Endoplasmic reticulum generates calcium signalling microdomains around the nucleus and spindle in syncytial Drosophila embryos

H. Parry, A. McDougall and M. Whitaker

Institute for Cell and Molecular Biosciences, Medical School, University of Newcastle upon Tyne, Newcastle upon Tyne NE2 4HH, U.K.

Abstract


Introduction

Calcium signals control the cell division cycle of early sea urchin [1–3], frog [4–6] and mammalian embryos [7,8]. There is evidence that blocking calcium signals prevents mitosis, but often mitotic calcium signals are small or undetectable [3,7,9–11].

In early syncytial nuclear divisions of Drosophila melanogaster embryos, ER (endoplasmic reticulum) is highly concentrated around the nucleus at prophase and is also associated with the spindle poles [12]. This configuration of ER creates a calcium signalling microdomain [13].

ER separates a calcium signalling microdomain around the nucleus from a cortical calcium domain

Early Drosophila development involves rapid nuclear divisions that occur in the same cytoplasm [14,15]. Dividing nuclei are first located deep within the embryo and migrate to the embryo cortex during cycles 8 and 9. In later cycles, the nuclei divide just beneath the surface of the embryo, making them amenable to confocal imaging. Co-labelling of microtubules and ER during mitosis shows that the mitotic spindle forms in an environment bounded by ER (Figure 1A), with ER particularly prominent at the spindle poles. [Ca]i (intracellular free calcium concentration) varies markedly in the embryo during successive nuclear cycles (see Supplementary Video 1 at http://www.biochemsoctrans.org/bst/034/bst0340385add.htm). Highest levels of cortical [Ca], occur during interphase [13]. Nonetheless, even in interphase, ER surrounds the nucleus and forms a privileged calcium environment in which calcium concentrations are lower in the nucleus than in the cortex [13]. Figure 1(B) shows simultaneous measurement of calcium and ER in the cortex of a syncytial embryo at the time of prophase and illustrates how the disposition of ER generates two calcium signalling domains within the cortical bud. As nuclei proceed through mitosis and chromosomes separate, the relation between ER and the spindle becomes quite complex, with invasion by ER of the spindle mid-zone and accretion around the separating chromosomes (Figure 1C).

Calcium release at mitosis in the nuclear microdomain requires both InsP3 and extracellular calcium

Cell cycle calcium signals are triggered by InsP3 [1,2,6]. Drosophila embryos express an InsP3 receptor [16,17]. Heparin and Xestospongin C are agents that inhibit the interaction of InsP3 with the InsP3 receptor [18,19]. Figure 2(A) shows that interfering with InsP3 signalling using either Xestospongin C [20] or heparin [1] as antagonists prevents the completion of nuclear division by interfering with chromatid disjunction, as visualized using histone-tagged chromatin. Complete arrest occurs over three nuclear cycles at a concentration of approx. 125 µg/ml heparin.

We investigated the requirement for extraembryonic calcium during the syncytial nuclear divisions. In Drosophila embryos, extracellular fluid is contained in the perivitelline space outside the plasma membrane, bounded by the vitelline membrane. We titrated the calcium concentration in the perivitelline space using dibromoBAPTA [where BAPTA is bis-(o-aminophenoxy)ethane-N,N,N′,N′-tetra-acetic acid]. Calcium Green Dextran; [Ca], intracellular free calcium concentration; ER, endoplasmic reticulum.

Abbreviations used: BAPTA, bis-(o-aminophenoxy)ethane-N,N,N′,N′-tetra-acetic acid; CaGr, Calcium Green Dextran; [Ca], intracellular free calcium concentration; ER, endoplasmic reticulum.

1 To whom correspondence should be addressed (email michael.whitaker@ncl.ac.uk).

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as judged by the fall in the signal from the calcium indicator CaGr (Calcium Green Dextran) (Figure 2B, i). Microinjection of dibromoBAPTA to a concentration of 5 mM did not alter the CaGr signal (Figure 2B, ii), demonstrating that the calcium concentration in the perivitelline space was greater than 5 mM, but less than 10 mM. We found that injection of 5 mM dibromoBAPTA into the perivitelline space did not affect the nuclear division cycle, but that reducing perivitelline calcium concentrations to micromolar levels with 10 mM dibromoBAPTA led to cell cycle arrest (Figure 2B, iii).

