Transformation and Oncogenesis

Lecture 18
Biology 3310/4310
Virology
Spring 2017

Cause and effect, means and ends, seed and fruit, cannot be severed; for the effect already blooms in the cause, the end pre-exists in the means, the fruit in the seed.
RALPH WALDO EMERSON
Transformation

Hamster embryo → Trypsin → Single cells → Chemical → Seed plates → Transformed clones identified morphologically

Normal
Transformed

Loss of growth potential → Appearance of immortal cells

Cell growth rate

Days after initiation of culture

30 60

Crisis
The puzzling properties of transformed cells in the laboratory

- Immortal: Grow indefinitely (HeLa)
- Loss of anchorage dependence
- Loss of contact inhibition
- Colony formation in semi-solid media
- Decreased requirements for growth factors (serum)
Oncogenesis

- Development of cancer
  - Tumor: swelling caused by abnormal growth of tissue, benign or malignant
- Cancer is a genetic disease
- 8.2 million deaths/yr developed countries
- Mutations (~12) affect signal transduction pathways that govern cell proliferation, survival, determination of cell fate, maintenance of genome integrity
- Mutations may be inherited, caused by DNA damage, environmental carcinogens, infectious agents including viruses
Transformation and oncogenesis are distinct

- Studying virus-transformed cells provides insight into molecular events that establish oncogenic potential
- *No virus can do it all*
# Human cancer viruses

<table>
<thead>
<tr>
<th>Virus</th>
<th>Cancer</th>
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<tbody>
<tr>
<td>Epstein-Barr virus</td>
<td>Burkitt's lymphoma</td>
</tr>
<tr>
<td></td>
<td>Nasopharyngeal carcinoma</td>
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<tr>
<td>Hepatitis B virus</td>
<td>Hepatocellular carcinoma</td>
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<tr>
<td>Hepatitis C virus</td>
<td>Hepatocellular carcinoma</td>
</tr>
<tr>
<td>Human T-lymphotropic virus-1</td>
<td>Adult T cell leukemia</td>
</tr>
<tr>
<td>Human immunodeficiency virus-1</td>
<td>Many tissues and organs</td>
</tr>
<tr>
<td>Human papillomavirus</td>
<td>Cervical, penile, anogenital, head and neck cancers</td>
</tr>
<tr>
<td>Kaposi’s sarcoma herpesvirus</td>
<td>Kaposi’s sarcoma</td>
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<tr>
<td></td>
<td>Primary effusion lymphoma</td>
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<tr>
<td></td>
<td>Multicentric Castleman’s disease</td>
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<tr>
<td>Merkel cell polyomavirus</td>
<td>Merkel cell carcinoma</td>
</tr>
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</table>

Contributing factor in ~20% of human cancers
Virus-induced cancer

*Transformation and oncogenesis is not required for replication of any* virus
On October 1, 1909, Dr. Peyton Rous removed a tumor from an English hen and injected a cell-free filtrate from the tumor into another healthy chicken, which later developed the same type of tumor.

*Cancer could be caused by a viral infection!*
It took the world almost 50 years to accept this idea

Dr. Rous lived long enough to be awarded the Nobel Prize for Physiology and Medicine in 1966 for his research

His legacy: RSV; Rous Sarcoma Virus, a key player in two more Nobel Prizes
“By the 1950s, cancer researchers had split into three feuding camps.

The virologists, lead by Rous, claimed that viruses caused cancer, although no such virus had been found in human studies.

Epidemiologists...argued that exogenous chemicals caused cancer, although they could not offer a mechanistic explanation.

The third camp possessed weak, circumstantial evidence that genes internal to the cell might cause cancer...
In 1951, a young virologist named Howard Temin arrived at Cal Tech to study the genetics of fruit flies. Restless and imaginative, he soon grew bored with fruit flies. Switching fields, he chose to study Rous sarcoma virus in Renato Dulbecco’s laboratory.

Until the late fifties, Rous sarcoma virus had been shown to cause tumors only in live chickens. Temin imagined creating cancer in a petri dish. In 1958, in his seventh year in Dulbecco’s lab, Temin succeeded.
He added Rous sarcoma virus to a layer of normal cells in a petri dish. The infection of the cells incited them to grow uncontrollably, forcing them to form tiny distorted heaps containing hundreds of cells that Temin called foci. The foci, Temin reasoned, represented cancer distilled into its essential, elemental form: cells growing uncontrollably, unstoppably - pathological mitosis.

