Cell Cycle Dynamics and Endoreplication In the Mosquito Midgut

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Abstract
The mosquitoes are the major transmitters of vector-borne infections, including virus, protozoans and helminths and, during the interaction it is critical for pathogens to accomplish immune evasion actions whereas the mosquito turn on immune mechanisms for its survival. The mosquito midgut is the organ where initial interactions with pathogens happen, and the insect has mechanisms to repair the tissue and to eliminate the invaders. For these tasks’ mosquitoes use different strategies including the specific DNA replication, amplifying specific genes and without cell division in order to have more DNA templates for transcription that allow, in a secondary infection, the fast production of immune proteins in a process known as priming. The study of these processes is necessary for the understanding of the mechanisms during insect-pathogens interactions, and to design future interventions to block mosquito pathogens transmission that could be environment-friendly and not depending on costly chemicals.

Introduction
The mosquito midgut is an essential tissue where pathogens and microorganisms interact. This organ is fundamental to transmit diverse diseases, including the virus causing dengue, zika, chikungunya, and the malaria causing parasite Plasmodium sp. During the interaction with pathogens, the midgut cells react; repairing cell damage and producing immune molecules to fight against invaders. However, we have limited knowledge of the mechanisms behind the response and the cell-damaged repair. The cell cycle is a finely regulated process that manages proliferation and differentiation in cells, which is controlled by a coordinated expression of specific molecules, notably proteins called cyclins. Cyclins are expressed in a particular sequence in specific cell cycle phases and are critical elements for cycle progression [1]. The conventional cell cycle phases include Gap 1 phase (G1), synthesis phase (S), Gap 2 phase (G2), and end in mitosis (M) where the cell splits into two daughter cells. In insects exist specific situations where cells can vary the plan and generate cells with different ploidy and/or DNA content, as in polytenic cells happening in dipteran salivary glands. Interestingly, the machinery that manages the conventional cell cycle is the same, participating in different options of the cell cycle [2-4].

One of the versions of a specialized cell cycle is the endoreplication. Endoreplication is a variant of the normal replicative cell cycle, in which cells increase their genomic DNA content without division. Endoreplication can enclose different options of the cell cycle; such as endocycles, re-replication, and endomitosis. The first one consists of repeated successions of S – G phases of all genetic material, without cell or nuclei division. In re-replication DNA synthesis is initiated multiple times at individual origins of replication within the same S phase, provoking site-specific replication of a unique sequence. In endomitosis an entry into mitosis occurs, the cells condense the chromosomes but do not dissociate them to daughter cells. Instead, they re-enter a similar phase to G1 and S phase starts again resulting in a multiple nuclei cell. [5]. The typical illustration of endoreplication is the generation of polytenic chromosomes in Drosophila salivary glands, but in many others, insects occur endoreplication. For example, in the beetle Tribolium castaneum larval stages, intestinal stem cells (ISC) conduct endoreplication for adult midgut polypliod epithelium formation [6]. Also, in the flour moth Euphestia kuehniella, nuclei in Malpighian tubules and silk glands increase in size through larvae in stars. Even, in the last instar,
larvae nuclei are polyploid with a high DNA content, provoking a branched nucleus. This polyploidy that results in branched nuclei, could be considered an adaptation because the distance between the nuclear area and the cytoplasmic zone is increased to permit traffic of molecules produced in high quantities [7].

Here, we discuss the cell cycle in the midgut development and the mechanisms conducting normal cell cycle into endoreplication. We describe endoreplication in the mosquito midgut as a fundamental part of its homeostasis. Finally, we also consider infection as a stressor when tissue undergoes damage and immune stimulation.

**Cell cycle regulation during development**

The regulation of the cell cycle depends on master mechanisms on several levels. Hormones that coordinate the cell cycle between different body parts and more fine mechanisms which act at the cellular/molecular level. Signaling pathways that detect environmental cues, including the hormones and chemicals, master molecules, as cyclins, kinases and phosphatases, that are synthesized and activated/desactivated or destroyed in a strict sequence in order to stop or induce the change from one to next cell cycle phases [8-11] The two main insect hormones, the Juvenile Hormone (JH) and the steroid hormone ecdysone and its active metabolites as 20-hydroxyecdysone (20E) regulates development and reproductive maturation in insects.

