

Lecture 14. Cell cycle I

Outline

- I. Overview of the cell cycle
- II. The cell cycle control system
- III. S phase

Reminder: cell theory

Cell comes from cell

I. Overview of cell cycle

- ♥ Phases of cell cycle
- ♥ Model systems for cell cycle studies
- ♥ Common methods to study cell cycle

1. Cell cycle

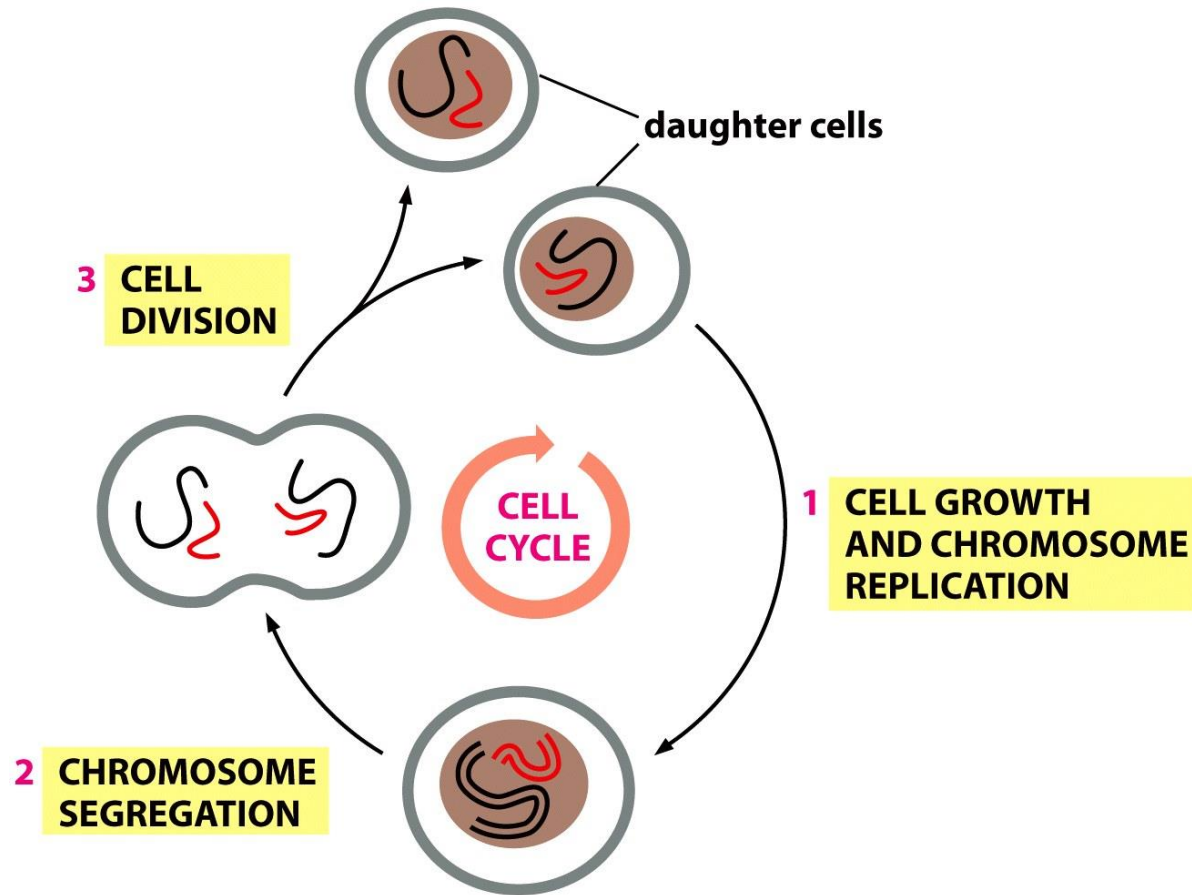
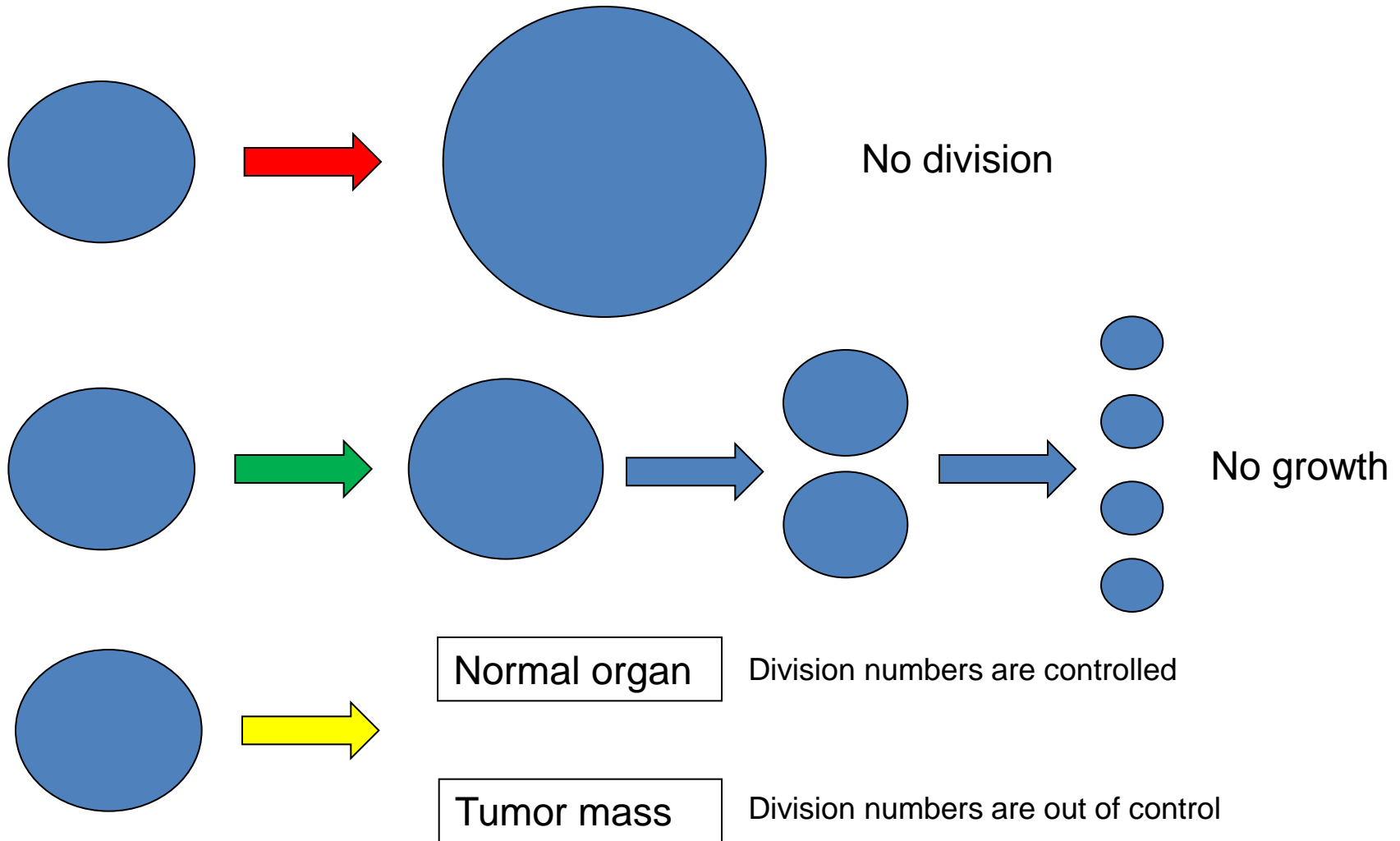


Figure 18-1 Essential Cell Biology 3/e (© Garland Science 2010)

What happens if this is dysregulated?



The major events in cell cycle

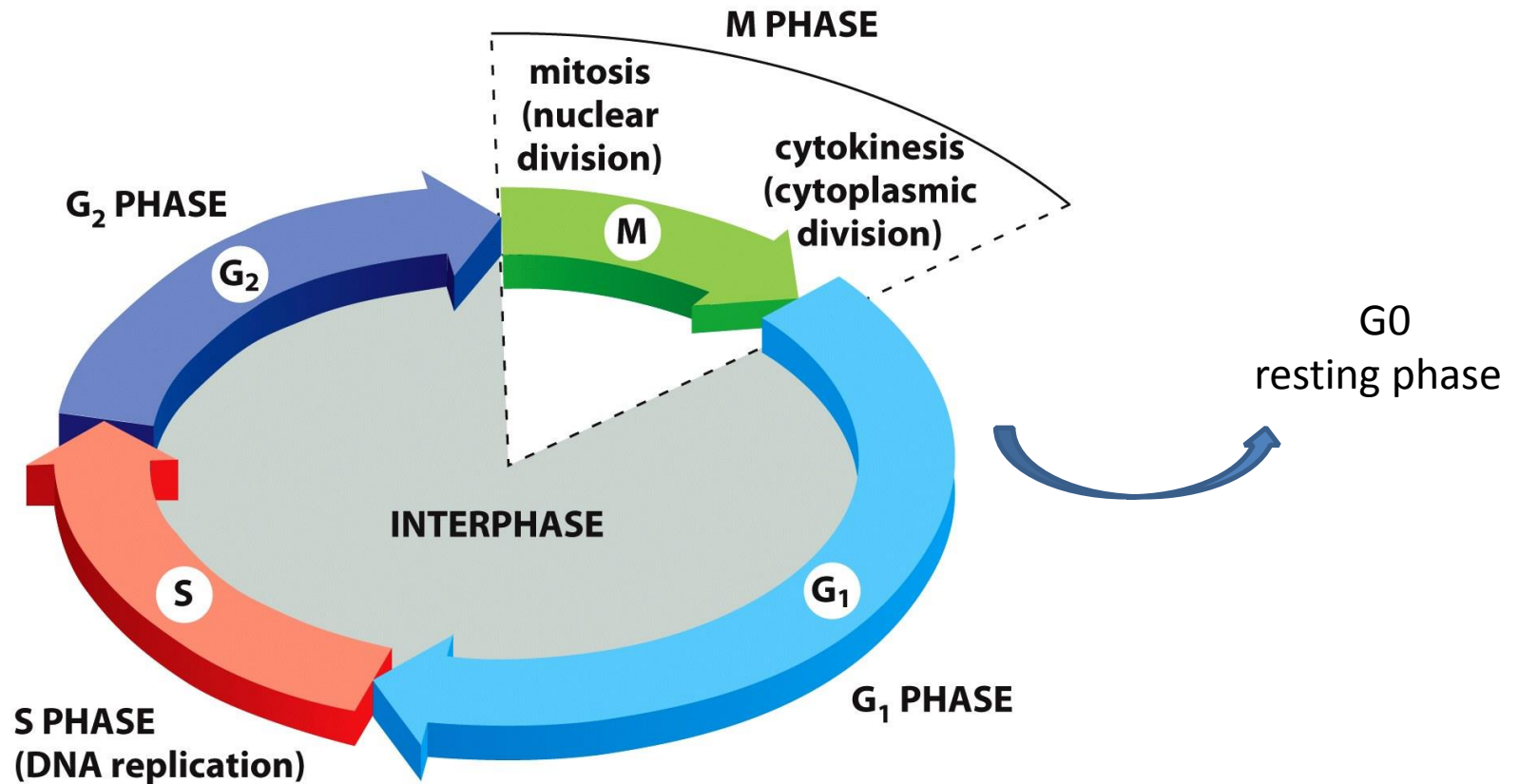
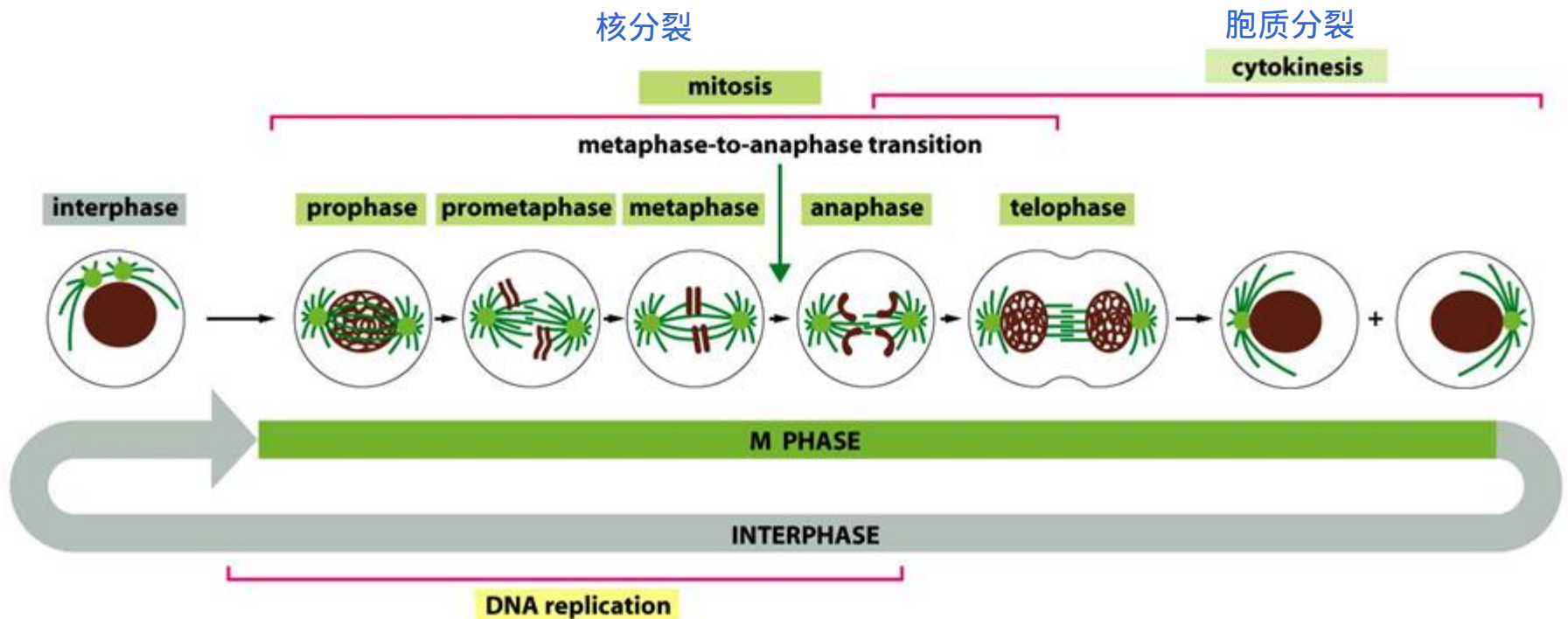


