



BHARATHIDASAN UNIVERSITY

Tiruchirappalli- 620024, Tamil Nadu,
India

**Programme: M.Sc., Biomedical Science
(5 Year Integrated Program)**

Course Title : Stem Cell Biology and Tissue engineering
Course Code : 18BMS48C14

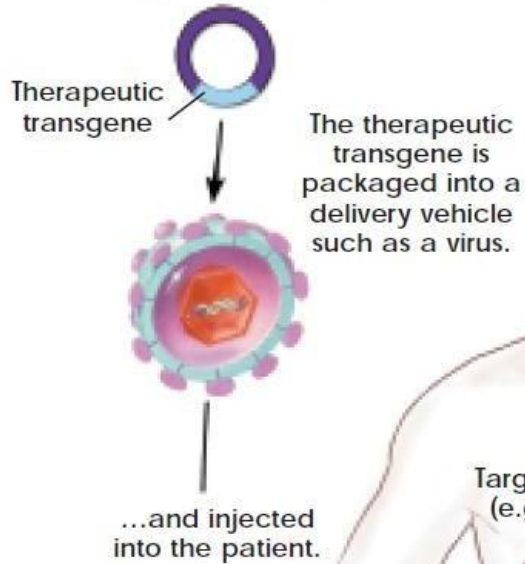
Unit-IV
Stem Cells in Gene Therapies

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STEM CELLS IN EXPERIMENTAL GENE THERAPIES

**WHY STEM CELLS ARE USED IN SOME
CELL-BASED GENE THERAPIES?**

DIRECT DELIVERY



CELL-BASED DELIVERY

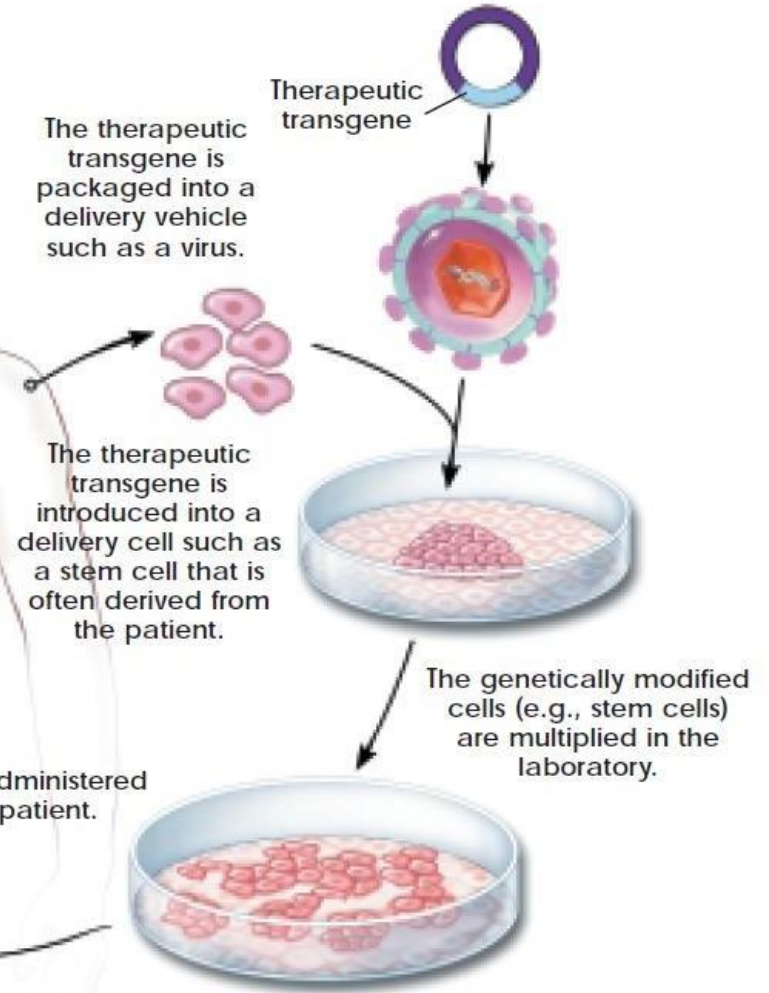


Figure 11.1. Strategies for Delivering Therapeutic Transgenes into Patients.

First, although small in number, they are **readily removed from the body via the circulating blood or bone marrow** of adults or the **umbilical cord blood** of newborn infants.

In addition, they are **easily identified and manipulated** in the laboratory and can be returned to patients relatively easily by injection.

The ability of hematopoietic stem cells to give rise to many different types of blood cells means that once the engineered stem cells differentiate, **the therapeutic transgene will reside** in cells such as T and B lymphocytes, natural killer cells, monocytes, macrophages, granulocytes, eosinophils, basophils, and megakaryocytes.