_Temin went on to discover RT in RSV_

**Avian cells transformed by RSV**

A  
B  
C  
D  

- Fusiform
- Round, refractile
Transformation of cells by viruses

- 1962: After infection with polyomavirus, rare BHK21 cells changed shape, kept growing.
- 1964: After infection of Swiss 3T3 cells with SV40, rare cells grew as colonies. Most of the infected cells died, but rare cells did not. They were “transformed”.
How can a viral infection transform a cell?

- Cytopathic effects must be reduced or eliminated
  - *The infected cell does not die*

- Viral replication must be reduced or eliminated
  - *Transformed cells do not produce virions*

- The cell must continue to divide
  - *It becomes immortal*
Which of the following is not a property of transformed cells?

A. Increased requirements for growth factors
B. Immortality
C. Loss of anchorage dependence
D. Loss of contact inhibition
E. Colony formation in semi-solid media
Route to understanding viral transformation of cells in culture and relationship to cancer was convoluted.
How does Rous sarcoma virus cause tumors in chickens and transform cells \textit{in vitro}?

<table>
<thead>
<tr>
<th>(1900s) Retroviruses</th>
<th>(1943) in vitro cancer biology</th>
<th>(1920s) DNA viruses</th>
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<tbody>
<tr>
<td>(1950’s) in vitro studies with RSV</td>
<td>Convergence (1960s, 1970s)</td>
<td>(1959) in vitro studies with polyomaviruses</td>
</tr>
</tbody>
</table>

\textit{(present) Unified theory of cell growth control}
Avian leucosis retroviruses (ALV) are endemic in virtually all chicken flocks

- Ellerman & Bang 1908
- Most chickens infected with ALV within a few months of hatching
- Leucosis (leukemia) occurs sporadically in infected birds >14 wk old (3%)
- 97% of birds have transient viremia, become immune, don’t get leukemia
Infected birds develop other cancers as they age

- Connective tissue tumors or sarcomas (solid tumors)
- Viruses isolated from these solid tumors rapidly cause sarcomas, not leucosis
- Rous isolated one of these viruses: Rous sarcoma virus, RSV
- Most of these sarcoma viruses are defective

Rous was lucky!
How does RSV, but not ALV, cause sarcomas?

- Key finding: the viral genomes from solid tumors were recombinants!
- A piece of the ALV genome is replaced with a segment of host DNA called an oncogene

J. Michael Bishop and H. Varmus identified the oncogene (v-SRC) carried by Rous sarcoma virus in 1976

Nobel Prize to both in 1989 for this discovery

http://www.microbe.tv/twiv/twiv-400/
https://youtu.be/frbMV-YGgQU
Major insight

- ALV infected birds came down with a variety of tumors
- These rare tumors all contained retroviruses derived from ALV, but most were defective and all were different
- Rous was lucky - his RSV isolate was not defective

The retrovirus genomes isolated from each new solid tumor had different host DNA, NOT the v-SRC gene found in RSV

Each new DNA segment had a novel chicken oncogene

A gold mine for molecular oncology
Genomes of transducing retroviruses

Avian transducing retroviruses

- **gag**
- **pol**
- **env**
  
  **Typical progenitor**
  
  5'→ Avian leukosis virus

Rous sarcoma virus

Avian myeloblastosis virus BA1

Avian myeloblastosis virus E26

Avian myelocytoma virus MC 29

Avian myelocytoma virus MH2

Avian sarcoma virus Y73

Avian erythroblastosis virus ES4

Avian reticuloendotheliosis virus

Mammalian transducing retroviruses

- **gag**
- **pol**
- **env**
  
  **Murine leukemia virus**

Abelson murine leukemia virus

Moloney murine sarcoma virus

3611 murine sarcoma virus

Gardner-Arnstein feline sarcoma virus

McDonough feline sarcoma virus

Simian sarcoma virus

HZ4 feline sarcoma virus

Harvey murine sarcoma virus
Defective vs non-defective retroviruses

- Defective viruses require helper virus to produce more virus
- Usually missing envelope proteins
- Envelope genes deleted during oncogene capture
Mechanism for oncogene capture

1. Integration within a proto-oncogene
2. Packaging of deleted and wild-type genomes
3. Nonhomologous recombination during reverse transcription in newly infected cell