Figure 18-2 Essential Cell Biology 3/e (© Garland Science 2010)

M phase can be further divided into:
mitosis and cytokinesis



Different Cell cycle time (doubling time) for some eukaryotic cells

Fertilized xenopus oocytes	30 min
Yeast cell	1.5-3 hours
Mammalian intestine epithelial cells	~12 hours
Mammalian fibroblasts	20 hours
Human liver cell	~ 1 year

2. Model systems to study cell cycle

Cell cycle control mechanisms are conserved during evolution

♥ Yeast:

fission yeast (*S.pombe*)

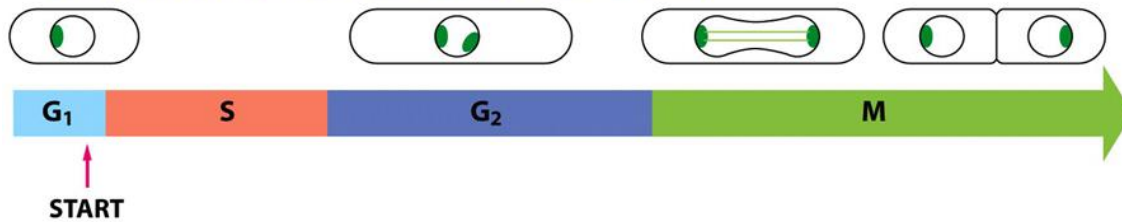
budding yeast (*S.cerevisiae*)

♥ Xenopus oocyte

♥ Mammalian cells

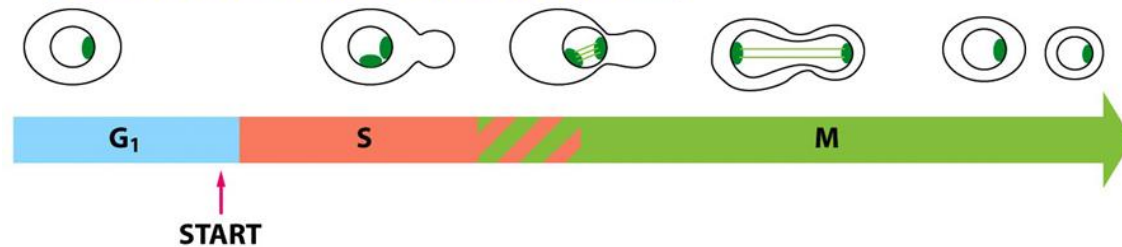
1) Yeast

(A) FISSION YEAST (*Schizosaccharomyces pombe*)



Fission yeasts are rod, septum
Divides into two daughter cells

(B) BUDDING YEAST (*Saccharomyces cerevisiae*)

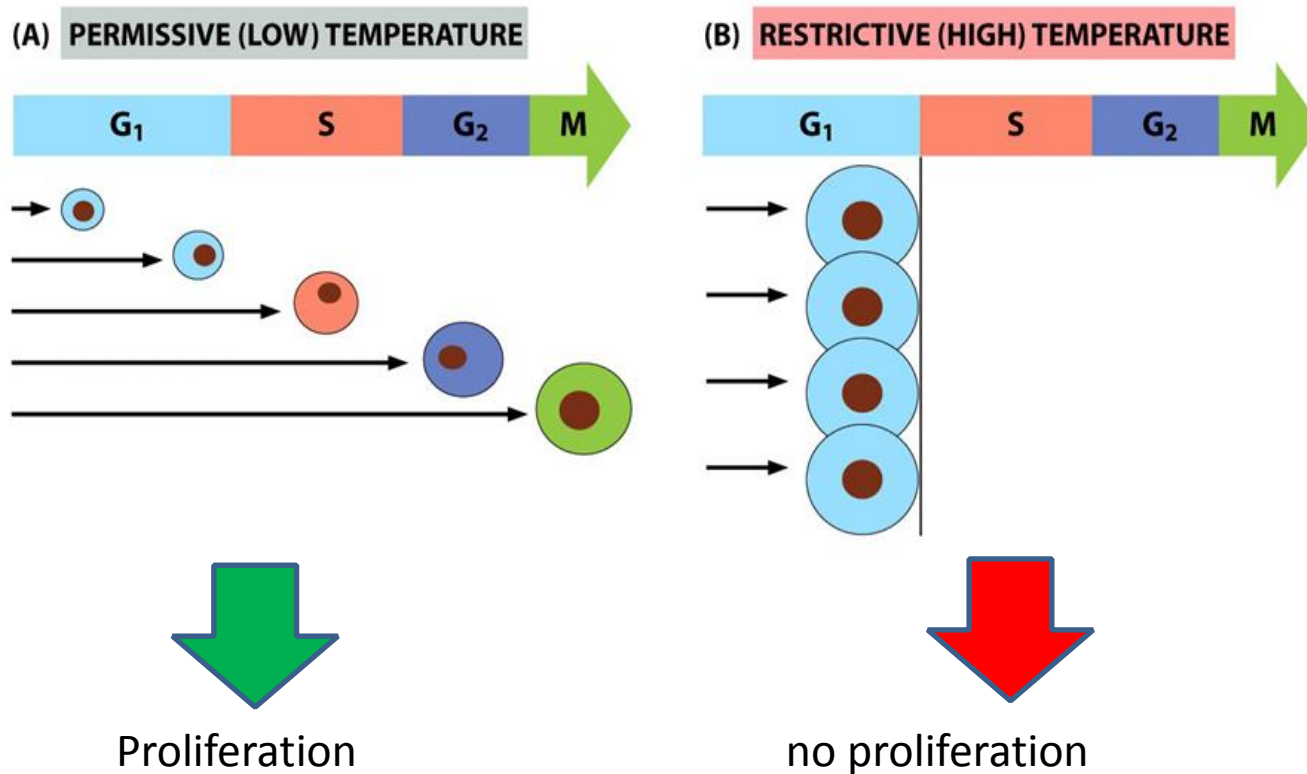


Budding yeasts are oval, daughter
cell buds from mother cell.

Conditional mutation in yeast cells for cell cycle studies

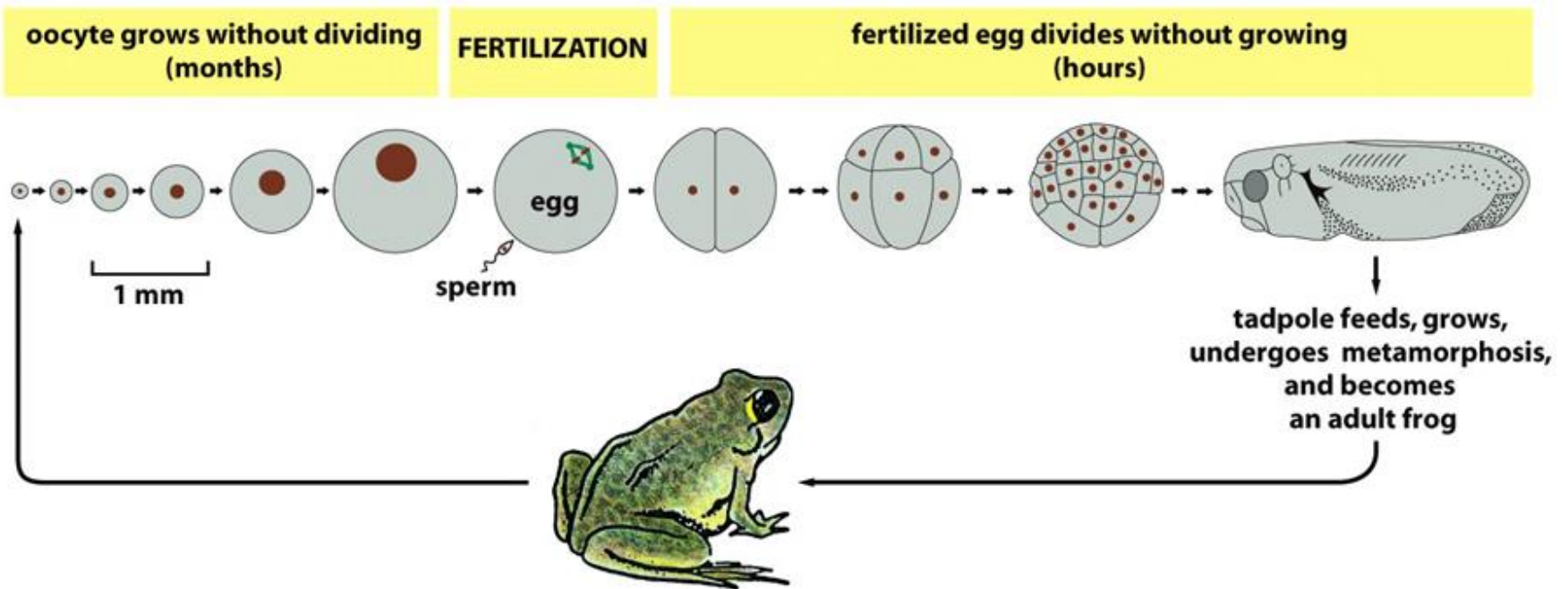
Start with haploid cells, perform genetic mutation

???