The clinical applications of hematopoietic stem cell-based gene therapies are thus also diverse, extending to **organ transplantation, blood and bone marrow disorders, and immune system disorders.**

In addition, hematopoietic stem cells “**home,**” or **migrate,** to a number of different spots in the body—primarily the bone marrow, but also the liver, spleen, and lymph nodes. These may be strategic locations for localized delivery of therapeutic agents for disorders unrelated to the blood system, such as liver diseases and metabolic disorders.

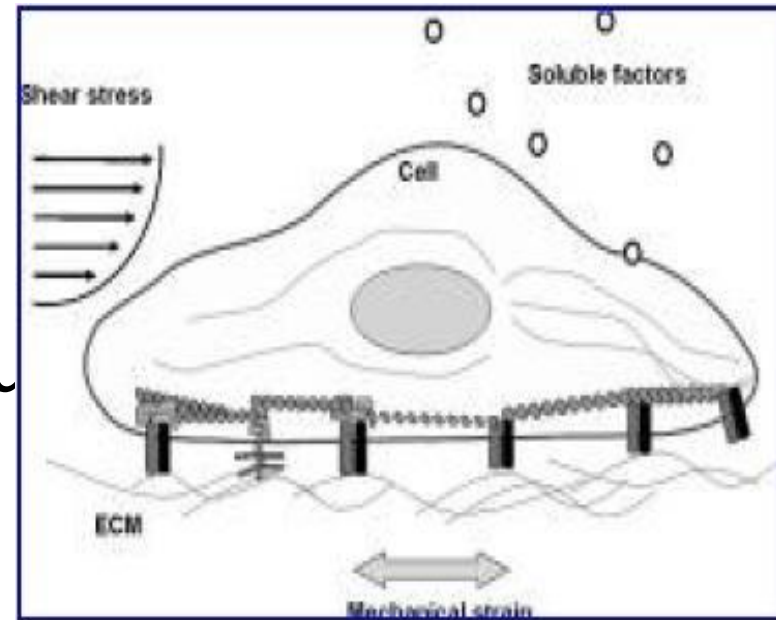
Myoblasts appear to be good candidates for use in gene therapy because of an unusual and advantageous biological property: when injected into muscle, they fuse with nearby muscle fibers and become an integral part of the muscle tissue.

In a series of experiments in rodents, a team of investigators has been testing **neural stem cells** as vehicles for cell-based gene therapy for brain tumors known as **gliomas**.

Another cell-based gene therapy system under investigation involves the use of **osteoblasts**, or bone forming stem cells.