Discussion
The syncytial nuclear divisions of early Drosophila embryos are well suited to investigation of the role of calcium in mitosis because ER does not invade the nucleus, instead surrounding the nucleus and mitotic spindle [12]. This disposition of ER defines spatially identifiable domains in which we have shown that calcium concentrations are differentially regulated. While calcium signalling microdomains have never previously been identified during mitosis, there are reports that calcium signals during mitosis are confined to the mitotic spindle pole [21,22]. The principle that calcium signalling subdomains can exist within a single cell bounded by ER has been well established in pancreatic acinar cells [23].

Here, we show that ER separates two calcium signalling domains within the cortical buds of syncytial Drosophila embryos. The dominant and striking calcium signal in these embryos is cortical and is associated with actin cytoskeletal rearrangements [13]. High [Ca], correlates with the phase of cortical contraction associated with interphase nuclei [14] and inhibited by cytochalasins [24]. The ER forms a
Figure 2 | Block of nuclear division by inhibition of InsP₃ signalling and by removal of extracellular calcium

(A) In separate experiments, spindles and chromosomes were tagged with fluorescein-labelled tubulin or fluorescein-labelled histones. (i) Control to show normal progress through mitosis, with a doubling of nuclei. (ii, iii) The InsP₃ antagonist Xestospongin C prevents anaphase onset: metaphase spindles form normally, but chromosomes fail to segregate at anaphase. (iv, v) Heparin leads to a similar outcome as in (ii) and (iii); the arrow in (v) identifies a single nucleus for clarity. (vi) Inhibition of mitosis by Xestospongin C is progressive, with 75% of nuclei arrested after one cell cycle and the remainder in the next cell cycle. (vii) Dose-response curve for heparin. (B) The perivitelline space was first labelled by injection of CaGr. The calcium chelator dibromoBAPTA was then injected to a final concentration of (i) 10 mM or (ii) 5 mM. DibromoBAPTA (10 mM) markedly reduced the CaGr signal, indicating that the chelator reduced the calcium concentration of the perivitelline space to micromolar levels, while 5 mM dibromoBAPTA was unable to effect this. These results indicate that the calcium concentration in the perivitelline space is between 5 and 10 mM. (iii) The nuclear division cycle is fully disrupted by reducing extracellular calcium to micromolar levels. For methods see [13].

![Image](image_url)

Bar chart showing the percentage of nuclei arrested at different stages of the nuclear cycle.

Bar chart showing the percentage of embryos arrested at different doses of heparin.

**Bar**

Barrier between the cortex and the nucleus. During episodes of high cortical [Ca], it preserves a lower calcium concentration within the nucleus; at a time when cortical [Ca] is low, the ER generates a calcium signal within the spindle that triggers anaphase [13].

These observations in Drosophila embryos demonstrate that calcium concentrations in the nucleus and spindle are controlled independently of bulk cytoplasmic calcium concentrations. It is known that in most cell types, ER associates intimately with the nucleus and spindle [25]. This has been interpreted as a mechanism to ensure proportionate inheritance of ER when cells divide [26]. The intimate association of ER with the nucleus and spindle suggests that similar calcium signalling microdomains are present during mitosis in other cell types. However, the close intermingling of ER and spindle microtubules in most cell types may make very local calcium signals within the dividing nucleus more difficult to resolve and explain why such signals are not always readily detectable [3,7,9–11].

We also show here that the InsP₃-dependent calcium signals that control entry into and exit from mitosis in syncytial Drosophila embryos are dependent on extraembryonic...
calcium. Calcium from the perivitelline space is essential to maintain the nuclear division cycles. It will be interesting to determine whether calcium from the extracellular space can tunnel through ER directly to the mitotic apparatus, as had been shown to occur in the apical pole of pancreatic acinar cells [27,28].

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References


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