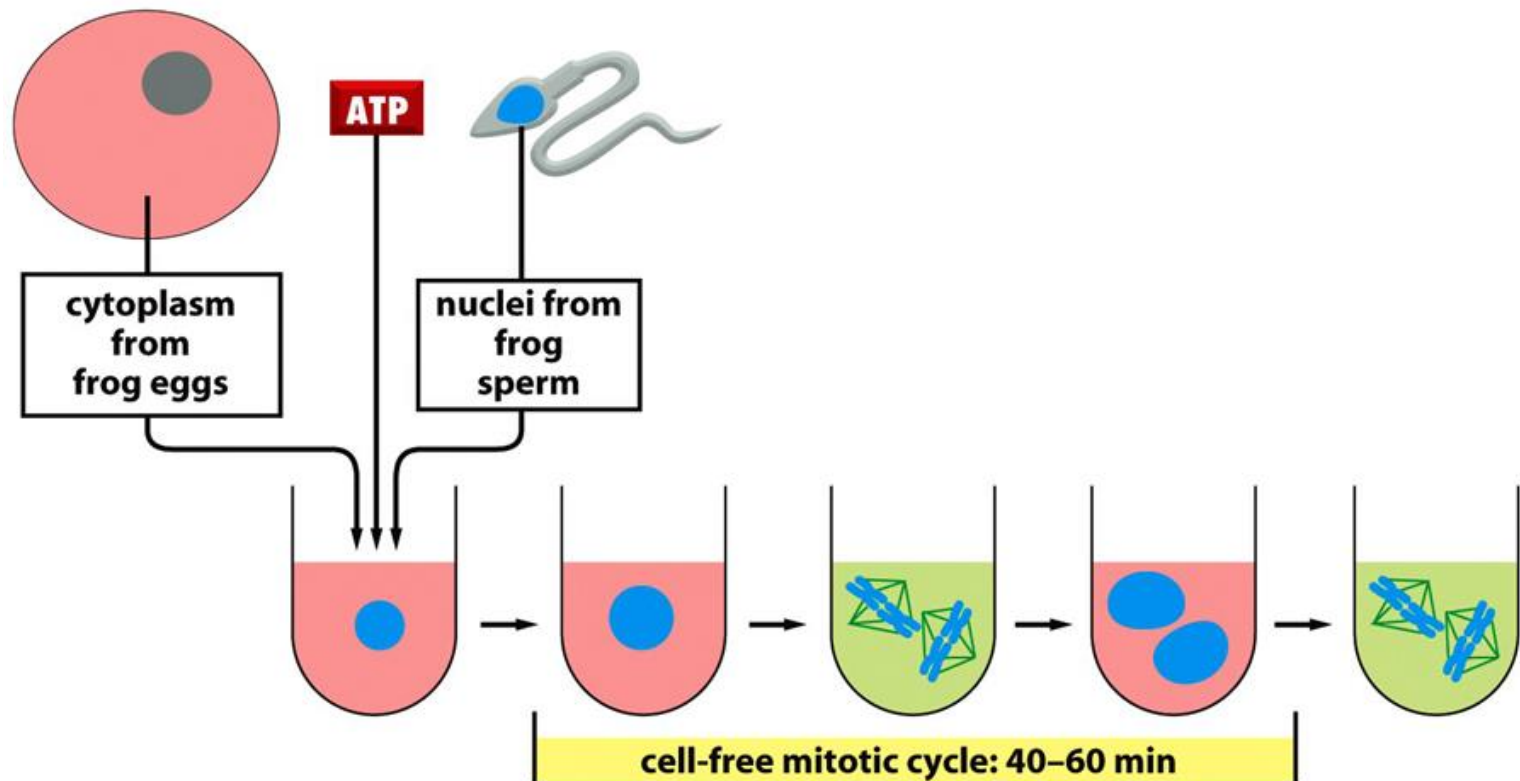


2). *Xenopus* oocytes

Xenopus oocytes have rich source of cell division proteins.

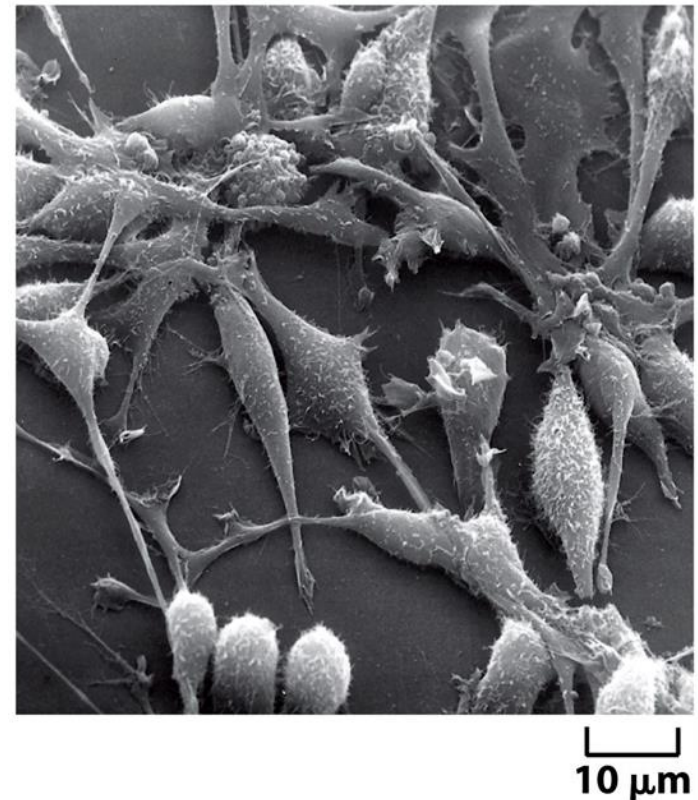


Xenopus oocytes are frequently used in cell-free system for *in vitro* studies



3). Cultured mammalian cells

- ♥ Normal primary cell culture
- ♥ Transformed immortal culture
- ♥ Cancer cell lines

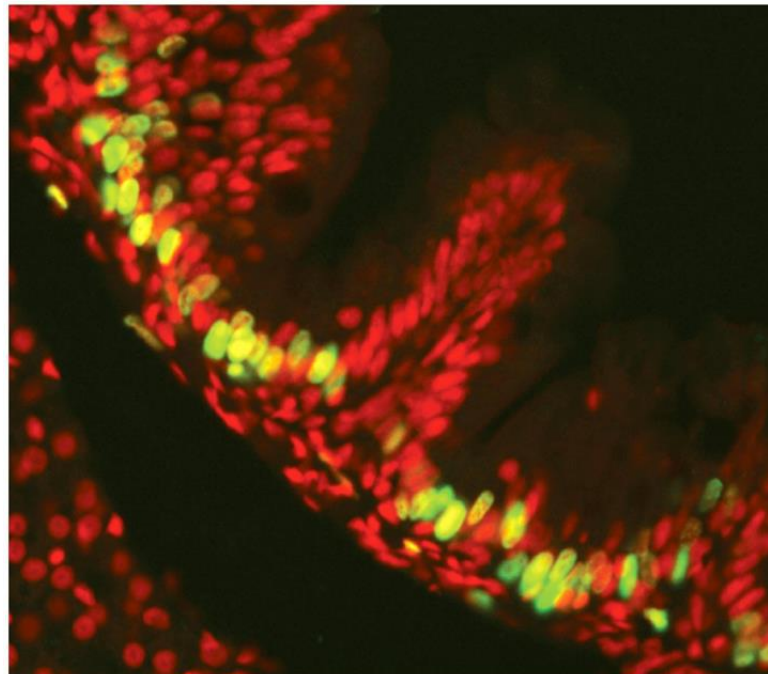


3. Various methods to study cell cycle

- ♥ Visualization under microscope
- ♥ BrdU/EdU incorporation assay
- ♥ Cell cycle distribution assay

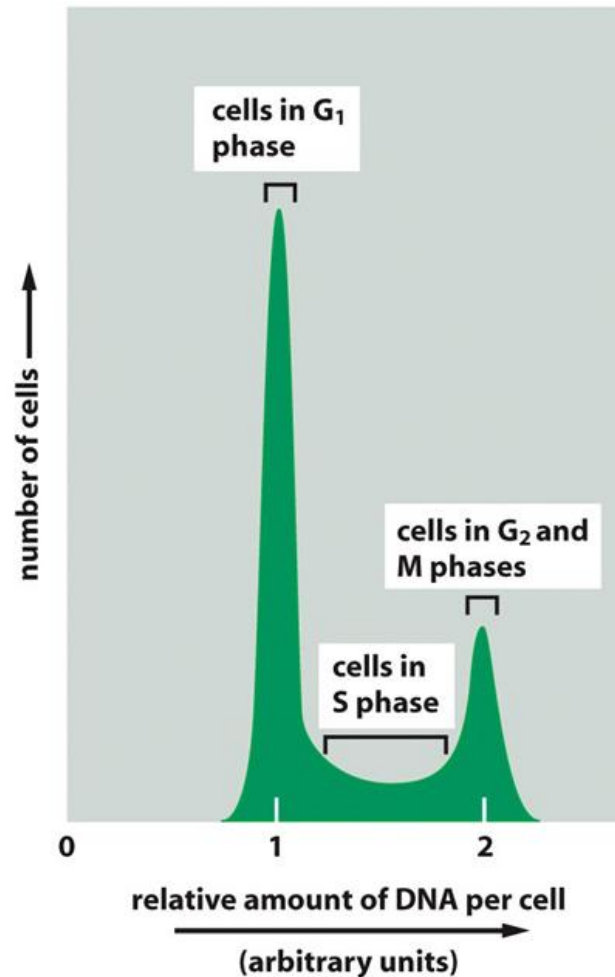
T analog incorporation analysis

treat cells with T analog BrdU (or EdU)
Stain by BrdU antibody (or with Click labeling)



Mitotic index:
 $\frac{\text{mitosis cell numbers}}{\text{Total cell numbers}}$

Flow cytometry to detect cell cycle phase

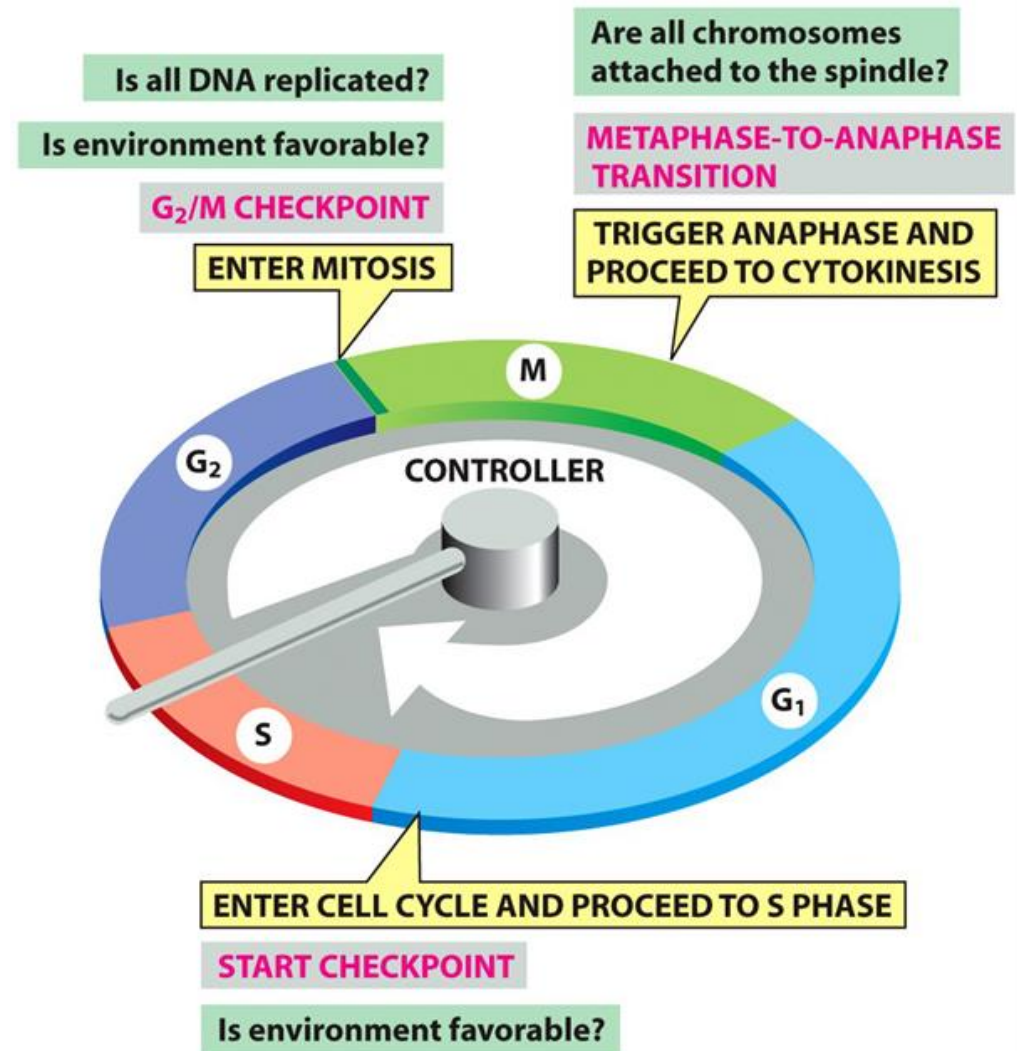


Use fluorescence dye to bind to DNA quantitatively, followed by flow cytometry

II. The cell cycle control system

Three major control checkpoints:

1. G₁/S transition
2. G₂/M Transition
3. Metaphase-to-anaphase transition



Cyclins and Cyclin-dependent kinase (Cdk) are two major players in cell cycle control

♥ Cyclins:

--- different cyclins oscillate in cell cycle and bind/control different cdk activity; it decides cdk substrates specificity and activates cdk.