Tissue Engineering

- Repair/replace damaged tissue
 - Enhance natural regeneration



Cell Source

Embryonic stem cells
Adult stem cells
Progenitor cells

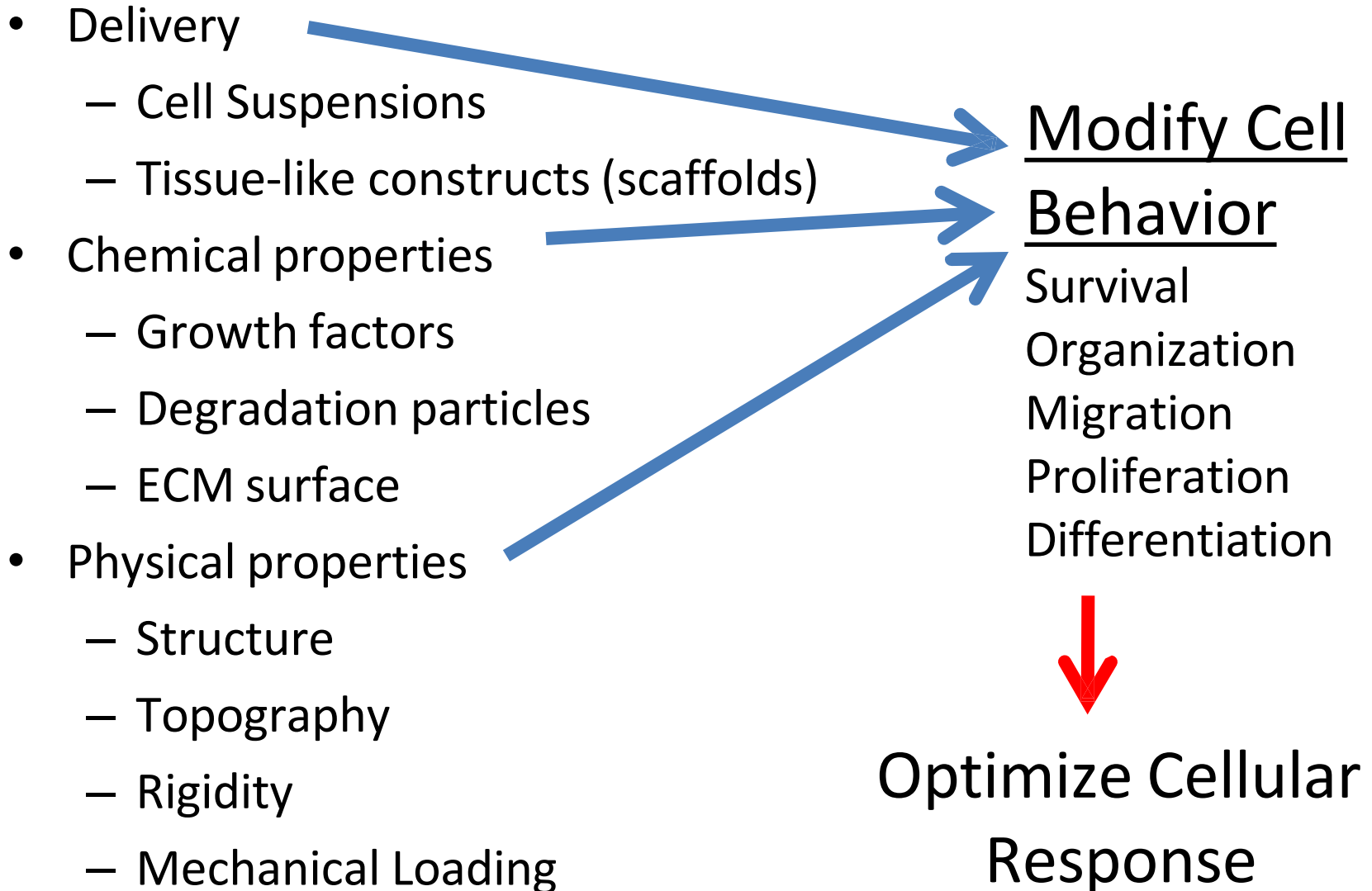
Signals

Growth factors
Drugs
Mechanical forces

ECM

Metals
Ceramics
Synthetic polymers
Natural polymers

Important Variables



Models for Tissue Engineering

- *In vitro* differentiation
 - Construct tissues outside body before transplantation
 - Ultimate goal
 - Most economical
 - Least waiting time
- *In situ* methodology
 - Host remodeling of environment
- *Ex vivo* approach
 - Excision and remodeling in culture

