin-3¹⁶. Strikingly, fertilization triggered the repetitive release of zinc into the extracellular milieu (Fig. 1a-b). These events were termed zinc sparks for their brevity and intensity. Successful fertilization was confirmed by extrusion of the second polar body (Fig. 1c-d, PB2). These events were recapitulated during parthenogenesis of mouse eggs (Fig. 1e, Video S1). Importantly, zinc sparks are an evolutionarily conserved phenomenon, as they were also observed during parthenogenetic activation of two different non-human primate species, *Macaca mulatta* (rhesus macaque) and *Macaca fascicularis* (crab-eating macaque) (Fig. S1, Video S2). Additional characterization of the zinc sparks was completed using parthenogenetically activated mouse eggs.

The dynamics of the zinc sparks were variable from egg to egg. Zinc sparks occurred during the first 90 min of activation, and individual eggs exhibited between one and five zinc sparks, which were counted in a plot of fluorescence intensity over time (see representative plot in Fig. 1a). Eighty-one percent of all eggs examined (N = 28) exhibited two or three exocytosis events, while eleven percent exhibited one event and eight percent exhibited four

or five events. No more than five zinc sparks were observed in any of the samples. The interval between each exocytic event was 9.5 ± 0.8 min (mean \pm SEM, N=36), although the range of intervals was quite broad, between 3.46 and 26.00 min.

It is notable that the zinc sparks could be induced by fertilization or by parthenogenesis using two different reagents: strontium chloride in rodents and ionomycin in two non-human primate species. While strontium chloride induces multiple calcium transients⁶ whereas ionomycin only induces a single calcium transient¹⁷. Accordingly, multiple zinc sparks were observed in rodents whereas only a single zinc spark was seen in the non-human primate species. This result speaks to a possible calcium dependence of the zinc sparks, and this was further investigated in the following experiments.

Initiation of the zinc sparks is dependent on intracellular calcium transients

To understand the role of zinc sparks within the temporal context of the calcium oscillations, we simultaneously monitored intracellular calcium oscillations alongside zinc sparks using the calcium fluorophore Calcium Green-1 AM (ex. 506, em. 531), which has similar excitation and emission spectra to FluoZin-3 (Fig. 2a, Video S3). Each zinc spark was closely associated with an intracellular calcium transient (Fig. 2b); upon closer inspection, it was found that each zinc spark was immediately preceded by an elevation in intracellular calcium (Fig. 2c). Parallel experiments employing a membrane permeable zinc fluorophore, FluoZin-3 AM, did not reveal clear intracellular zinc transients (Fig. S2a), ruling out the possibility that large oscillations in intracellular free zinc contributes to the intracellular transients detected by Calcium Green-1 AM. We confirmed that this was representative of successfully activated eggs by simultaneously monitoring second polar body (PB2) extrusion by brightfield imaging (Fig. S2a). Further control experiments employing one fluorophore at a time reveal that the zinc sparks are not dependent upon or affected by the presence of the calcium probe; for example, detection of zinc sparks using FluoZin-3 in the absence of Calcium Green-1 AM is shown in Fig. 1a (fertilization) and Fig. 1e (parthenogenesis).

The observation that calcium transients immediately preceded each zinc release event supports the idea that the zinc sparks were directly dependent on intracellular calcium fluxes. Two lines of evidence provided additional support. First, zinc sparks did not occur in the absence of an activating agent and hence, a lack of calcium oscillations (Fig. S2b). Second, zinc sparks were absent in eggs treated with the membrane permeable acetoxymethyl ester (AM) derivative of 1,2-bis(o-aminophenoxy)ethane-N,N,N',N'-tetraacetic acid (BAPTA). This reagent is commonly used to suppress calcium oscillations in eggs^{6, 7} and as such, both calcium transients and zinc sparks were absent in BAPTA AMtreated eggs even after extended culture in strontium chloride (Fig. S2c). Furthermore, these eggs did not extrude a second polar body or form a pronucleus (Fig. S2d-e). Taken together, these data indicate that the zinc sparks are preceded by and may depend upon the calcium oscillations that begin early in the egg activation process.