♥ Cdk:

--- protein kinase, phosphorylates a subset of substrates to control cell cycle progression at specific checkpoint.

Cyclin/cdk complex

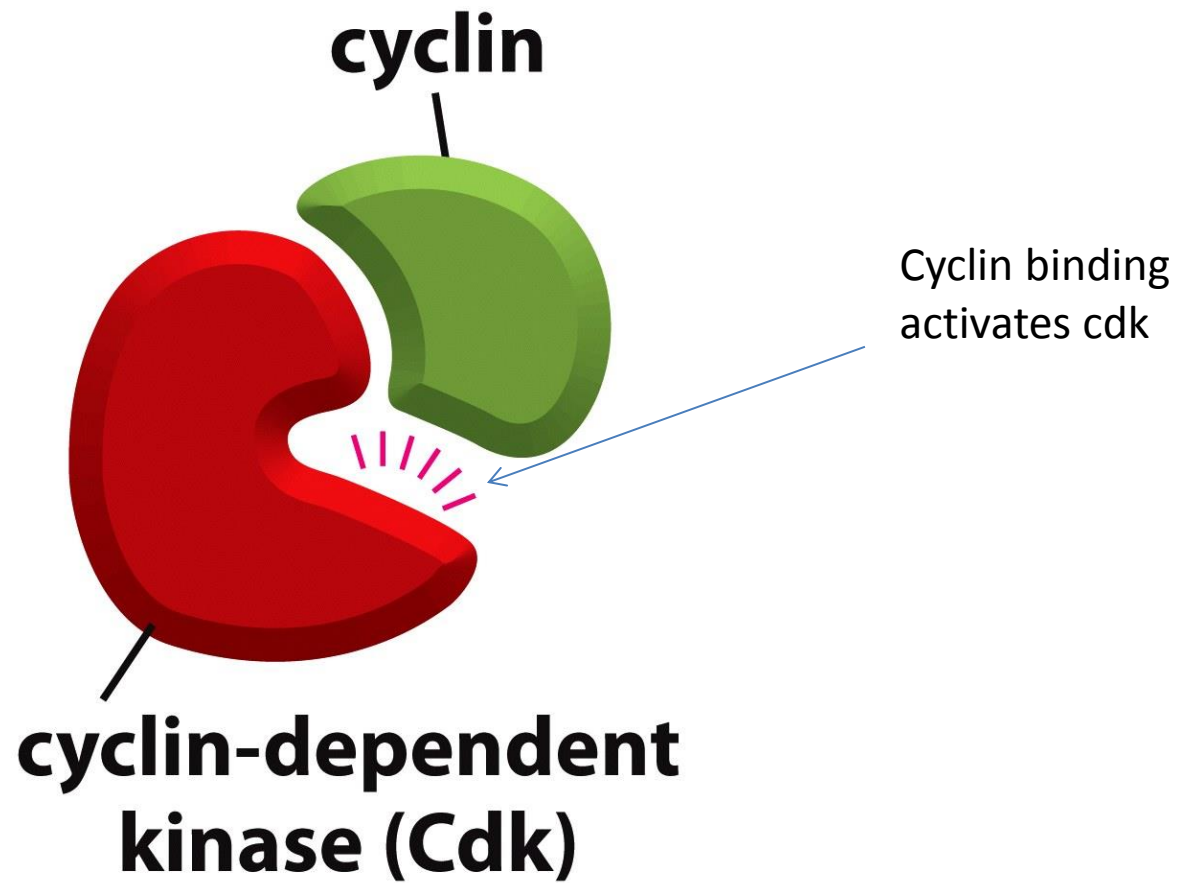


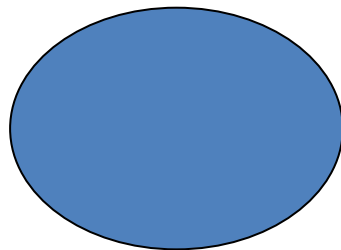
Figure 18-4 Essential Cell Biology 3/e (© Garland Science 2010)

The discovery of cyclins

- ♥ 1983, by Joan Ruderman and Tim Hunt
- ♥ Use oocytes from sea urchins and surf clams

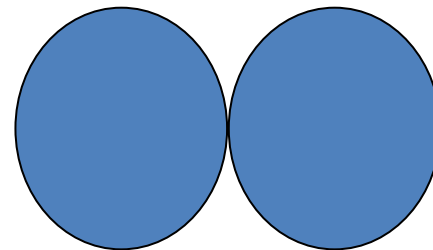
海胆

蛤蚌



oocyte

Add sperm



Fertilized egg division

Addition of sperm causes synchronized division for oocytes

♥ Synchronized

cells start from the same point in cell cycle

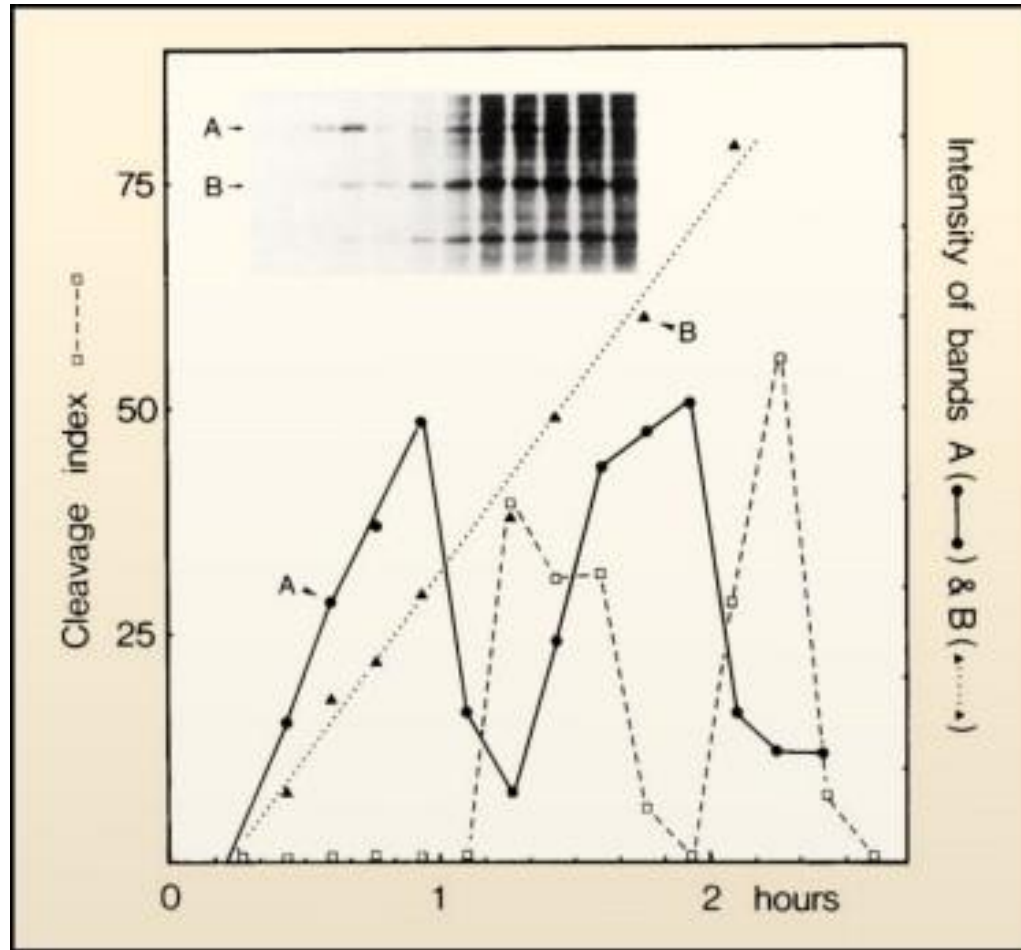
♥ Unsynchronized

cells start cell cycle differently

In oocytes: many mRNAs were not translated.

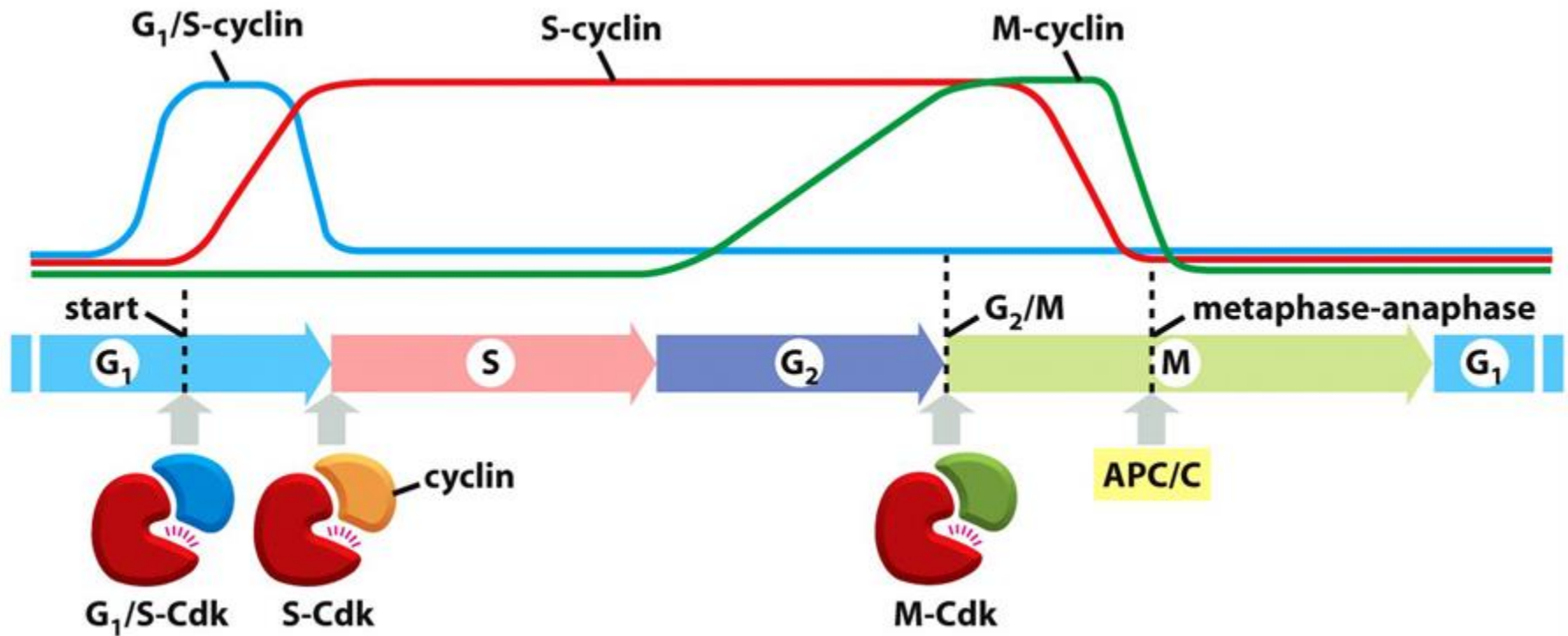
In fertilized eggs: many mRNA began to be translated.