Combine physical and
chemical factors



Optimize stem cell
differentiation and
organization

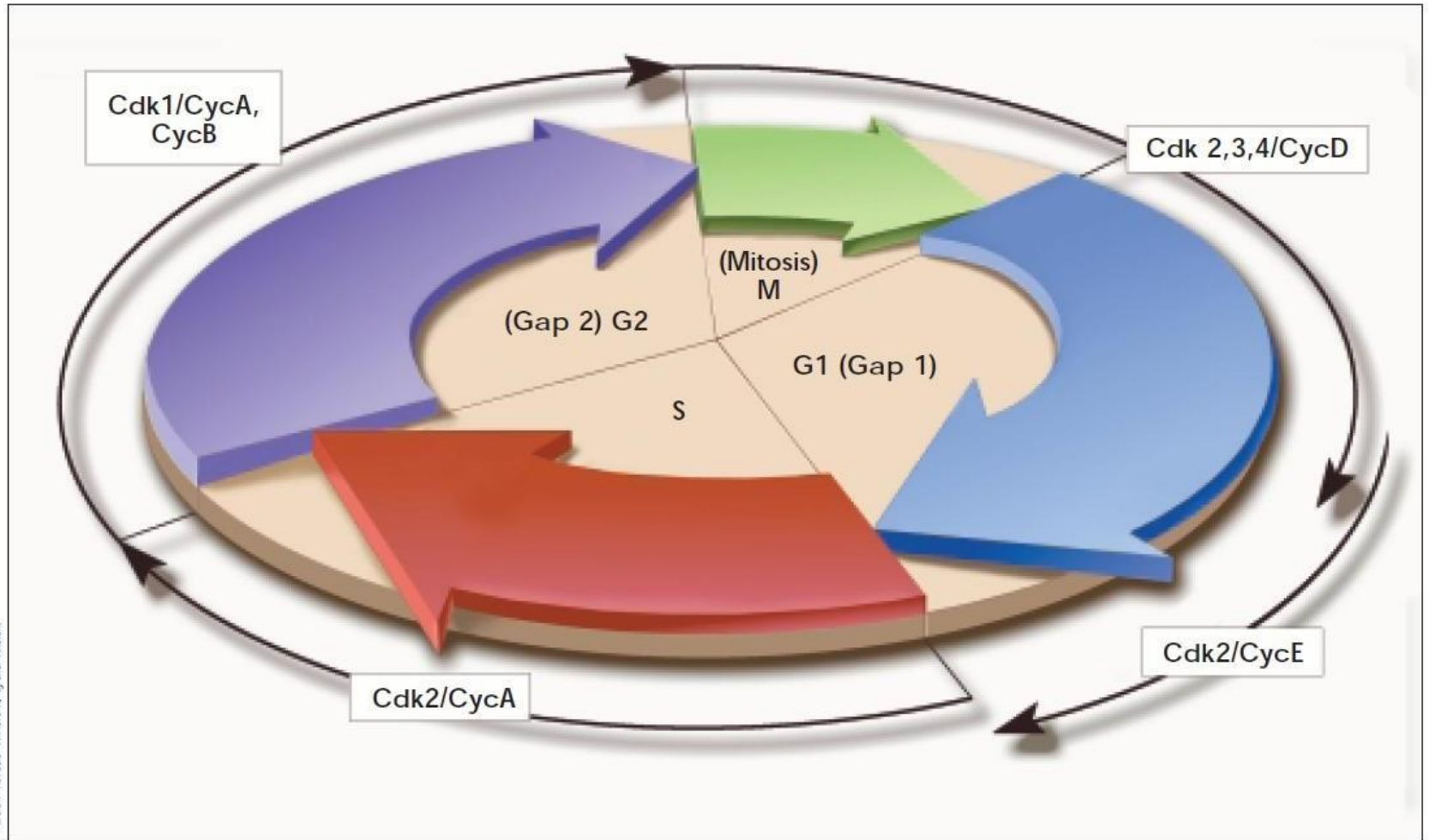
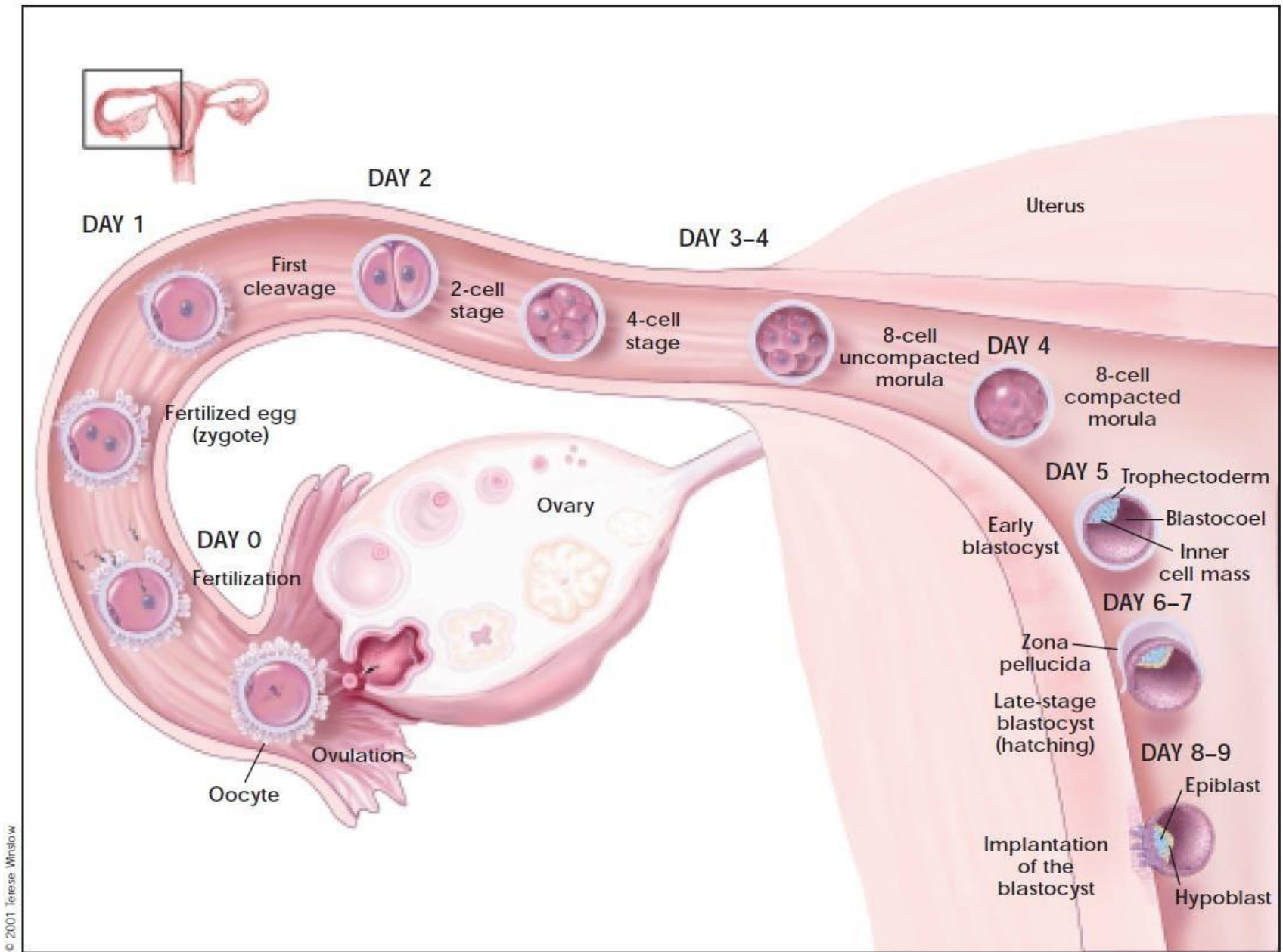


Figure A.1. Cell Cycle.