Localization of zinc is cortically polarized in mature eggs

We noted that the zinc sparks were excluded from the region corresponding to the location of the meiotic spindle, as confirmed by the extrusion of the second polar body at that site (Fig. 2a, arrowheads). To investigate possible cellular sources of the zinc release, the distribution of zinc was determined in unfertilized eggs by synchrotron-based x-ray fluorescence microscopy (XFM). This technique provides the localization and abundance of the total zinc content of a sample, including both the loosely bound (labile or chelatable) and tightly bound cellular pools, and was previously used to quantify zinc and other transition metals in maturing oocytes ¹⁴. The XFM images revealed distinct regions of high zinc

concentration that formed a polarized and hemispherical pattern (Fig. 3a; i-iii represent three independent samples). Other essential transition metals such as iron exhibit a homogeneous distribution across the entire cell (Fig. 3b; i-iii represent three independent samples). Quantitative analysis revealed that zinc was an order of magnitude more abundant than either iron or copper at all points between the mature, *in vivo* ovulated (IVO) MII stage egg and the two-cell embryo (24 hours post-fertilization, or hpf). There is a downward trend with a mean loss of about six billion atoms of zinc upon fertilization, and this trend continued at 6 hpf and 24 hpf (Fig. S3). Assuming different standard deviations for each elemental group, ANOVA analysis noted a significant difference in the mean zinc content across the four timepoints (p = 0.019 for zinc). We also confirmed by fluorescence imaging of intracellular zinc that chelatable zinc, as detected by FluoZin-3 AM, also decreased in activated eggs following the zinc sparks (Fig. S4).

We next asked whether the polarized pattern for total zinc correlated with the distribution of labile (or chelator-accessible) zinc in the egg. Two chemically distinct, zinc-selective fluorophores were used to visualize this loosely bound zinc in live, unfertilized eggs. Projected images of confocal Z-stack series revealed that zinc fluorescence was concentrated away from the meiotic spindle, which was marked with fluorescent DNA probes (Fig. 3c-h). This was also evident when analyzing the Z-series slice by slice, which clearly illustrated higher cortical localization of zinc relative to the cytoplasm (Fig. S5). Furthermore, zinc was present in distinct intracellular compartments as indicated by punctate foci of fluorescence. This pattern was independent of the molecular properties of the zinc probe: both zinquin ethyl ester (Fig. 3c) and FluoZin-3 AM (Fig. 3f) yielded the same general pattern wherein a majority of the foci have a peripheral localization. These probes have pM values of 9.3¹⁸ and 8.8^{16} , respectively, where pM = $-\log[M]$ when [L] = $10 \mu M$ and [M] = $1 \mu M$ at pH 7.4¹⁹. Given that the pM value of zinc-binding metalloproteins are significantly higher (for example, carbonic anhydrase has a pM value of 12.4¹⁸), it is unlikely that the probes are detecting zinc tightly bound to metalloproteins. Rather, these and related fluorescent chelators compete by mass action for zinc ions that are otherwise weakly bound to biopolymers and low molecular weight compounds. Some of the latter have been proposed to buffer intracellular zinc concentrations in various compartments^{20, 21}. While the zinc pools in different compartments of eukaryotic cells are not well understood in either kinetic and thermodynamic terms, the fluorescent probes are generally thought to be delineating compartments with elevated concentrations of "chelator accessible" zinc ions^{22, 23}. In addition to their punctate distribution, the pools of chelator-accessible zinc were concentrated towards the vegetal pole and away from the meiotic spindle, whose location was marked with live-cell nuclear stains Syto 64 (Fig. 3d) or Hoechst 33342 (Fig. 3g) as seen in the merged images (Fig. 3e and 3h). The polarized distribution detected both by the zinc fluorophores and by XFM corroborates a significant cortical compartmentalization of zinc in the egg. Since the polarized distribution of labile zinc mirrors the polarized release of zinc during the zinc sparks, it is likely that some fraction of these zinc-enriched compartments act the source of the coordinated zinc release events, i.e., the zinc sparks.