The discovery of cyclins



Cyclin upheaval: gene transcription
Cyclin down: protein degradation

Cyclin-cdk complexes of the cell-cycle control system



Major cyclins and cdks of vertebrates and budding yeast

Table 17–1 The Major Cyclins and Cdks of Vertebrates and Budding Yeast

CYCLIN–CDK COMPLEX	VERTEBRATES		BUDDING YEAST	
	CYCLIN	CDK PARTNER	CYCLIN	CDK PARTNER
G ₁ -Cdk	cyclin D*	Cdk4, Cdk6	Cln3	Cdk1**
G ₁ /S-Cdk	cyclin E	Cdk2	Cln1, 2	Cdk1
S-Cdk	cyclin A	Cdk2, Cdk1**	Clb5, 6	Cdk1
M-Cdk	cyclin B	Cdk1	Clb1, 2, 3, 4	Cdk1

* There are three D cyclins in mammals (cyclins D1, D2, and D3).

** The original name of Cdk1 was Cdc2 in both vertebrates and fission yeast, and Cdc28 in budding yeast.

A fourth class of cyclin, cyclin G1, helps to govern the activities of the G1/S cyclins.

Cyclin degradation through ubiquitination

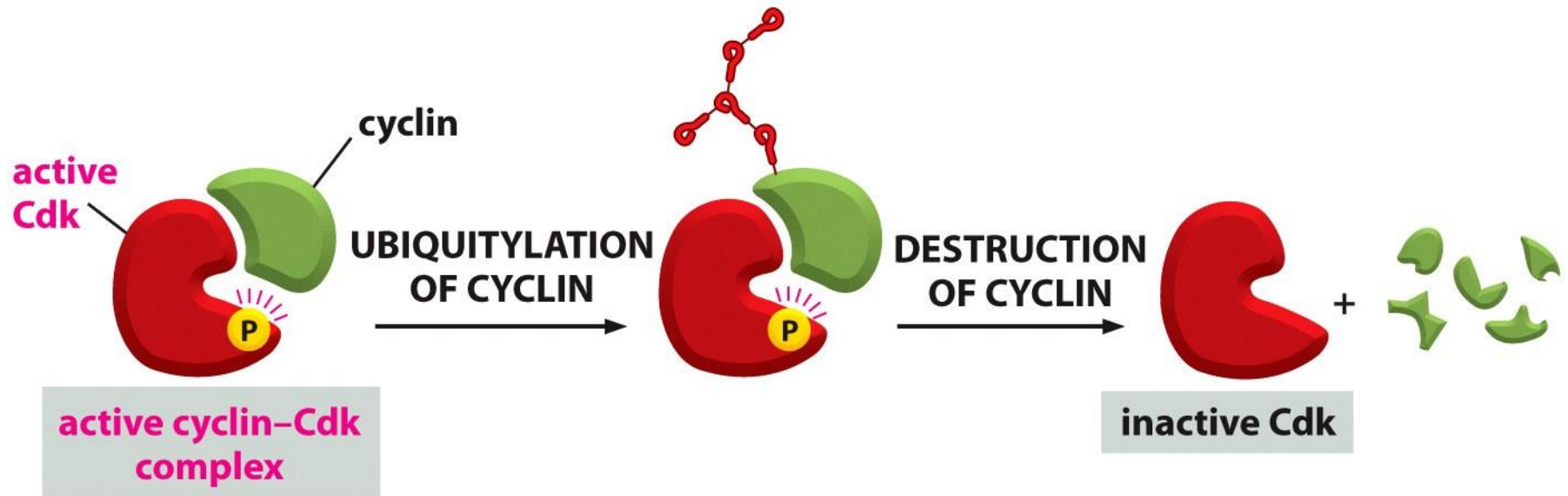


Figure 18-11 Essential Cell Biology 3/e (© Garland Science 2010)

Activation of cdk is additionally controlled by phosphorylation

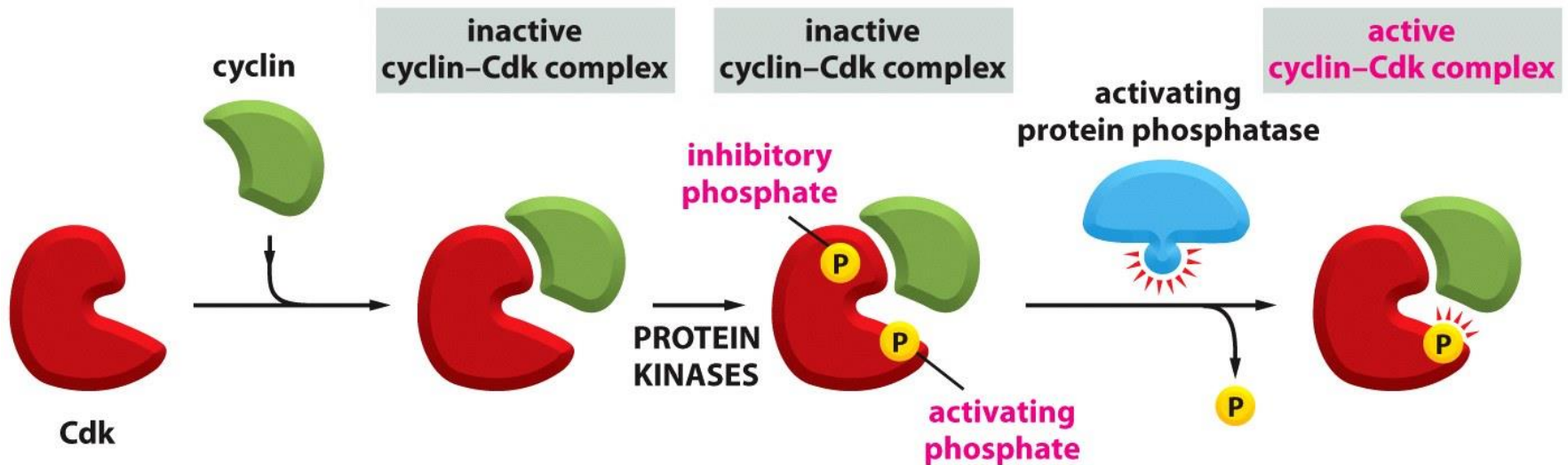
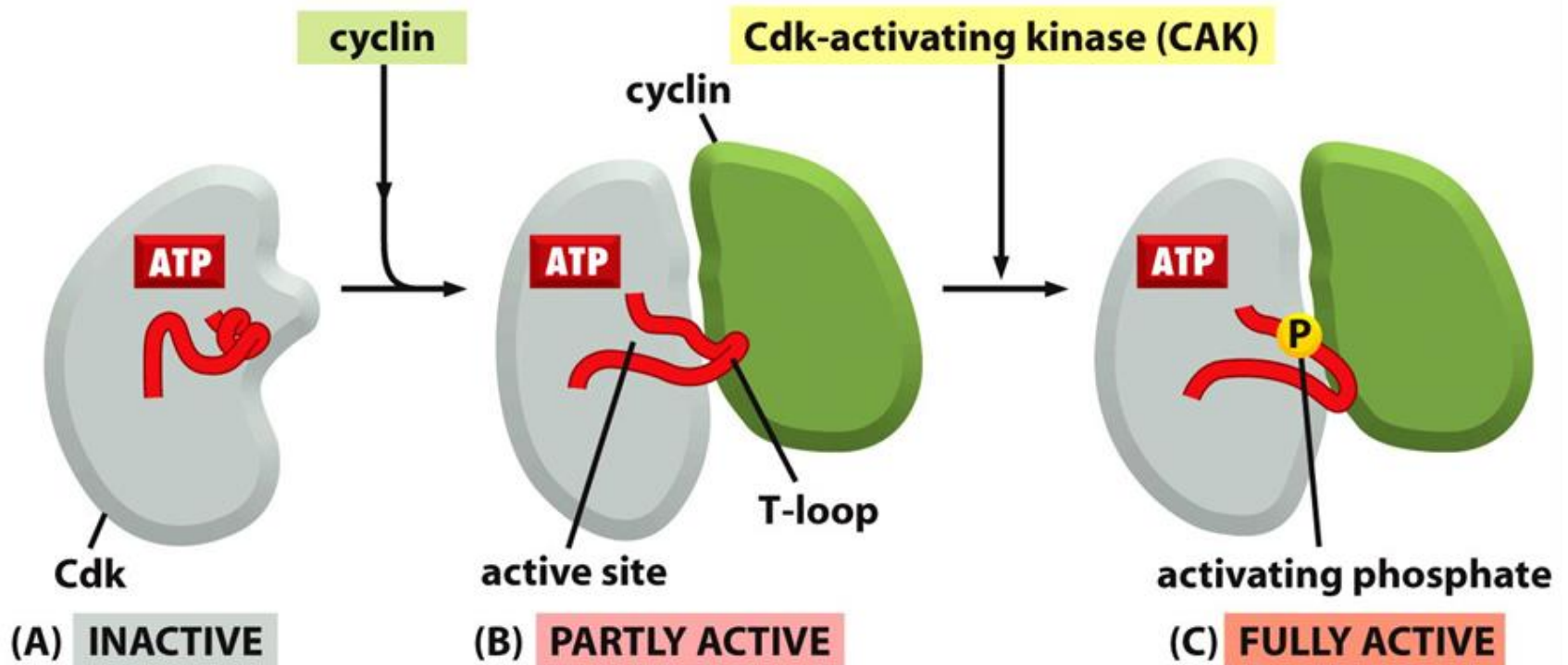


Figure 18-9 Essential Cell Biology 3/e (© Garland Science 2010)

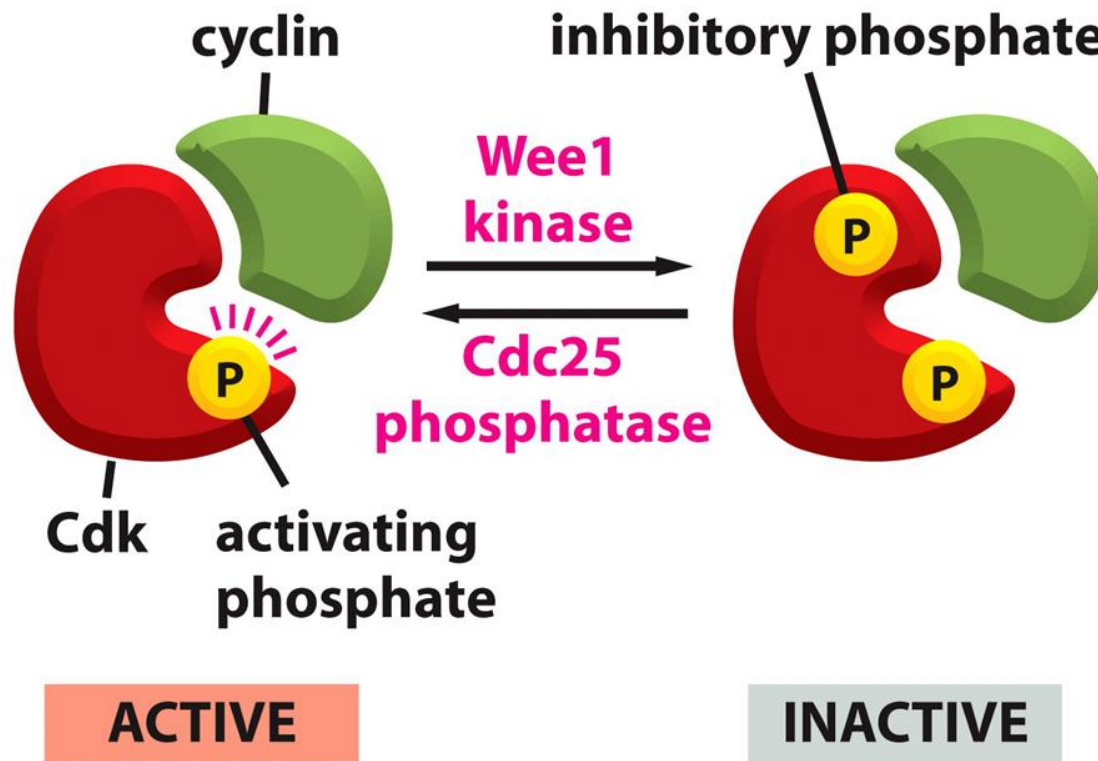
Protein kinases and protein phosphatases that modify Cdks