Additional rearrangements and point mutations may occur
Proto-oncogenes

- >60
- In all cells, control cell growth
- Highly regulated
- Normal cellular genes abbreviated as c-ONCS, eg c-SRC, c-MYC, c-MOS, C-RAS
- Certain retroviruses isolated from tumors carry altered copies of c-ONCS abbreviated as v-ONCS, eg v-SRC, v-MYC, v-MOS, v-RAS
Subcellular location of major classes of oncoproteins

Growth Factors
- sis

Growth factor receptors
- erbB, fms, kit, ros, sca

Membrane-bound protein kinases
- abl, src, yes

G-proteins
- ras

Cytoplasmic protein kinases
- fps, mos, raf

Nuclear oncogenes, transcriptional regulators, cell cycle regulators
- erbA, ets, fos, jun, myb, myc, rel, ski

Original phenotype in cultured cells:
TRANSFORMATION
Uncontrolled cell growth
The cell cycle

Proto-oncogenes

Mitogenic signals revealed by studying transforming retroviruses

Dominant oncogenes
Retroviruses transform cells by three mechanisms

- Rapid tumor formation: eg RSV; 2 weeks
  - *RSV has activated dominant oncogene in genome (v-SRC)*
  - *Protein produced immediately when virus replicates*
- Intermediate kinetics of tumor formation: eg ALV; months
  - *ALV carries no dominant v-ONC gene*
  - *cis-activation: provirus turns on expression of endogenous oncogene*
- Slow kinetics of tumor formation; eg HTLV; years
  - *HTLV carries no dominant v-ONC gene*
  - *Does not cause cis-activation of local oncogenes*
  - *A viral regulatory protein activates oncogenes by trans-activation*
Proviruses with different transforming potential

e.g., IL2 and the IL2 receptor
Mammalian transforming retroviruses

- Retroviruses transform cells as a mistake or byproduct of their life cycle - they must integrate their DNA
- No obvious viral requirement for transformation or for oncogenesis
Which of the following allows Rous sarcoma virus to transform cells?

A. Presence of the env gene  
B. Presence of a pol gene  
C. Presence of a src gene  
D. Presence of LTRs  
E. None of the above
How study of DNA virus transformation revealed how the cell cycle is regulated

(1900s) Retroviruses

(1950s) *in vitro* studies with RSV

(1943) *in vitro* cancer biology

Convergence

(1960s, 1970s)

(1920s) DNA Viruses

(1959) *in vitro* studies with polyomavirus

(present) *Unified theory of cell growth control*
DNA tumor viruses

First oncogenic DNA virus discovered was *papillomavirus* that causes warts (papillomas) in cottontail rabbits - Richard Shope, 1933
DNA tumor viruses: *Polyomaviridae*

- Ludwig Gross discovered murine polyomaviruses in 1953
- Caused *rare* tumors under certain conditions
  - Natural host is the mouse
  - Ubiquitous in mice; no role in mouse cancer
  - Makes tumors of many tissues (poly-oma) in infant hamsters, rats, rabbits
DNA tumor viruses: *Polyomaviridae*

- Eddy and Hilleman showed that SV40, a contaminant of early poliovirus vaccines, induced *rare* tumors in newborn hamsters - 1962

- Several million Americans were infected with SV40 by poliovirus immunization
  - *Natural host is monkey*
  - *Causes no tumors in monkeys*
  - *Does not transform monkey cells in culture*
Response of different cells to infection

<table>
<thead>
<tr>
<th>Species</th>
<th>SV40</th>
<th>Mouse polyomavirus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Monkey</td>
<td><em>Permissive</em></td>
<td>Non-permissive</td>
</tr>
<tr>
<td>Mouse</td>
<td>Non-permissive</td>
<td><em>Permissive</em></td>
</tr>
<tr>
<td>*Hamster</td>
<td>Semi-permissive</td>
<td>Semi-permissive</td>
</tr>
<tr>
<td>*Rat</td>
<td>Semi-permissive</td>
<td>Semi-permissive</td>
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</table>

* Tumors
Polyomaviral transformation of cultured cells is rare

- 1 transformed cell per 100,000 infected cells
- Why is it so rare?
- How does this property relate to rare tumor formation in animals?
Adenoviridae: Another family of transforming DNA viruses

- Many human serotypes, do NOT cause cancer in humans
- Ad 12-18 tumors in hamsters
- Ad 7-11 poorly tumorigenic in hamsters
- Tumors and transformation of cells: like polyomaviruses and papillomaviruses, very rare events
Key finding: Viral T antigens in tumors and transformed cells

- SV40: Large T, small T
- Polyomaviruses: Large T, middle T, small T
- Papillomaviruses; T encoded by E5, E6, E7 genes
- Adenoviruses: T antigens are E1A, E1B

All different proteins!
T antigens are encoded by essential viral genes

- Required for replication
- Activate viral transcription
- Required for viral DNA synthesis
- Only viral genes *always* retained in tumor cells or transformed cells
- T antigen alone can transform cultured cells
Three seemingly unconnected discoveries in DNA virus biology were critical to understanding the link between viruses, transformation, and the cell cycle.