The polarization of the zinc-enriched compartments mirrors that of the cortical granules (CG), which are exocytic vesicles that participate in the cortical reaction, which involves the hardening of the zona pellucida to establish a block to polyspermy at fertilization²⁴. We hypothesized that the zinc sparks would mirror the mechanisms regulating the cortical reaction and first investigated whether the zinc sparks were the product of an exocytic event. To inhibit exocytosis, eggs were treated with cytochalasin B (CytoB), an agent that disrupts actin reorganization and has previously been shown to disrupt CG exocytosis in mouse and hamster eggs^{7, 25}. CytoB-treated eggs still undergo normal calcium oscillations upon activation but only exhibit a single zinc spark concomitantly with the first calcium transient (Fig. S6). In most cases, subsequent sparks were inhibited (Fig. S6). It is likely that a cohort

of the zinc-containing vesicles is pre-fused with the egg membrane, allowing the first spark to occur even in the presence of CytoB. This type of phenomenon has been described for cortical granules in the sea urchin egg, where one subset of CGs are in a hemifused state prior to activation²⁶.

To determine whether the zinc sparks participate in zona hardening like the cortical reaction, we utilized a zinc-insufficient egg model that was obtained by treating oocytes with the heavy metal chelator TPEN during *in vitro* maturation¹⁴. These eggs did not undergo zinc sparks, even when activated in the same dish as control eggs (Video S4). We previously reported that these zinc-insufficient eggs fertilize at rates comparable to control eggs. Additionally, zinc-insufficient eggs form the normal number of pronuclei, as indicated by the presence of only one male and one female pronucleus¹⁴. Therefore, we find no evidence that the zinc sparks participate in the block to polyspermy.

Meiotic resumption at egg activation is dependent on the zinc sparks

The zinc-insufficient egg model revealed a critical function for zinc in the proper establishment of cell cycle arrest in the mature egg¹⁴, which was supported by the observation that chelation of zinc initiated egg activation²⁷. Given these results, we considered the possibility that a decrease in bioavailable zinc was necessary to permit downstream developmental events upon egg activation. To address this possibility, we treated activated eggs with zinc pyrithione (ZnPT), a zinc ionophore that elevates the intracellular free zinc content of mammalian cells²⁸. First, we performed a titration on unfertilized eggs and did not observe cytotoxic effects until concentrations of ZnPT reached 50 μM (Fig. S7). To minimize such cytotoxic effects, a five-fold lower concentration of ZnPT (10 μM) was used for subsequent experiments. Eggs were subjected to a 10 min ZnPT treatment after the completion of the zinc sparks at 1.5 hours post-activation (hpa) (Fig. 4a). Control (untreated) and ZnPT-treated eggs were then cultured until pronuclear formation was observed in the control group (6-8 hpa; Fig. 4b, c). Ninety percent of control eggs formed an organized pronucleus that was visible by brightfield (Fig. 4b, arrowheads) and also by fluorescence (Fig. 4d). F-actin was homogeneous around the cell (Fig. 4e) and α tubulin was compact in a spindle remnant (Fig. 4f) between the egg and the second polar body (merged image in Fig. 4g).

In contrast, only seventeen percent of ZnPT-treated eggs formed a pronucleus; the majority did not have a pronuclear structure that was clearly visible by brightfield (Fig. 4c). In eighty-four percent of these eggs lacking a pronucleus, we noted the presence of aligned chromosomes (Fig. 4h), instead of the membrane-enclosed nucleus observed in the control eggs. Notably, we also observed an actin-enriched region in the ZnPT-treated eggs (Fig. 4i, arrow) at the cortical region overlying the chromosomes (Fig. 4j). This "actin cap" is a polarized feature thought to be unique to unfertilized eggs^{29, 30}. Most strikingly, the chromosomes and α -tubulin (Fig. 4j) were organized into a metaphase-like spindle (merged image in Fig. 4k), which resembled a freshly ovulated, MII-arrested egg (Fig. 4l-o). It should be noted that only eggs with a clear second polar body were selected for ZnPT treatment; thus, the spindle configuration in ZnPT-treated activated eggs, while metaphase-like, is not that of metaphase II. An analogous metaphase-like spindle and arrest has been seen in parthenogenetically activated rodent eggs that receive insufficient activating stimulus: such cells complete meiosis II but stall at a so-called third metaphase (MIII) instead of progressing to interphase^{15, 19}.

