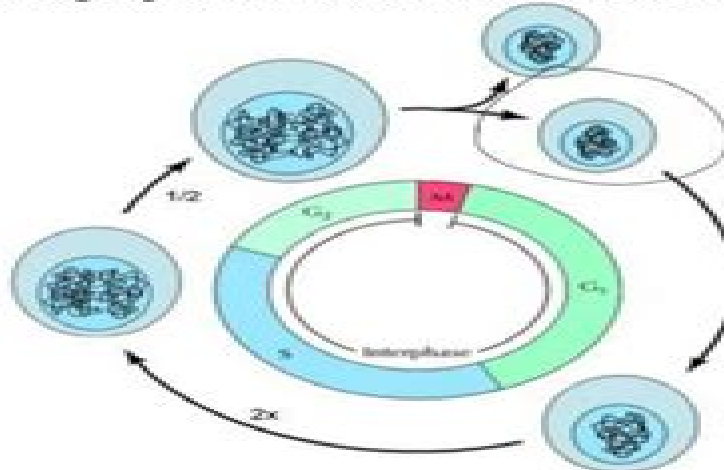


**The Cell Cycle – How One Cell Becomes Two Cells**

This figure gives more information about how one cell becomes two daughter cells.



Interphase includes G<sub>1</sub>, S and G<sub>2</sub>. Throughout Interphase the cell makes more cytoplasm and grows larger.

During the S phase the DNA in each chromosome is replicated to produce two copies of each chromosome. Each copy has the same genes as the original chromosome.

The M phase includes mitosis and cytokinesis. During mitosis, the two copies of each chromosome are separated to opposite ends of the cell. During cytokinesis the cell divides into two daughter cells, each with a complete set of chromosomes.

4. The cell cycle begins with a single cell and produces two daughter cells; each daughter cell can repeat the cell cycle. Circle the cell in the figure that represents both a daughter cell that has been produced by the cell cycle and a cell that is beginning the cell cycle. Explain why the cell is smallest at this time in the cell cycle.

**The cell is the smallest at this time because it has not fully grown and developed through the cell cycle.**

5a. Use 2x to label the arrow(s) when the amount of DNA doubles.

5b. Use 1/2 to label the arrow(s) when the amount of DNA in the cell is halved.

6. Complete this table to explain why specific phases of the cell cycle are required for successful cell division.

What would go wrong if the cell cycle occurred without the S phase?	Without the S phase, the daughter cells would have half of the amount of chromosomes.
What would go wrong if the cell cycle occurred without mitosis?	If M fase did not occur, then cells would not be very strange as they may not be identical to the parent cell. This is because in this stage the copple of the cell is one side of the cell, and if these mixed while dividing then the cells would not be exact clones.

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Leong, 2003 The rapid acceptance of immunohistochemistry as an important and even indispensable adjunct to morphological examination and diagnosis requires the modern anatomical pathology laboratory to be conversant with and proficient in immunostaining procedures as well as methods of tissue processing and antigen retrieval and the underlying characteristics of the increasing number of antibodies and antisera available. This fully revised expanded and updated edition provides a comprehensive list of antisera and monoclonal antibodies that have useful diagnostic applications in tissue sections and cell preparations. Various clones which are commercially available to detect the same antigen are listed and the sensitivities and specificities of the antibodies are discussed. Importantly the authors all pathologists with a wealth of experience in immunostaining procedures provide fully referenced details and expert advice on each reagent with the reader will find invaluable. Finally the appendices provide easily accessible and clear summaries of selected antibody panels for specific diagnostic situations details of heat induced antigen epitope retrieval including the use of microwaves and a useful reference to the websites of the main antibody suppliers.

**Leong's Manual of Diagnostic Biomarkers for Immunohistology**

Runjan Chetty, Kumarasen Cooper, Carol Cheung, Srinivas Mandavilli, 2022-07-21 With the interest in biomarkers available for diagnostic therapeutic prognostic and research applications growing rapidly it is now expected that every pathologist should be aware of the correct application of immunohistochemistry in daily practice. This fully updated fourth edition provides practical guidance about preparation key uses and pitfalls of antibodies for immunohistology in a unique A-Z format.

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*Readers' Guide to Periodical Literature*, 1922 An author subject index to selected general interest periodicals of reference value in libraries.

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pathologists as well as researchers residents and trainees     *Student Handbook and Solutions Manual* Harry Nickla,Klug,2002-08 Harry Nickla Creighton University This valuable handbook provides detailed step by step solutions or extensive explanations for every problem in the text Additional study aids include extra study problems chapter outlines vocabulary exercises and an overview of how to study genetics     **Techniques in Cell Cycle Analysis** Joe W. Gray,Zbigniew Darzynkiewicz,2008-02-24 Quantification of the proliferative characteristics of normal and malignant cells has been of interest to oncologists and cancer biologists for almost three decades This interest stems from the fact that cancer is a disease of uncontrolled proliferation b the finding that many of the commonly used anticancer agents are preferentially toxic to cells that are actively proliferating and c the observation that significant differences in proliferation characteristics exist between normal and malignant cells Initially cell cycle analysis was pursued enthusiastically in the hope of generating information useful for the development of rational cancer therapy strategies for example by allowing identification of rapidly proliferating tumors against which cell cycle specific agents could be used with maximum effectiveness and by allowing rational scheduling of cell cycle specific therapeutic agents to maximize the therapeutic ratio Unfortunately several difficulties have prevented realization of the early promise of cell cycle analysis Proliferative patterns of the normal and malignant tissues have been found to be substantially more complex than originally anticipated and synchronization of human tumors has proved remarkably difficult Human tumors of the same type have proved highly variable and the cytokinetic tools available for cell cycle analysis have been labor intensive as well as somewhat subjective and in many cases inapplicable to humans However the potential for substantially improved cancer therapy remains if more accurate cytokinetic information about human malignancies and normal tissues can be obtained in a timely fashion     **Reader's Guide to Periodical Literature Supplement** ,1917 These vols contain the same material as the early vols of Social sciences humanities index

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