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Figure A.2. Development of the Preimplantation Blastocyst in Humans.

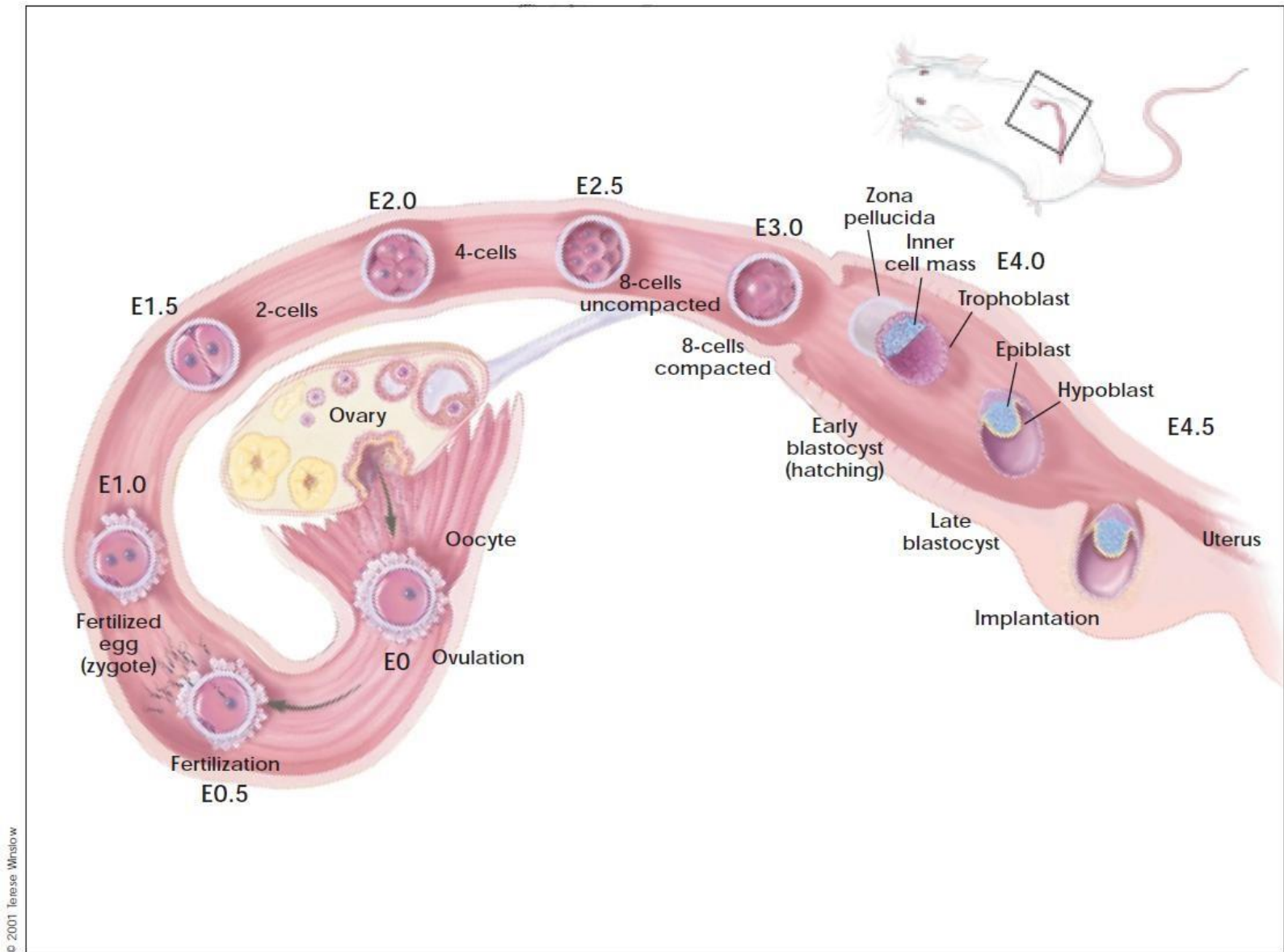


Figure A.3. Development of the Preimplantation Blastocyst in Mice from Embryonic Day 0 (E0) Through Day 5 (E5.0).