In the reverse of the zinc-loading experiment, the intracellular availability of zinc in the egg was decreased by continuous culture in the presence of 10 μ M TPEN; we previously confirmed the specificity of this chelator for zinc in mouse oocytes¹⁴. After 8 hrs of culture, most control eggs maintained metaphase II arrest while those exposed to TPEN had a

pronucleus-like structure at significantly higher rates (Fig. S8). A similar observation was made by Suzuki and colleagues with much higher concentrations of TPEN, and they further reported that other heavy metal chelators including p-aminosalicylic acid, ammonium tetrathiomolybdate, and 2,2'-bipyridine, were unable to cause egg activation²⁷, thus speaking to the unique role of zinc in driving this process. These complementary experiments support the idea that a significantly elevated zinc quota is associated with metaphase arrest and that the coordinated exocytosis of zinc upon egg activation relieves this arrest. Furthermore, these results also distinguish the intracellular target of the calcium-selective chelator BAPTA AM (Fig. S2) from that of TPEN. Given that BAPTA is also known to have a substantial affinity for zinc $^{31-33}$ (K_d = 107 nM for Ca $^{2+}$ and 8 nM for Zn $^{2+}$), a potential question is whether the BAPTA AM treatment is influencing zinc- or calcium-dependent intracellular processes. At the 10 µM concentration used in our studies, BAPTA and TPEN had significantly different effects on egg physiology. This is illustrated by the fact that longterm exposure to BAPTA did not parthenogenetically activate eggs (Fig. S2) but TPEN did (Fig. S8). Under these conditions, we conclude that BAPTA is scavenging intracellular free calcium and thereby blocking calcium oscillations. Taken together, the results support a model in which the calcium oscillations initiate and are required for the programmed loss of cellular zinc via the zinc sparks and that in turn drives cell cycle resumption.

Summary

It was previously suggested that only a modulation of intracellular zinc, not calcium, was essential for the resumption of the cell cycle following egg activation²⁷. In contrast, we show here that within the physiological context of fertilization, calcium transients immediately precede each zinc spark event. These apparently paradoxical results can be explained as thus: both studies achieve a decrease in the intracellular bioavailability of zinc but through different means (summarized in Fig. 5). Whereas we reached this endpoint through a physiological means, the calcium-triggered zinc spark, the previous report bypassed the need for a calcium-dependent event by artificially lowering zinc availability (TPEN) within the mature egg²⁷. Thus, we conclude that the calcium transients are involved in the physiological mechanism by which the lowering of intracellular zinc availability is initiated: namely, by inducing the zinc sparks and permitting cell cycle resumption to occur. One way that this change in the intracellular zinc status may regulate the cell cycle is by affecting the activity of the anaphase promoting complex/cyclosome (APC/C) inhibitor Emi2 (early mitotic inhibitor-2), which has been shown to be important for the establishment of metaphase II arrest³⁴ and contains a conserved zinc-binding region that is required for activity²⁰. Although we have reported trends in the egg's total zinc content preand post-fertilization, additional studies are required to define these vital fluxes in greater detail. Methods that quantify changes in the subset of the total zinc pool that is compartmentalized, labile and/or subject to modulation in response to egg activation will provide greater mechanistic insight in conjunction with the pathways that are activated at the protein level.

We postulate that in the egg, intracellular zinc fluxes play a gatekeeping role between cell cycle arrest and resumption, and that this pathway is controlled in part by intracellular calcium signaling (Fig. 5). Ultimately, the fertilized mammalian oocyte faces a physiological imperative to lower intracellular zinc availability in order to permit cell cycle resumption, and this is achieved in part via the calcium-regulated zinc sparks. Indeed, the same effect can be achieved pharmacologically through the intracellular chelation of zinc²⁷. Importantly, the abundance of zinc appears to act as both a permissive factor and a maintenance factor in metaphase arrest. During oocyte maturation, an elevated level of zinc must be accumulated to establish metaphase arrest¹⁴. Conversely, during egg activation, the zinc sparks are used to decrease bioavailable zinc, but metaphase arrest can be reestablished

if the levels of intracellular zinc are raised. These results implicate zinc as a key regulatory element in egg activation and are consistent with a specialized role for this transition metal in the regulation of developmental processes during the earliest stages of life.