Cdk-activating kinase (CAK)	phosphorylates an activating site in Cdks
Wee1 kinase	phosphorylates inhibitory sites in Cdks; primarily involved in suppressing Cdk1 activity before mitosis
Cdc25 phosphatase	removes inhibitory phosphates from Cdks; three family members (Cdc25A, B, C) in mammals; primarily involved in controlling Cdk1 activation at the onset of mitosis

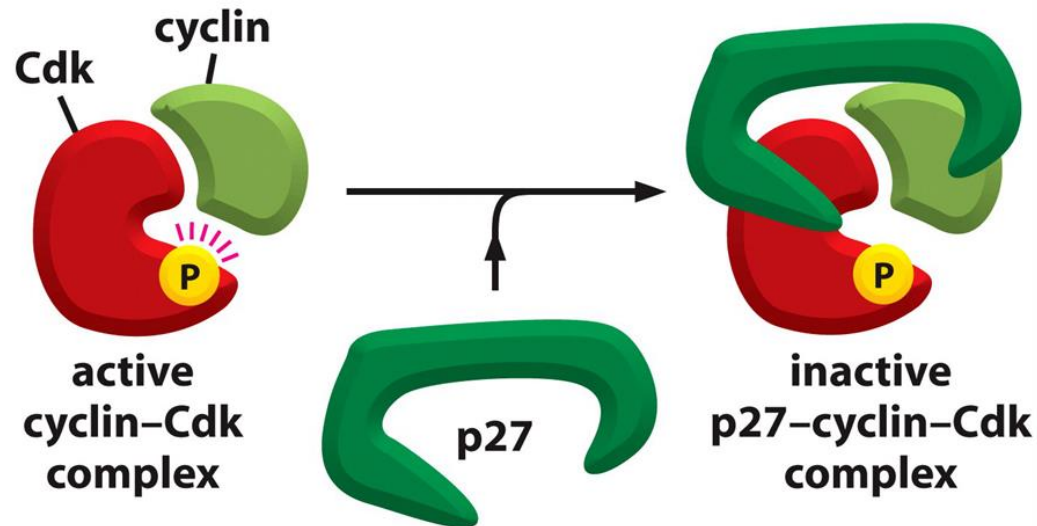
CAK activates the activation loop of Cdk through phosphorylation



Cdk phosphorylation can be regulated by Wee1/Cdc25



Cdk inhibitor proteins (CKI) inhibits Cdk kinase activity



Cdk inhibitor proteins (CKIs)

Sic1 (budding yeast)

p27 (mammals)

p21 (mammals)

p16 (mammals)

suppresses Cdk1 activity in G_1 ; phosphorylation by Cdk1 at the end of G_1 triggers its destruction

suppresses G_1 /S-Cdk and S-Cdk activities in G_1 ; helps cells withdraw from cell cycle when they terminally differentiate; phosphorylation by Cdk2 triggers its ubiquitylation by SCF

suppresses G_1 /S-Cdk and S-Cdk activities following DNA damage

suppresses G_1 -Cdk activity in G_1 ; frequently inactivated in cancer

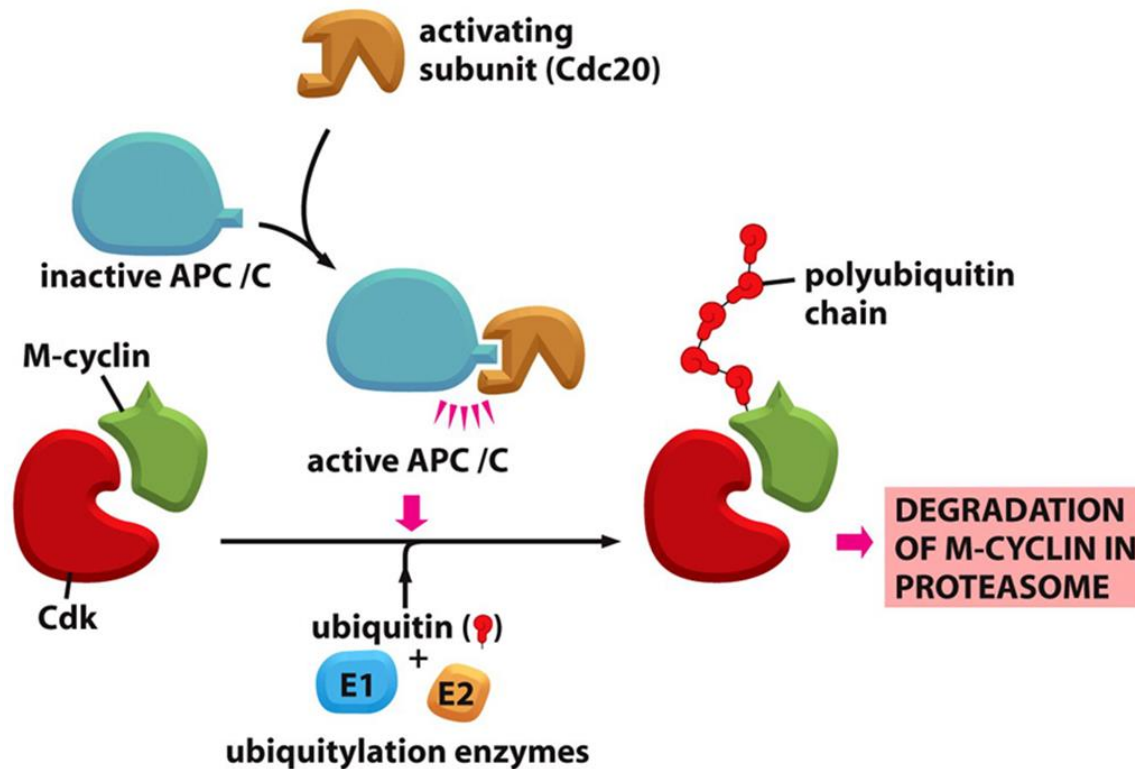
Metaphase to anaphase transition is controlled by proteolysis

- One Key player: anaphase-promoting-complex, or cyclosome (APC/C), a ubiquitin ligase

Reminder: protein degradation mediated by ubiquitination

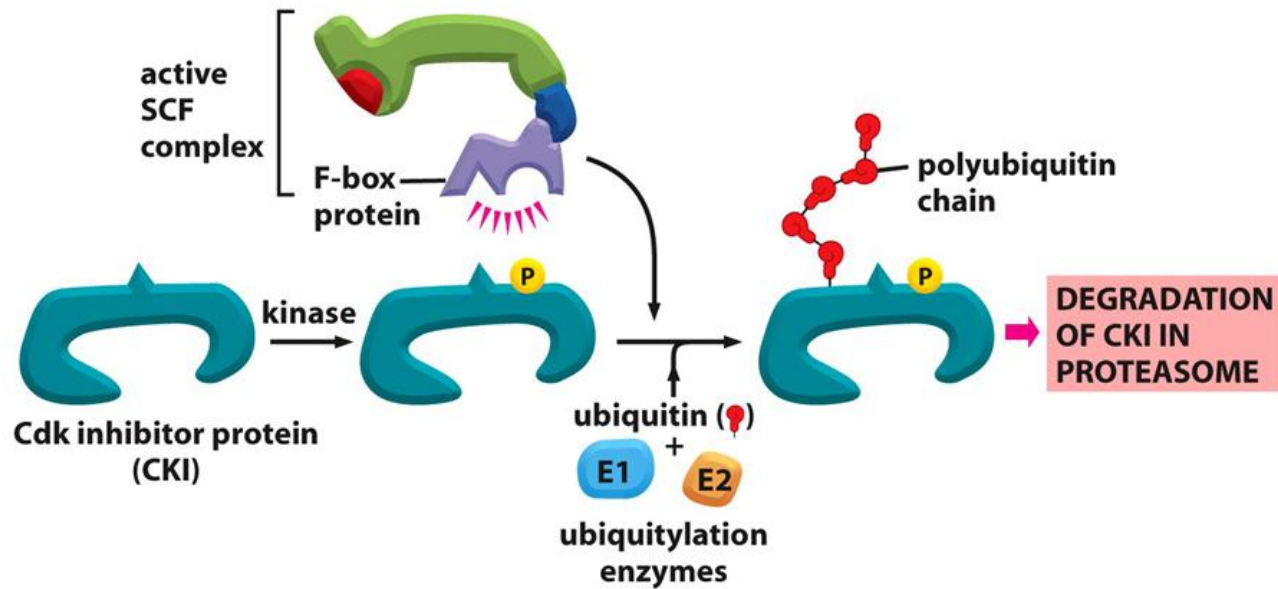
APC/C functions as ubiquitination ligase

control of proteolysis by APC /C

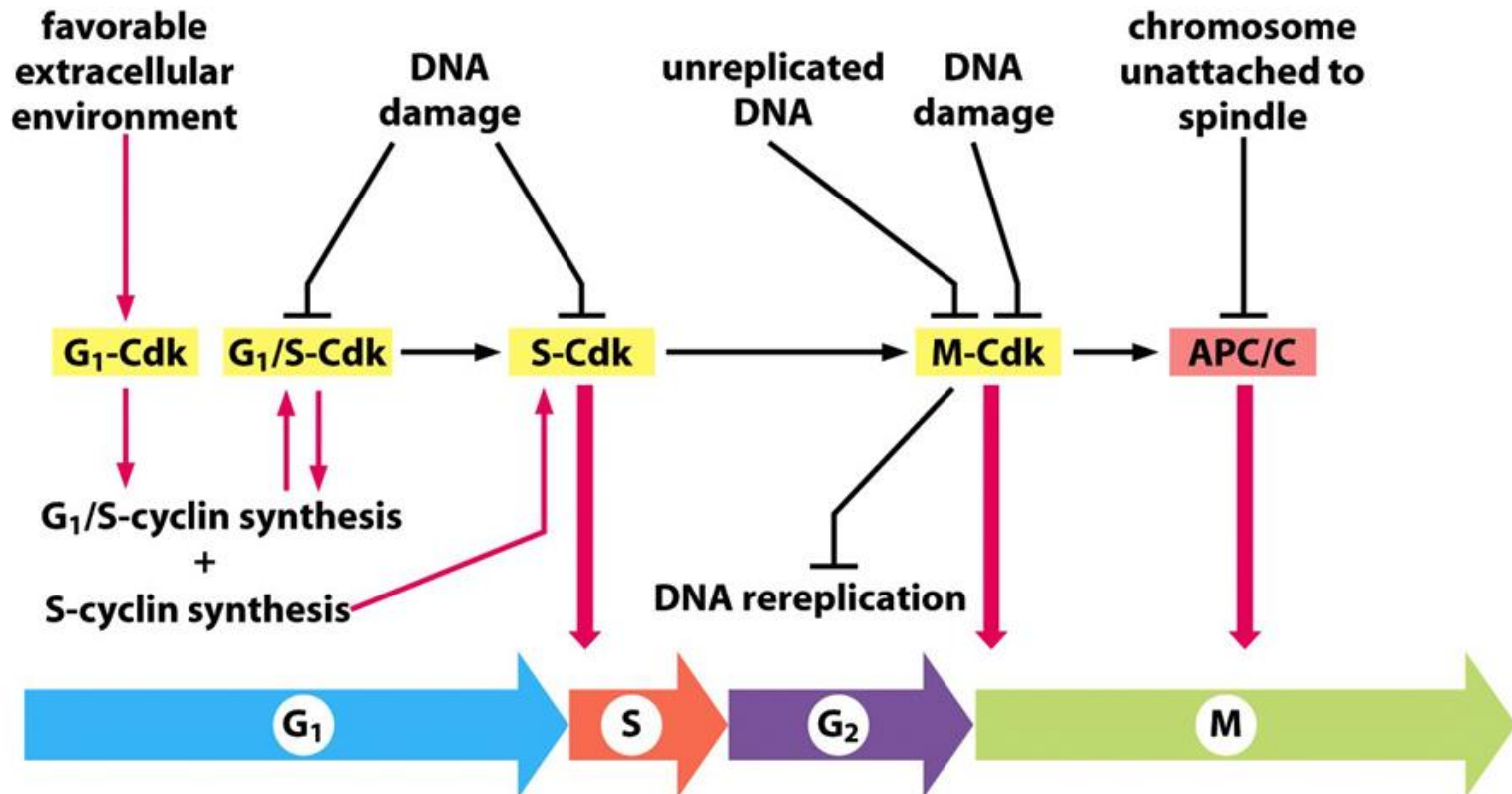


CKI can be degraded by SCF complex

control of proteolysis by SCF



The overview of the cell cycle control system



III. S phase

1. **DNA replication once per cycle**
2. Chromatin protein replication
3. Chromatin structure duplication (histone modification, heterochromatin, euchromatin packaging)
4. Cohesins hold sister chromatids.
5. Centrosome duplication : semiconservative manner, triggered by Cyclin E/cdk2.