JH is produced and released by the corpora allata (CA), endocrine glands connected brain [12], and their actions regulate reproduction, behavior and diapause, between others. In Locusta migratoria, the allatotomy which leads to depletion of CA, inhibits de novo DNA synthesis. Application of Methoprene, a JH analog, restores JH-induced activities, including DNA synthesis and polyplidization in fat body cells. The mechanism involves methoprene-tolerant (Met), a JH receptor that induces transcriptional activation in minichromosome maintenance genes 4 and 7 (Mcm4, Mcm7), both genes involved in genome replication in fat body cells [13]. JH regulates the machinery of the cell cycle to originate polyplid cells supporting the massive synthesis of vitellogenin, one of the main proteins in eggs maturation. Also, another mechanism in fat body cell suggests that JH directs Cdk6 and E2f expression causing polyplid cells [14]. Methoprene is an analog of JH, used for insect control, that is not degraded and maintains its effect for a long time and is useful to evaluate JH actions. High methoprene concentrations avoid cell division in the midgut of larval-pupal transition, and it produces different grades of polyploidy in Ae. aegypti [10].

On the other hand, ecdysone signaling is an essential mediator in the switch between endocyte and site-specific endoreplication, trough the binding to the ecdysone receptor (EcR). At signaling pathways level, in epithelial follicle cells during oogenesis, Notch signaling is a mechanism that modulates endocycles, switching to site-specific gene amplification. Notch is down regulated while ecdysone signaling is activated and cooperate with Tramtrack (Tkt), a transcription factor that induces endocyte exit and entry in site-specific endoreplication [15]. Moreover, steroid hormone 20-hydroxyecdysone (20E) arrests in G1 phase the mosquito cell line C7-10 from Aedes albopictus and at the same time occurs down-regulation of cyclin A [16].

A **“low-cost” strategy**

Cell proliferation is a demanding process in material and energy terms. During oogenesis in Drosophila melanogaster, a stable nutritional status is important in ovariole to continue with the conventional proliferation program [17]. Also, in oogenesis, after mitotic proliferation occurs a transition to endoreplication that causes an amplification of specific DNA regions in which genes for essential situations are located, and that need to be mass-produced [18]. In nature, many organisms carry out endoreplication to provide a massive production of molecules. This strategy was eventually developed in cell types with high metabolic activity [19]. This hypothesis was proposed in the seeds of plants with endosperm. Endoreplication can enhance metabolic capacity because it occurs an increase in the gene copy numbers of genes related to metabolism, allowing synthesis of high quantities of relevant molecules which will be stored inside seed [20].

Replication, transcription, and translation have different energetic costs inside a cell. However, replication occurs at least one time, and transcription and translation are constant processes. An analysis in energetic terms from bacteria to eukaryotes indicates that the cost of a gene activity at the DNA, RNA, and protein level decline in relation with cell mass [21]. According to this, in specialized organs or cells, endoreplication would be, and actually it is, a very useful strategy to increase the cell and/or tissue size by polyploidy, generating cells efficient for specialized production during a short time avoiding mitotic cycles and spending less energy.

**Cell cycle dynamics after infections**

The hematophagous insect’s midgut is a monolayer of cells that can distend under the stretching of a large blood volume and absorb the nutrients from the blood meal. In mosquitoes, as in other insects, the midgut cells are differentiated in endocellular, columnar, and regenerative cells.