Materials and Methods

Mouse egg collection

Eggs were collected from adult (6-8 weeks old) female mice of the CD-1 strain. Mice were superstimulated with an i.p. injection of 5 IU pregnant mare's serum gonadotropin (PMSG, EMD Biosciences, San Diego, CA), followed 46-48 h later by an i.p. injection of 5 IU human chorionic gonadotropin (hCG, Sigma-Aldrich, St. Louis, MO). Mice were sacrificed by CO_2 asphyxiation and cervical dislocation 13-14 h post-hCG administration. Clutches of cumulus-oocyte complexes were isolated from the oviducts. Cumulus cells were denuded using 30 μ g/ml hyaluronidase and gentle aspiration through a narrow-bore pipette. Animals were treated in accord with the National Institutes of Health Guide for the Care and Use of Laboratory Animals. Food and water were given ad libitum. The Northwestern University Institutional Animal Care and Use Committee (IACUC) approved all protocols.

Time-lapse imaging of mouse eggs during in vitro fertilization

Sperm from the cauda epididymides of proven breeder CD-1 males were used for sperm in vitro fertilization (IVF). The sperm population was purified using Percoll gradient centrifugation (PGC) as described previously³⁵ and capacitated for up to 3 h in IVF medium composed of potassium simplex optimized medium (KSOM, Millipore, Billerica, MA) supplemented with 3 mg/ml bovine serum albumin (BSA, MP Biomedicals, Solon, OH) and 5.36 mM D-glucose (Sigma-Aldrich). Superovulated eggs were collected as described above. The zona pellucida (ZP) was removed by brief treatment in acidic Tyrode's solution (pH 2.5, Millipore). The ZP-free eggs were allowed to settle in a 50 µl drop of calcium- and magnesium-free Dulbecco's PBS (DPBS, Invitrogen, Carlsbad, CA) on a glass-bottom dish (Bioptechs Inc., Butler, PA) coated with poly-L-lysine. After 10 min, IVF medium containing 20 uM FluoZin-3 (Invitrogen) was slowly added to a final volume of 1 ml, taking care not to disturb the eggs. The final DMSO concentration was 0.1% (v/v), as higher concentrations were cytotoxic to sperm. Capacitated sperm were added to the eggs to a final concentration of 1.0×10^6 sperm/ml and image acquisition began immediately upon addition of sperm. Images were acquired every 4 sec for a total duration of 3 hours. All images were acquired on a TCS SP5 confocal microscope (Leica Microsystems, Heidelberg, Germany) equipped with a stage top incubator (Tokai Hit, Shizuoka, Japan), 20× objective, and an Ar (488 nm) laser line.

Parthenogenetic activation and pyrithione treatment of mouse eggs

Strontium chloride (SrCl₂) was prepared as a 1 M stock solution in calcium-free KSOM (Millipore) and stored as ready-to-use aliquots at $-20\,^{\circ}\text{C}$. SrCl₂ was diluted to a working concentration of 10 mM in calcium-free KSOM. Eggs were incubated in SrCl₂ for 90 min. In some cases, eggs were then treated with zinc pyrithione (ZnPT, Sigma-Aldrich). Dose response experiments were performed to test cytotoxicity; a distinct cytotoxic effect was seen at concentrations above 20 μ M. Therefore, ZnPT concentrations were reduced to 10 μ M and treatment time was limited to 10 min at 37 °C in an atmosphere of 5% CO₂. Both control and ZnPT-treated eggs were washed and transferred to fresh KSOM for extended culture.

Non-human primate oocyte collection, in vitro maturation, and parthenogenetic activation