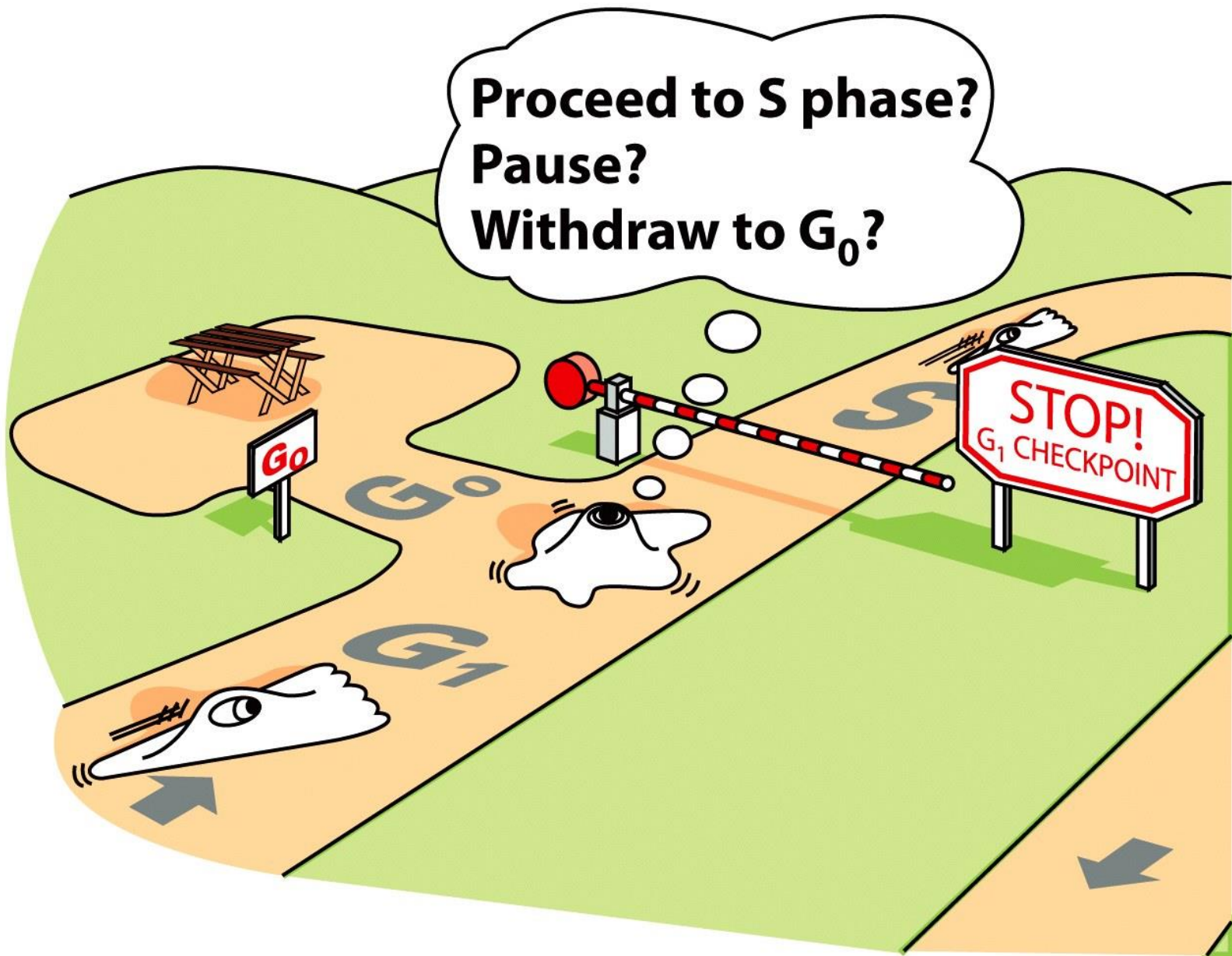
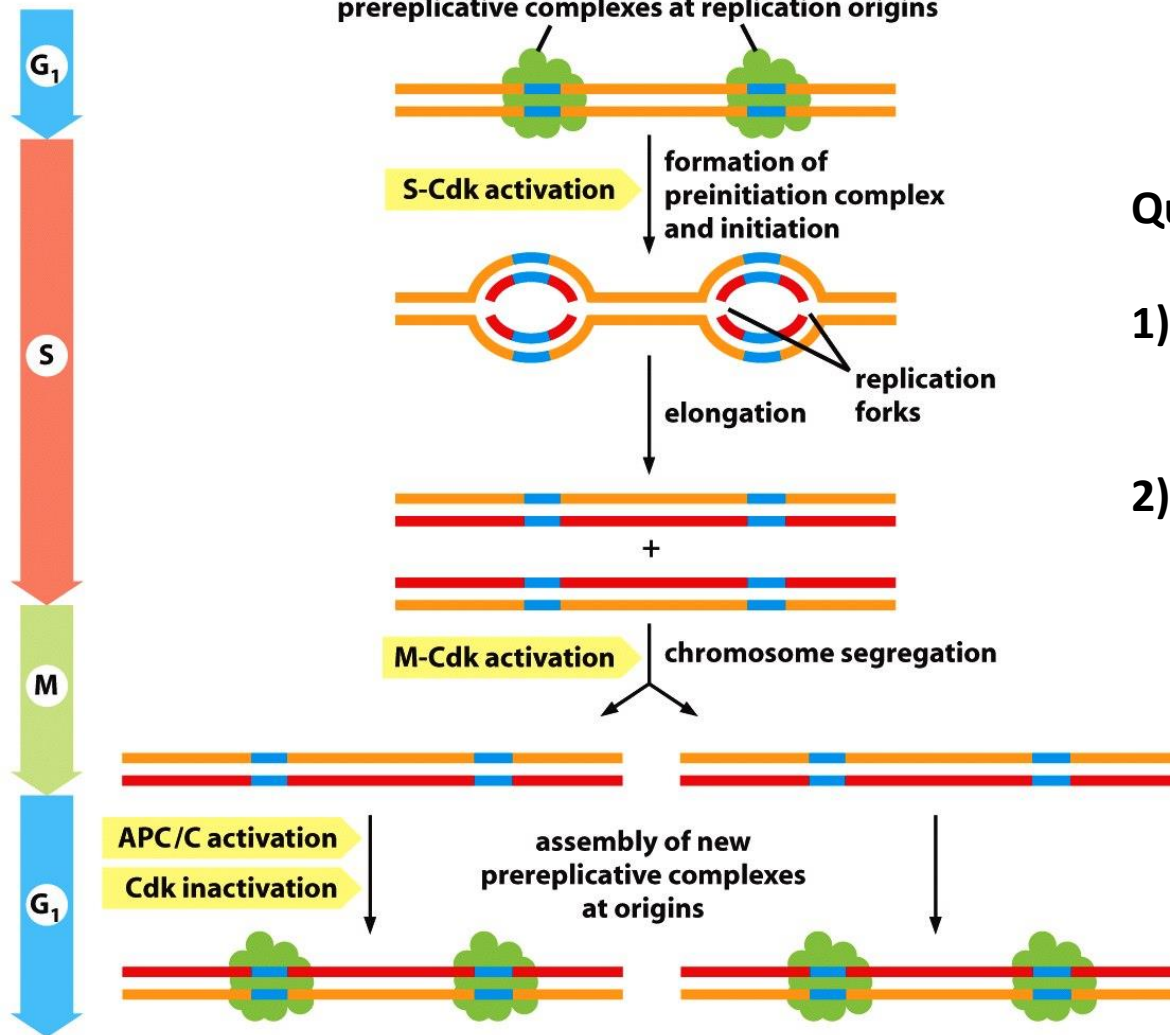


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Chromosome duplication in cell cycle



Questions to be asked:

- 1). How to ensure DNA is replicated only once?
- 2). How do Cdks and APC/C control timed replication?

Question 1. how to ensure DNA replication once per cycle?

Answer

- **Pre-replication complex** (pre-RC) - “prime and licensing”
activated by APC/C in late M and early G1 when APC/C activity is high.
- **Pre-initiation complex** – DNA unwinding, replication
activated by S-Cdk in late G1 when APC/C activity is low, pre-RC is partially dismantled.

S-Cdks and M-Cdks remain high until after late mitosis, when APC/C regains its activity, and start the next round of pre-RC formation.

