

**Reverse transcription (RT);  
Analysis of RNA ends;  
Techniques for studying transcription**

*Michał Koper, IGiB UW*

**RNA steady-state level =  
equilibrium between  
transcription, processing,  
and degradation**

# Methods of studying transcripts



## Steady state transcript level (Presence in the cel):

- Northern-blot technique
- RT-PCR
- RT-qPCR
- SAGE (Serial Analysis of Gene Expression)
- Microarrays
- *In situ* hybridization (microscopic technique)
- RNAseq: High-throughput sequencing of transcriptomes



## Nascent transcripts (RNA „in statu nascendi”):

- TRO (Transcription Run-On)
- FRAP (Fluorescence Recovery After Photobleaching; microscopic technique).
- GRO-Seq (Global Run-On Sequencing).



## In vivo Protein-DNA/RNA interactions:

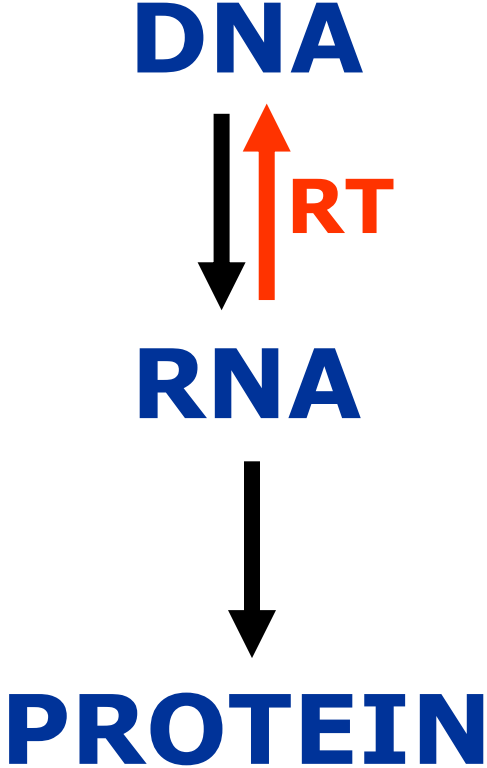
- **ChIP** (Chromatin Immunoprecipitation), e.g., presence of RNA Pol on gene; TFs-DNA interactions; "ChIP on ChIP"; ChIP-seq
- **RIP** (RNA Immunoprecipitation) – interactions of RRM proteins with RNA
- **DIP** (DNA Immunoprecipitation)
- **CLIP** (*in vivo* Cross-Linking and Immunoprecipitation; RNA-protein complexes).

# Reverse transcription (RT) reaction

# Discovery of reverse transcriptase



Nobel Prize in Medicine and Physiology 1975.



Howard  
M. Temin



David  
Baltimore



Renato  
Dulbecco

*„For their discoveries concerning the interaction between tumour viruses and the genetic material of the cell“*

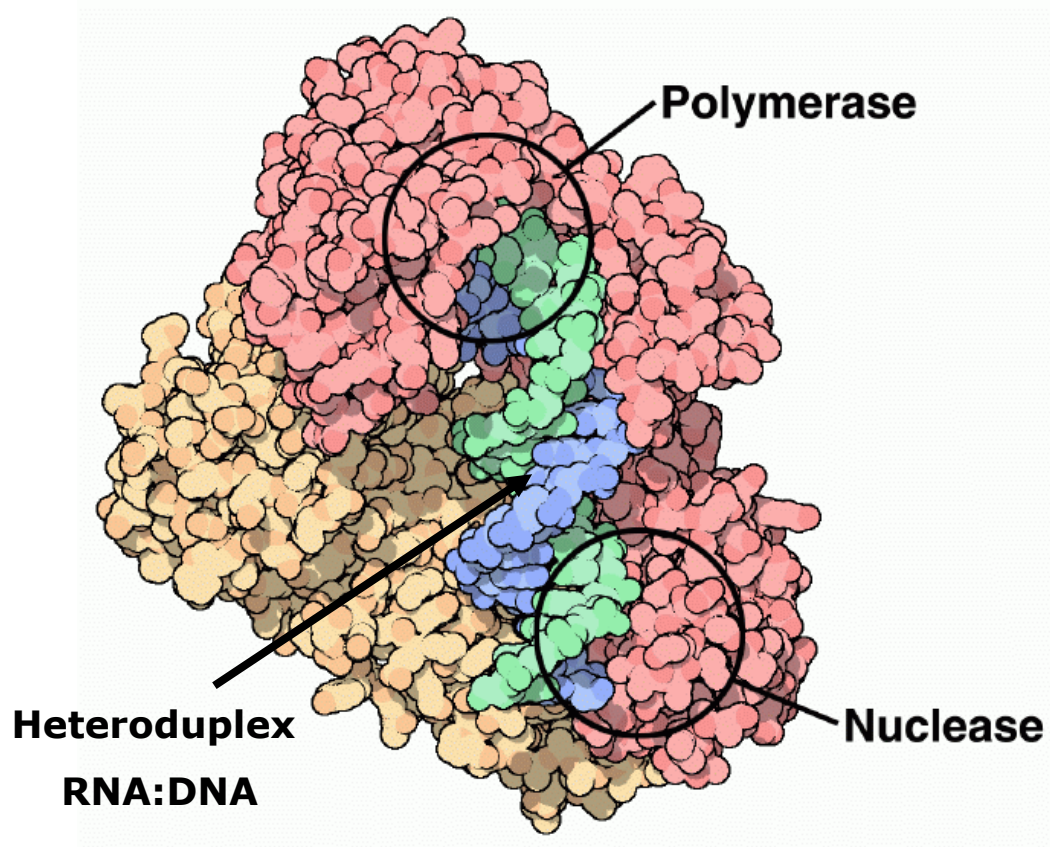
## RNA-DEPENDENT DNA POLYMERASE

Temin, H. M., and S. Mizutani. 1970. **RNA-dependent DNA polymerase in virions of Rous sarcoma virus.** Nature 226: 121-123.