Ovarian tissue was surgically dissected from *Macaca mulatta* (rhesus macaque, 3 years old) and *Macaca fascicularis* (crab-eating macaque, 3 years old) females at the Panther Tracks Learning Center (Immokalee, FL). The tissue was cut in half, placed in 15 ml of SAGE oocyte washing medium (Cooper Surgical, Trombull, CT), and shipped overnight at ambient temperature. The tissue was processed within 18 hours of surgery. Large follicles were opened by grazing with an insulin-gauge needle. Oocytes of diameter greater than 100 µm (including the zona pellucida) were collected for *in vitro* maturation (IVM), whether they were enclosed by cumulus cells or denuded. They were placed in an IVM medium composed of SAGE oocyte maturation medium (Cooper Surgical) supplemented with 75 mIU/ µl follicle stimulating hormone (FSH, a gift from Organon, Roseland, NJ), 75 mIU/ µl luteinizing hormone (LH, a gift from Ares Serono, Randolph, MA), 1.5 IU/ µl hCG (Sigma-Aldrich), and 5 ng/ml epidermal growth factor (EGF, BD Biosciences, San Jose, CA). Oocytes were observed up to 48 hours post-maturation and used immediately for imaging upon complete extrusion of a polar body.

Time-lapse imaging of parthenogenetically activated eggs

Eggs were first incubated in 10 μ M Calcium Green-1 AM (Invitrogen) and 0.04% (w/v) Pluronic F-127 (Invitrogen) for either 30 min (mouse) or 60 min (non-human primate) at 37 °C. Mouse eggs were washed in calcium-free KSOM (Millipore) and transferred to a 50 μ l drop containing 10 mM SrCl₂ (Sigma-Aldrich) and 50 μ M FluoZin-3 (Invitrogen) on a glass-bottom dish (Bioptechs Inc.). Non-human primate eggs were transferred to one end of a long and shallow drop (20 μ l) of calcium-free KSOM (Millipore) containing 50 μ M FluoZin-3 (Invitrogen) on a glass-bottom dish (Bioptechs Inc.), covered with oil for embryo culture (Irvine Scientific, Santa Ana, CA). Ionomycin was perfused in at the opposite end to a final concentration of 20 μ M. In both cases, image acquisition began immediately and occurred every 4 s for a total of 1-3 hours (mouse) or 15 min (non-human primate), unless noted otherwise. All images were acquired on a TCS SP5 confocal microscope (Leica Microsystems) equipped with a stage top incubator (Tokai Hit), 20× objective, and an Ar (488 nm) laser line.

Imaging of total and labile zinc in the mouse egg

Fertilized eggs were collected at two and six hours post-fertilization were prepared wholemount for synchrotron-based x-ray fluorescence microscopy (XFM). Cells were transferred with a minimal amount of media to an intact 5 mm × 5 mm silicon nitride window (Silson, Blisworth, U.K.) on a heated stage warmed to 37°C. When most of the media had evaporated without drying out the sample, 1 µl of ammonium acetate solution (100 mM, 4°C) was administered to each sample under a dissection microscope. This facilitated a quick wash and dehydration process, leaving the morphology of the sample intact without causing membrane rupture. XFM was performed at Beamline 2-ID-E at the Advanced Photon Source (Argonne National Laboratory, Argonne, IL). 10 keV x-rays were monochromatized with a single bounce Si(111) monochromator, and focused to a spot size of 0.5×0.6 µm using Fresnel zone plate optics (X-radia, Concord, CA). Raster scans were done in steps of 1 µm. Fluorescence spectra were collected with a 1 sec dwell time using a silicon drift detector (Vortex-EM, SII NanoTechnology, CA). Quantification and image processing was performed with MAPS software. The fluorescence signal was converted to a two-dimensional concentration in µg/cm² by fitting the spectra against the thin-film standards NBS-1832 and NBS-1833 (National Bureau of Standards). It was assumed that no elemental content was lost during sample preparation. The data was compared to the total zinc content in unfertilized eggs and two-cell embryos. The data for the two-cell embryo was already published in a previous report¹⁴. The distribution of labile zinc was interrogated in live, unfertilized mouse eggs using two independent zinc fluorophores, zinquin ethyl ester

(Sigma-Aldrich) and FluoZin-3 AM (Invitrogen). Eggs were incubated in either 20 μM zinquin ethyl ester in combination with 1 μM Syto 64 (Invitrogen) or 10 μM FluoZin-3 AM in combination with 10 $\mu g/ml$ Hoechst 33342 (Invitrogen) for 60 min in KSOM (Millipore). They were imaged in 20 μl drops of KSOM covered with oil for embryo culture (Irvine Scientific) on glass-bottom dishes (Bioptechs Inc.). Images were acquired as Z-stacks on a TCS SP5 confocal microscope (Leica Microsystems) equipped with a stage top incubator (Tokai Hit), 63× oil-immersion objective, and HeNe (543 nm), Ar (488 nm), and near-UV (405 nm) laser lines.