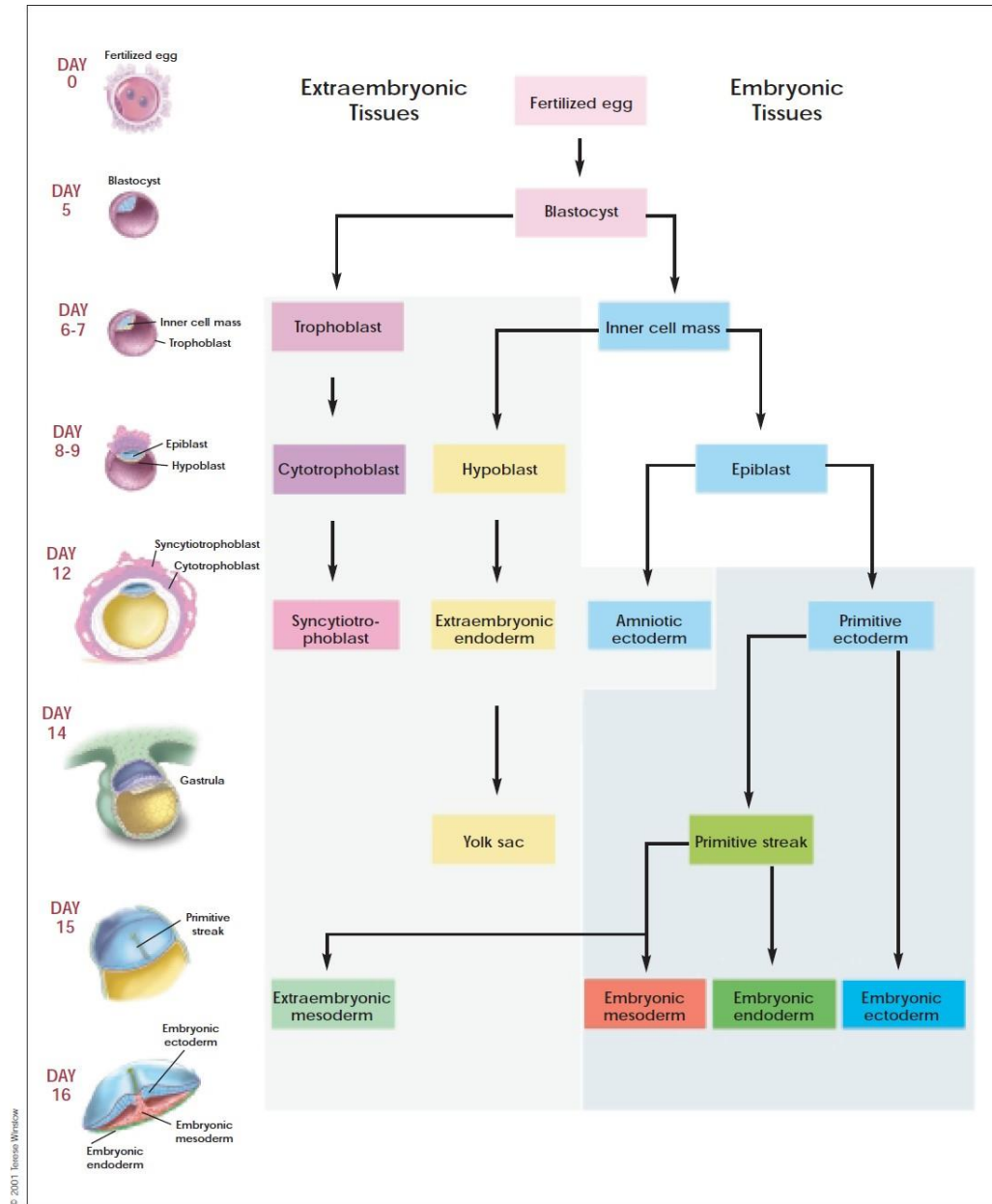


Figure A.4. Development of Human Embryonic Tissues.