1. 53 kDa cell protein binds SV40 T antigen

2. Transcription of a set of adenovirus early genes (the E2 gene cluster) requires cell protein E2f (E2 factor)
   - Now a family of proteins called the E2f family

3. E2f found to bind a cellular protein called Retinoblastoma protein (Rb)

p53, Rb and E2F were subsequently discovered to be critical players in control of the normal cell cycle.
The cell cycle

G is for “gap” Resting cells are in “G zero”

When stimulated to divide, cells enter the G1 phase and then into S where they replicate their DNA and prepare for cell division.
A go/no go decision is determined by nutrient concentration and growth factors

- Is the outside world rich enough to replicate the cell?
- Remember: Detectors and signaling proteins for growth were discovered as oncogenes carried by transforming retroviruses
If conditions are not right, cell cycle pauses at restriction point

- No DNA synthesis, no cell division
- The protein that regulates the restriction point decision is \textit{Rb}

\textit{Retinoblastoma protein (Rb)} – if both copies of the Rb gene are lost, develop retinal tumors of retinoblasts that form retina - these cells are gone by age 5

\textit{It is a recessive oncogene (the wild type protein is a tumor suppressor)}
But DNA viruses need cells in S phase so they can replicate their DNA

*Proto-oncogenes*

$\text{Go}$

$\text{Stop}$

*T antigens kick quiescent cells into S phase!!*
When viral T antigens bind to Rb, E2f proteins are released and initiate S phase transcription.
The entry into S decision is under MORE control

- DNA damage or unscheduled DNA synthesis is monitored by p53
- Don’t duplicate damaged genetic information!
- Viruses must counter p53
How do viruses counter p53?

Adenovirus, papillomavirus, or polyomavirus infection

Repression of transcription; binding to apoptosis-inducing proteins

Increased transcription of specific cellular genes

Apoptosis

Cell cycle arrest

Apoptosis

E4 ORF6 + E1B 55 kDa

HPV-16 or HPV-18 E6 protein

Sequestered p53

E1B 55 kDa

p53 accumulation

Increased transcription of specific cellular genes

p21GIP, Bax, Apaf-1

Apothosis

Cell cycle arrest

Apothosis

Principles of Virology, ASM Press
Go to:

b.socrative.com/login/student
room number: virus

**T antigens are:**

A. Encoded by viral genes that are essential for replication  
B. Present in tumors and transformed cells  
C. Encoded by viral genes that have been incorporated into the cell genome  
D. Antagonists of cell cycle checkpoint proteins  
E. All of the above
Two more mysteries remain

1. Why are all viral genes EXCEPT T-antigen encoding genes deleted or turned off in SV40, polyoma, and adenovirus- transformed cells?

2. Why is transformation by these viruses so rare?
Transformation is rare because two low probability events are required

1. Lethal late genes must not be expressed
   - Rare spontaneous deletion of late genes
   - Infection of semi-permissive cells; late gene expression blocked

2. T antigen must be on constitutively and transmitted to every cell
   - Viral DNA encoding T antigen must be integrated into the host DNA
Transformation and tumor formation are abnormal, epigenetic processes for these DNA “tumor” viruses. These events are not required for the normal viral life cycle or transmission. We now understand that even in the natural host, these rare events may occur, leading to tumorigenesis (e.g. HPV).
Transformation is an epiphenomena of a unique reproductive cycle

- DNA tumor viruses must start the cell synthetic machinery to make DNA
- T antigens turn on the cell cycle to start the G1 to S phases through inactivation of normal inhibitor (Rb)
- Inactivation of p53 blocks apoptosis

If lytic events are blocked, cells making T antigens continue to divide – they are transformed

They are on their way to becoming cancer cells
Proto-oncogenes

Revealed by studying transforming retroviruses

(Dominant oncogenes)

Tumor suppressor genes

Revealed by studying DNA tumor viruses

(Recessive oncogenes)