TPEN treatment and spindle imaging of mouse eggs

Superovulated eggs were collected and transferred to KSOM medium (Millipore) with or without 10 μ M TPEN and cultured up to 8 hrs at 37 °C in an atmosphere of 5% CO₂ in air. Previous work showed that treatment of maturing oocytes with 10 μ M TPEN could selectively limit the intracellular acquisition of zinc (relative to iron and copper) but without significant toxicity¹⁴. Eggs were fixed and permeabilized for 30 min at 37 °C in a solution containing 2% formaldehyde, 2% Triton X-100, 100 mM PIPES, 5 mM MgCl₂, and 2.4 mM EGTA. Eggs were then washed and blocked for 1-3 hrs in 1× PBS containing 0.1 M glycine, 3 mg/mL BSA, 0.01% Tween-20, and 0.01% sodium azide followed by incubation with anti- α -tubulin (1:100, Sigma) in blocking buffer for 1 hr at 37 °C. Eggs were washed again in blocking buffer, incubated in Alexa Fluor 488-conjugated goat anti-mouse IgG (Invitrogen) and rhodamine-phalloidin conjugate (Invitrogen) at 37 °C for 1 hr, washed three additional times, and mounted on microscopy slides with coverslips in Vectashield with DAPI. Images were acquired as Z-stacks on a TCS SP5 confocal microscope (Leica Microsystems) equipped with a stage top incubator (Tokai Hit), 63× oil-immersion objective, and HeNe (543 nm), Ar (488 nm), and near-UV (405 nm) laser lines.

Statistical Analysis

The abundance of each element was compared across time points using one-way analysis of variance assuming different standard deviations for each group. These analyses were followed by pairwise comparisons of each time versus each other time using the Tukey-Kramer method³⁶. All statistical tests were performed using the software Prism 4.0 (GraphPad Software, San Diego, CA). P < 0.05 was considered statistically significant.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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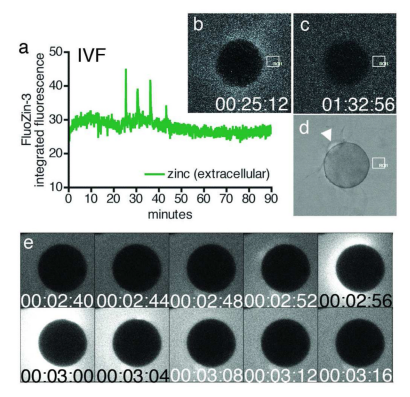


Figure 1. Zinc is released into the extracellular environment as an early fertilization event Changes in extracellular zinc concentration are readily monitored with FluoZin-3 during *in vitro* fertilization, and rapid, repetitive increases in fluorescence intensity were detected (**a**) by ROI analysis (denoted by white boxes in **b-d**). Each zinc spark can be distinguished in the time-lapse series (**b**, 00:25:12) against background fluorescence (representative example in **c**, 01:32:56). Successful fertilization was confirmed by the extrusion of a second polar body (**d**). Zinc sparks were also noted during strontium chloride-induced parthenogenesis (**e**, 00:02:56). Time is expressed as hh:mm:ss, wherein 00:00:00 represents the start of image acquisition.

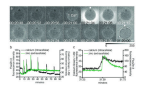


Figure 2. Zinc sparks are polarized and are immediately preceded by intracellular calcium transients

Shortly following activation, zinc sparks occur around the egg cortex with the exception of a zinc spark-free region ($\bf a$, 00:21:04, arrowhead). This spark-free region corresponds to the region containing the meiotic spindle, where the second polar body is extruded ($\bf a$, 01:20:00, arrowhead). Egg activation was confirmed by simultaneously monitoring intracellular calcium oscillations with Calcium Green-1 AM every 4 s ($\bf b$). Intracellular calcium increases immediately before a zinc spark, as evident when images were collected at a faster acquisition rate of every 100 ms in an independent experiment ($\bf c$). Time is expressed as hh:mm:ss, wherein 00:00:00 represents the start of image acquisition. In all cases, extracellular zinc was detected with FluoZin-3.