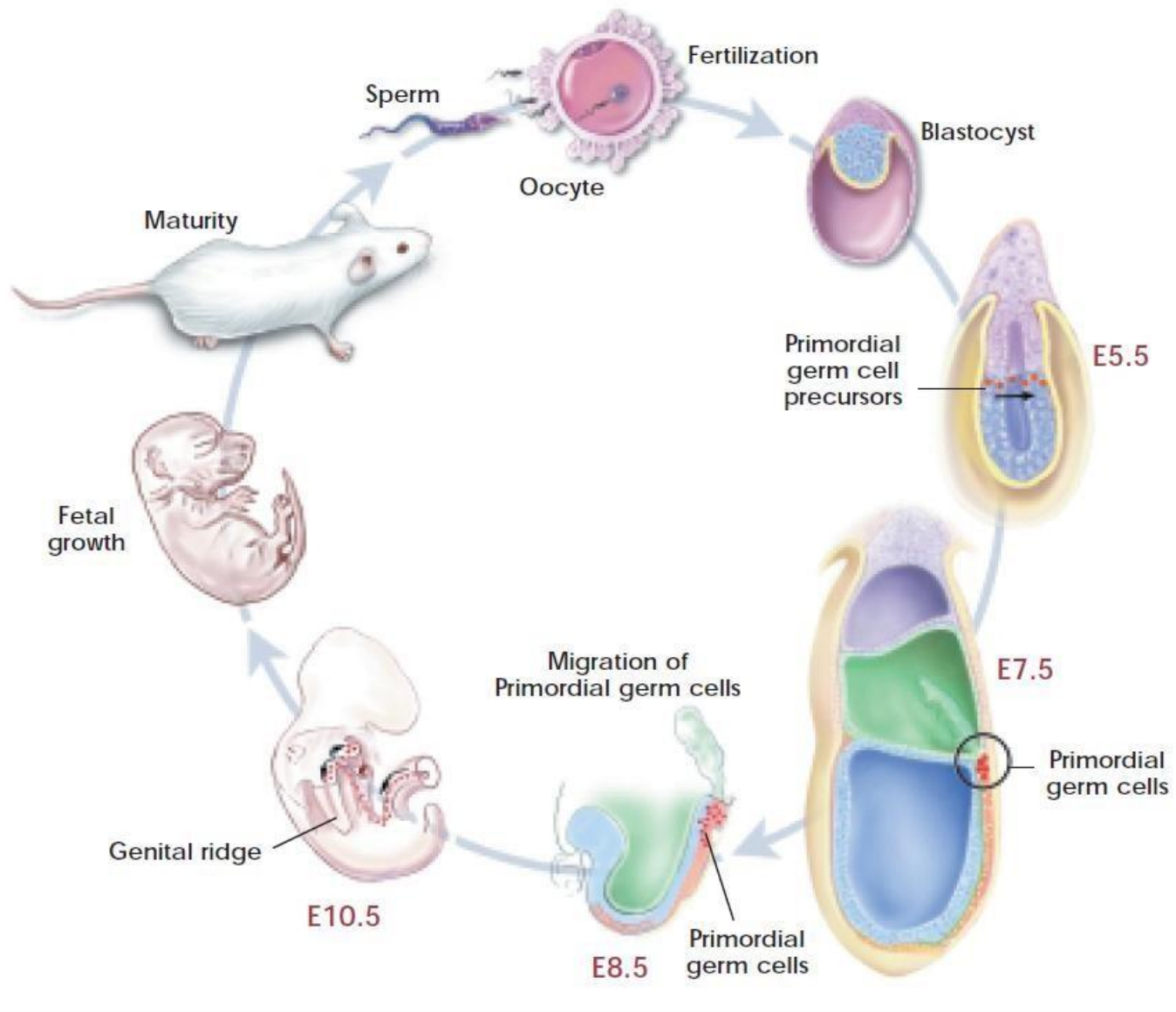
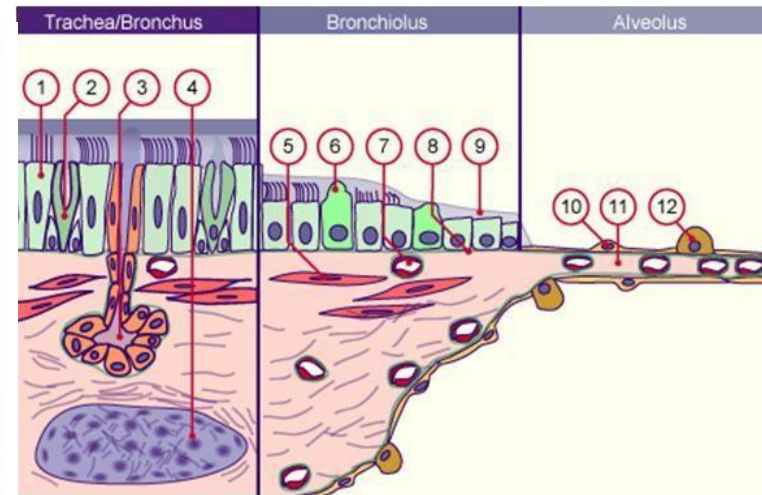
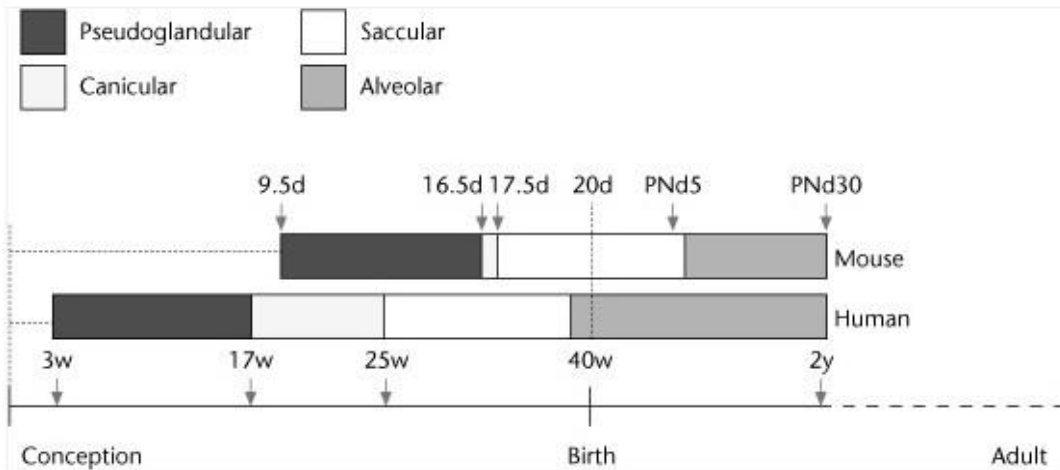