Mosquitoes are the principal bloodfed arthropods responsible for transmitting critical diseases (vector-borne diseases) such as malaria, dengue, Zika, chikungunya, between others. In mosquitoes, there are data suggesting that after infections or immune challenges in specific tissues, including the midgut, cell cycle dynamics change [22], initiating regeneration and/or immunity functions. Cell cycle activation has been documented in Aedes albopictus midgut after chemical and bacterial damage [23]. These stimuli produced an increase in regenerative cells, allowing to the midgut to equilibrate the homeostasis. Besides, after viral infections and oxidative stress, cell proliferation has been tested in Aedes aegypti midgut [24].
Evidence suggests that midgut infection is modulated by delayed or immediate activation of ISC to division. This activation is considered a strategy for tissue repair. However, polyploidization and growth by cell fusion also permit tissues to reach homeostasis with lower energy expenditure. It avoids the demand for an energetic process involving proliferation.

Metazoans maintain tissue size and morphology using cell division as a primary strategy to increase tissue size and repair. Although post-mitotic tissues can undergo compensatory cellular hypertrophy (CCI), consisting in hypertrophic growth without cell division, and this activation depends on the speed-up of endocycles rate, producing a cell with high DNA content and increased size to repair spaces in tissue after damage or apoptosis. Insulin/IGF (insulin-like growth factor)-like signaling pathway is involved in this strategy, intensifying endocycles rendering cells with increased cell size [25]. A hypothesis is that CCI strategy could be displayed during the insect immune response. In *Anopheles albimanus* tissues with a known relevant immune response and challenged with different microorganisms showed a significant DNA synthesis [22,26]. This DNA synthesis has been attributed to an endoreplication process because any cell division has been observed. DNA synthesis was evidenced by BrdU incorporation in various tissues after immune challenge with *Saccharomyces cerevisiae*. Besides, after colchicine treatments an inhibition in BrdU incorporation occurs [26]. These results suggest that DNA synthesis is an active process caused by an immune challenge. Likewise, in *Anopheles albimanus* DNA synthesis after *Plasmodium sp.*, challenge has been detected using BrdU incorporation [27]. This DNA synthesis is associated to an adaptive immune response, that in insects is known as immunological priming. Priming is defined as the ability to acquire a protective response to a pathogen as a consequence of previous exposure to the same organism. Such response has been probed in many invertebrates’ groups [28]. In *An. albimanus*, after a second exposure with *Plasmodium berghei* an increase in HNT, a protein that can regulate Notch pathway in proliferation and differentiation during development in follicular cells, [29] is detected. Similar results in *Ae. aegypti* after a second exposure to Dengue Virus [30] suggest an essential role of HNT in mosquito immunological priming and after oxidative stress [24]. The mechanism by which immunological priming occurs should be confirmed, but in mosquitoes, DNA synthesis and the expression of molecules related to endoreplication and stress-immunity response as antimicrobial peptides (AMP’s), are exciting results indicating that during the immune response the cell cycle is activated (Figure 1). In addition, morphologic evidence suggests that after *P. falciparum* invasion and damage of mosquito midgut epithelium, cells undergo division and differentiation leading to regeneration [31]. However, the mechanism in epithelial repair after cell damage by *Plasmodium gallinaceum* invasion in *Aedes aegypti* midgut involves an actin cone that mediates cells displacement without cell division evidence [32].

**Conclusion**

In insects, particularly in dipterans as flies and mosquitoes, cells needing the massive synthesis of molecules, required during reproduction and other tasks, carry-out the amplification of DNA segments, bearing genes specific to be used as templates for extensive transcription of messengers that will be used to produce useful molecules. This dynamic response is “low-cost” compared with other strategies, including the cell division, which needs a lot of material, energy, and time. There is increasing evidence that in mosquitoes, this strategy can be deployed to face immune challenges, and to prepare the individuals against repetitive pathogens assaults. These actions lead the mosquitoes to a “primed” status, in which the insect will be resistant to specific infections. Although there is a lot of molecules and mechanisms that are unknown in this critical group of insects. This is an exciting field that should be explored; to understand the mosquito-pathogens relations, and to design creative strategies to interfere with disease transmission.

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**Conflict of Interest**

Authors declare that do not have any conflict of interest.

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