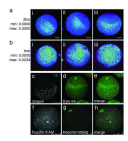


Figure 3. Zinc ions are cortically polarized in the mouse egg

Total zinc, as detected by synchrotron-based x-ray fluorescence ($\bf a$, $\bf b$), is uniquely polarized in the unfertilized egg ($\bf a$; i-iii represent replicates). This distribution is absent in the other essential transition elements, such as iron ($\bf b$; i-iii represent replicates). The minimum and maximum range of each group of images are given units of $\mu g/cm^2$. Labile (chelatoraccessible) zinc, as detected by confocal microscopy ($\bf c$ - $\bf h$), also has a hemispherical distribution in the live egg, as detected by two chemically distinct zinc fluorophores: zinquin ethyl ester ($\bf c$ - $\bf e$) and FluoZin-3 AM ($\bf f$ - $\bf h$). Co-staining with a DNA marker (Syto 64 in $\bf d$, Hoechst 33342 in $\bf g$) revealed that zinc was concentrated at the vegetal pole away from the meiotic spindle. The images depicted here are projected images of the complete Z-series; representative slices from these same Z-stacks confirming cortical localization of zinc are shown in Fig. S5. Scale bar = 20 μ m ($\bf a$, $\bf b$) or 25 μ m ($\bf c$ - $\bf h$).

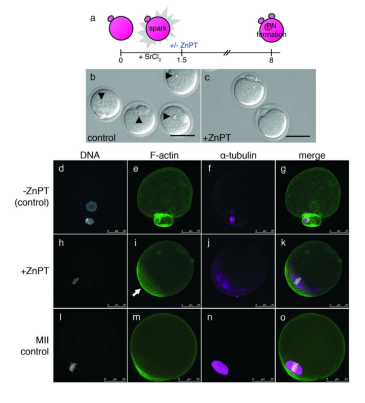


Figure 4. Sustained elevation of intracellular zinc availability following egg activation leads to reestablishment of metaphase arrest

Eggs were activated with strontium chloride (SrCl₂) then treated with zinc pyrithione (ZnPT) 1.5 hours later (a). At 6 h post-activation, control eggs form pronuclei (b, arrowheads) whereas the majority of eggs treated with ZnPT do not (c). When visualized by fluorescence, control eggs display decondensed DNA organized within a defined nucleus (d). F-actin is homogeneous around the egg's cortex (e) and α -tubulin remains organized as a spindle midbody remnant (f), which is visible between the egg and the second polar body (g, merged image). In contrast, ZnPT-treated eggs display condensed chromosomes (h) adjacent to an area of concentrated, cortical F-actin (i). α -tubulin is organized in a metaphase-like configuration (j) around the chromosomes (k, merged image). This layout mirrors the subcellular arrangement in unfertilized eggs (l-o), which also display condensed chromosomes (l) overlaid with an actin cap (m), surrounded by a metaphase spindle (n). Scale bar = 80 μ m (b, c) or 25 μ m (d-o).

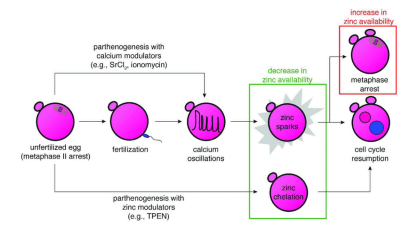


Figure 5. Zinc plays a gatekeeping role, switching between cell cycle arrest and resumption The unfertilized egg can now be activated by three different approaches, all of them leading to a decrease in intracellular zinc availability. The first is through the physiological pathway, fertilization (center row). This induces calcium oscillations in the egg, leading to the zinc sparks and subsequent cell cycle resumption. The two other approaches induce parthenogenesis and are pharmacological, either using modulators of intracellular calcium (top arrow) or zinc (bottom arrow). The calcium-based approaches induce zinc sparks, while the zinc-based approach bypasses the need for calcium oscillations by directly decreasing the availability of zinc in the egg (green box). Furthermore, cell cycle arrest at metaphase can be reestablished by raising zinc availability (red box), thus implicating zinc as a central regulator of the cell cycle during oocyte maturation and fertilization.