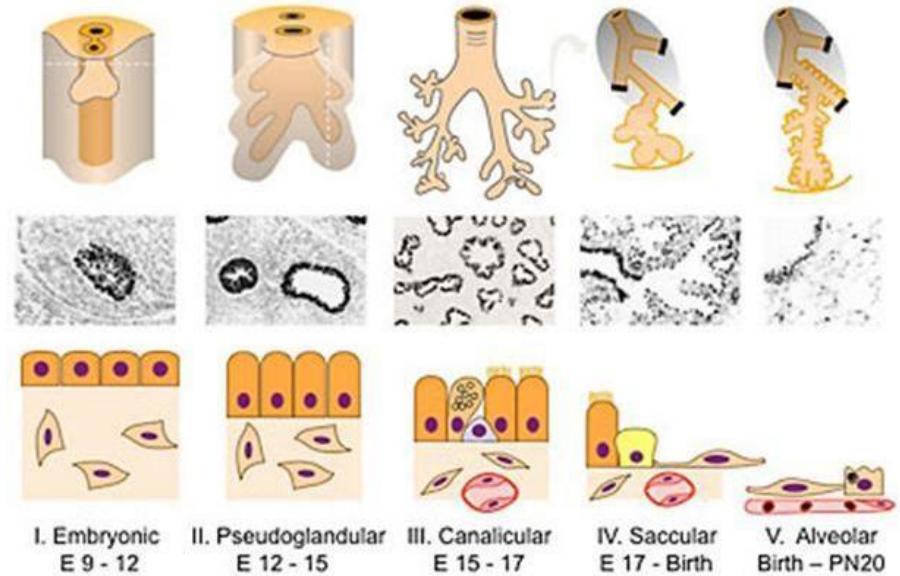
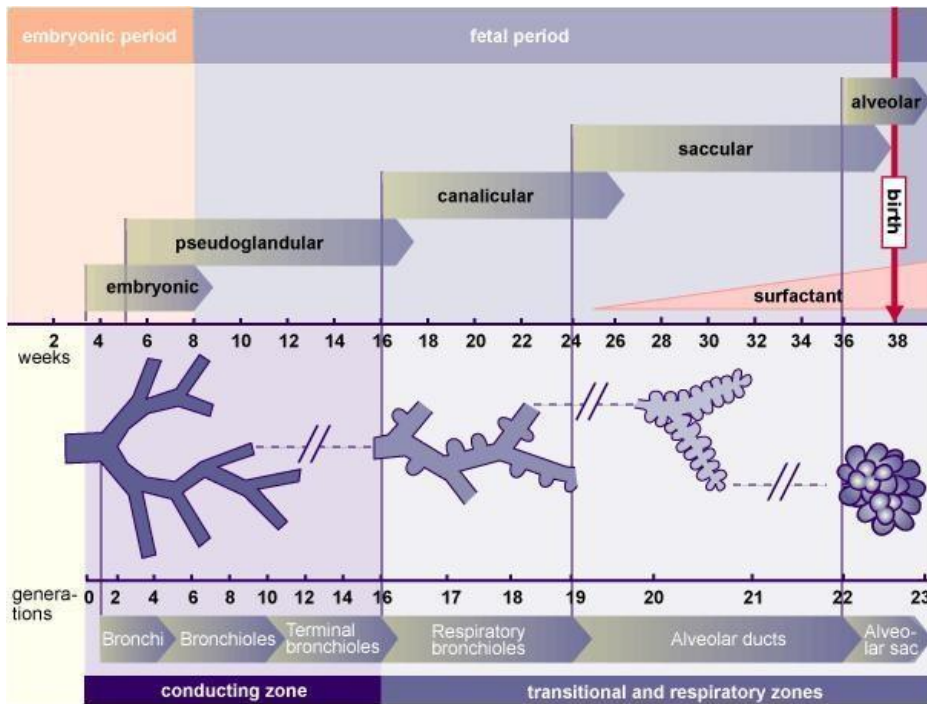


Figure A.5. Development of Mouse Embryonic Primordial Germ Cells.



THANK YOU