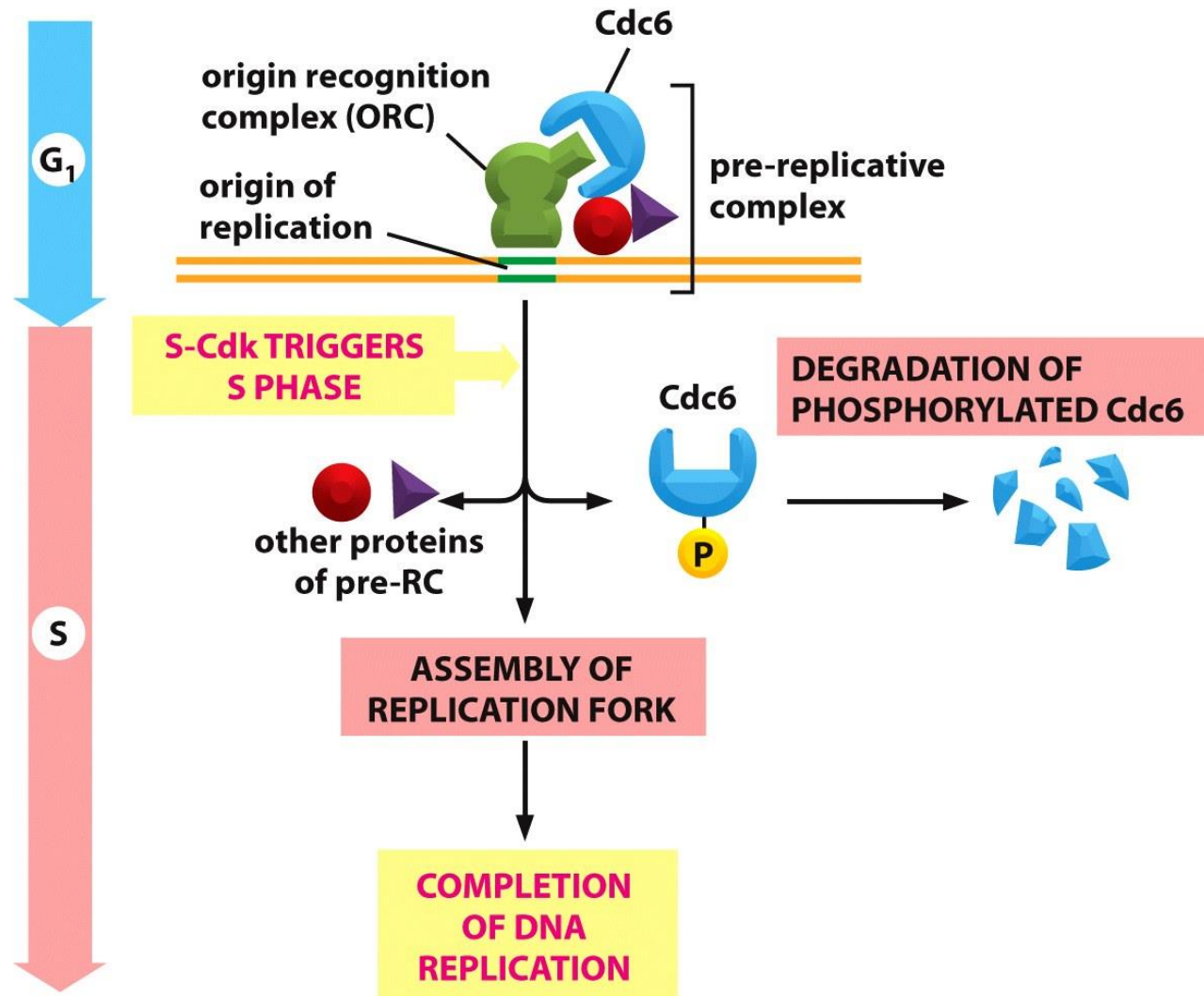
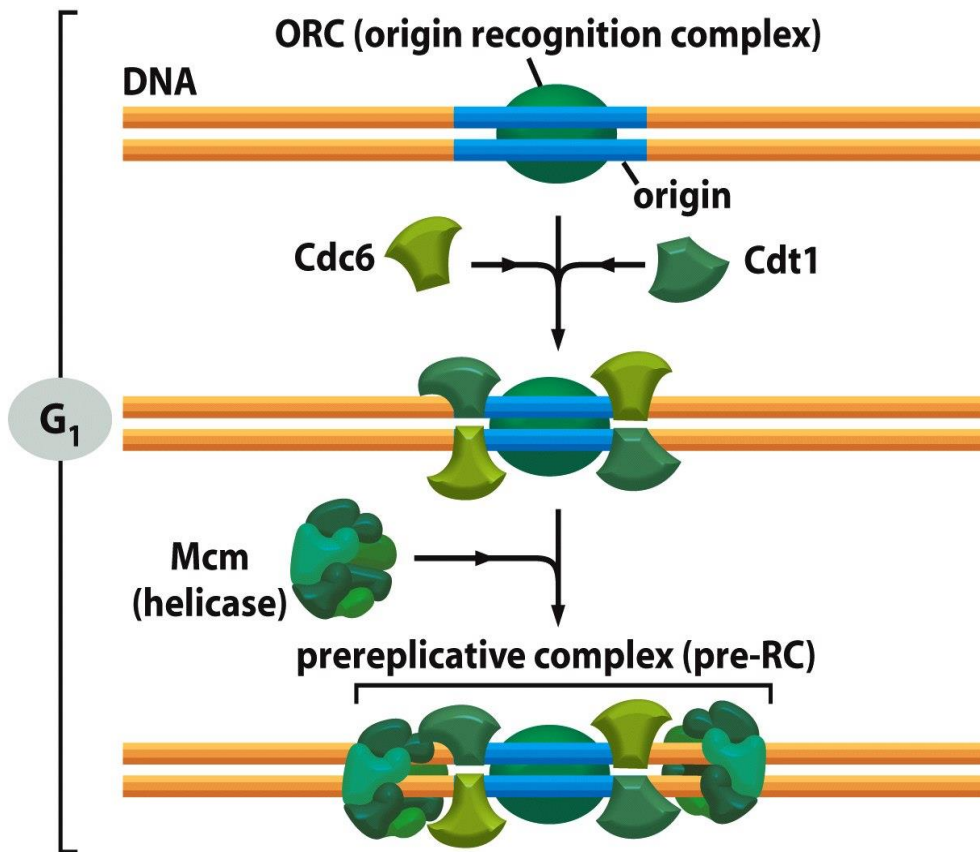


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Question 2: how do Cdks and APC/C control DNA replication in a timed manner?

Stage 1: formation of pre-RC



Pre-RC is inhibited by Cdk
And promoted by APC/C

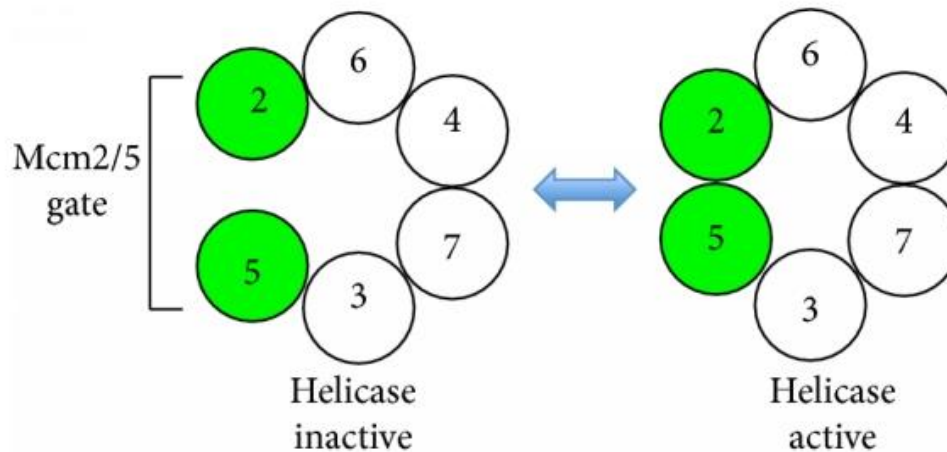
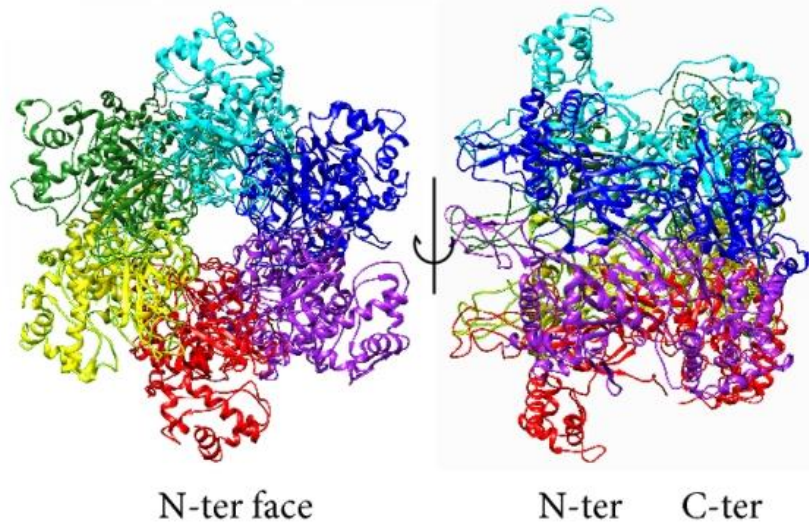
Late M/Early G₁: low Cdk activity,
And high APC/C activity, pre-RC
Is formed.

Firstly, cdc6 and cdt1 associate
with ORC

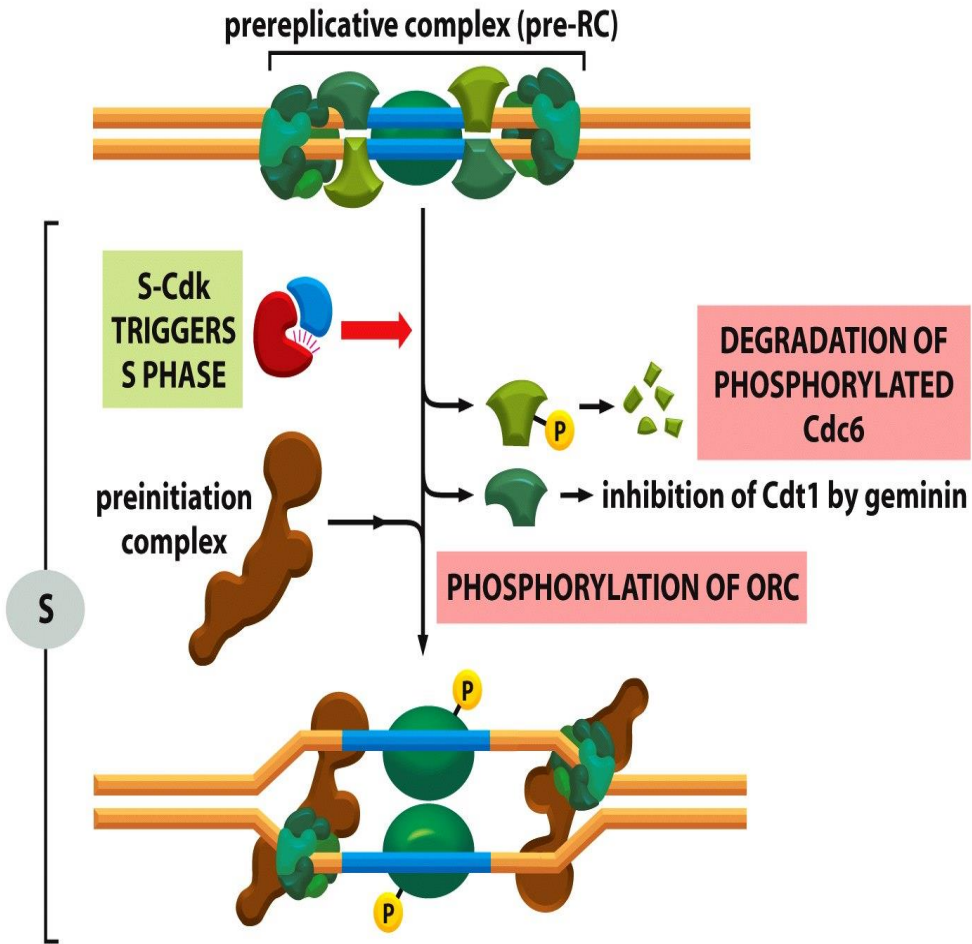
Secondly, MCM ring load onto DNA

Thirdly, pre-RC is formed:
MCM/cdc6/cdt1/ORC

MCM2-7 ring complex



Stage 2: formation of Pre-initiation complex



Firstly, S-Cdk Trigger inactivation Of Cdc6 AND Cdt1, dismantles pre-RC

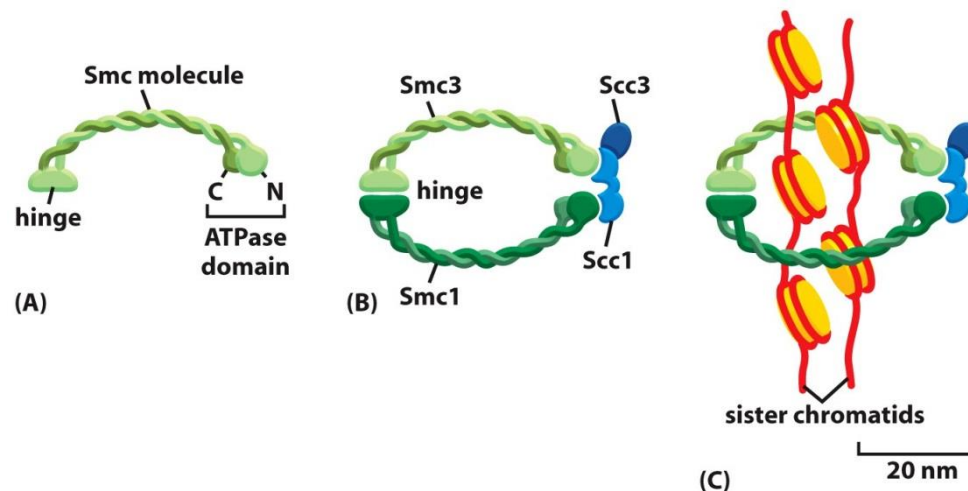
Secondly, APC/C triggers degradation of geminin, but when S-Cdk is high and APC/C is low in activity, geminin is stabilized.

Thirdly, S-Cdk phosphorylates ORC And load preinitiation complex onto ORC to initiate DNA replication.

Cohesins and DNA catenation help hold sister chromatids together

- ♥ Cohesins are deposited at many locations along the length of each sister chromatids
- ♥ Prevent drifting apart for sister chromatids after DNA replication.
- ♥ DNA catenation is interwining of sister DNA molecules which can be resolved by DNA topoisomerase II

Cohesins



Centriole replication happens during S phase,