# Reverse transcriptase: 2 enzymes in 1

## HIV-1 RT

(heterodimer)



*Ding J. et al, 1998, J. Mol. Biol. 284; 1095-1111*

**RNA-dependent DNA  
polymerase activity**

(and DNA dependent DNA Pol)

**Ribonuclease H  
(RNase H) activity**

# RT requires a primer to start DNA synthesis

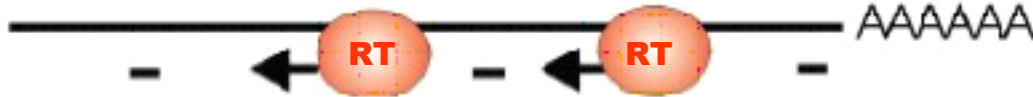
**gene-specific primers**



**Oligo-dT primers,  
Can be "anchored", i.e.  
TTTTTTTTTTT(A/C/G)**



**short random primers,  
e.g. hexamers, octamers**



# Applications of reverse transcription

## **QUALITATIVE techniques („presence“):**

- **structure of the genes (introns)**
- **determination of the RNA 5' ends**
- **structure of the RNA 3' ends**
- **detecting the expression of constructs**

## **QUANTITATIVE techniques („levels“):**

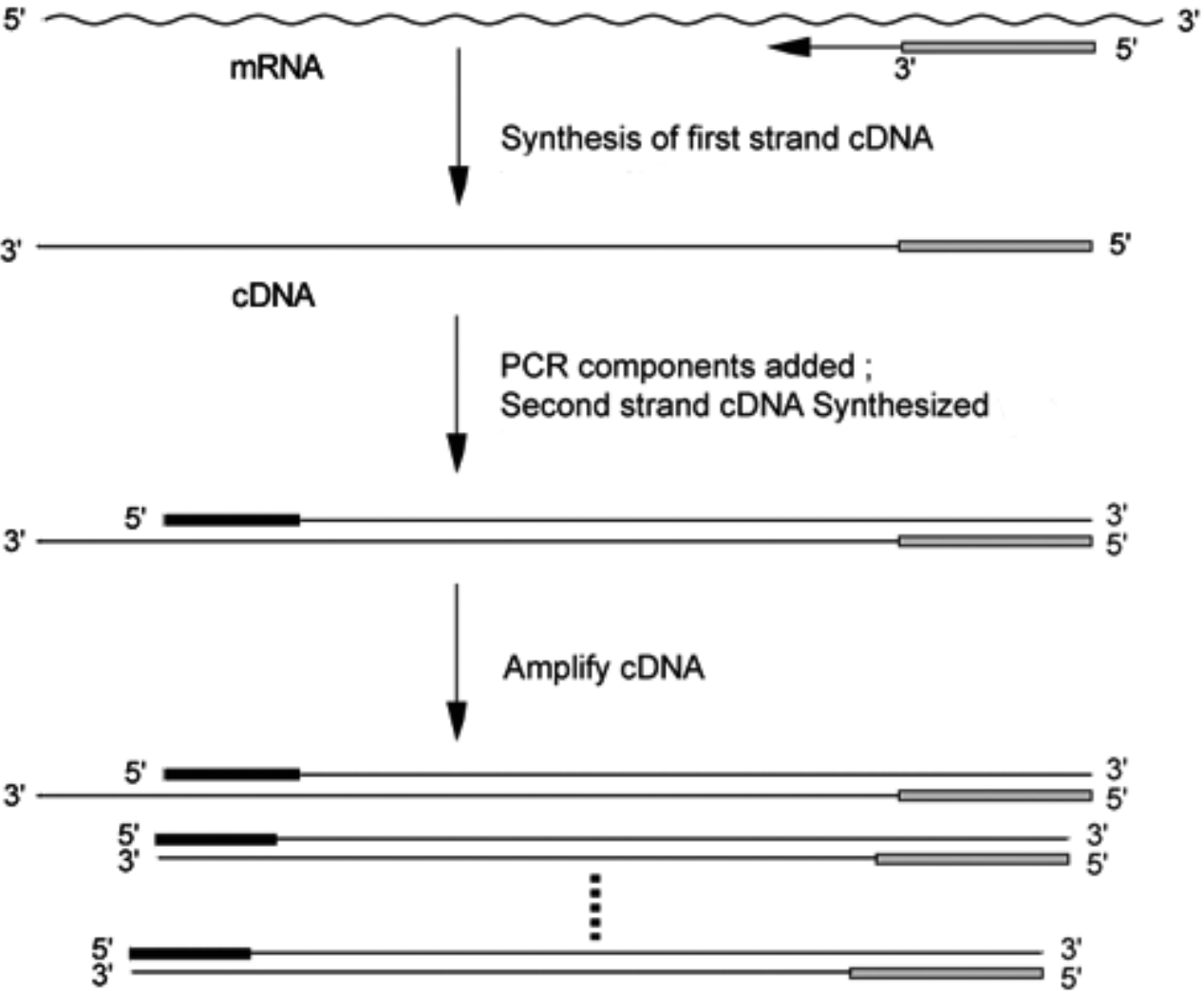
- **semi-q RT-PCR**
- **RT-qPCR**

## **BOTH – QUALITATIVE and QUANTITATIVE techniques**

- **RNAseq - transcriptome sequencing**



# Principle of RT-PCR



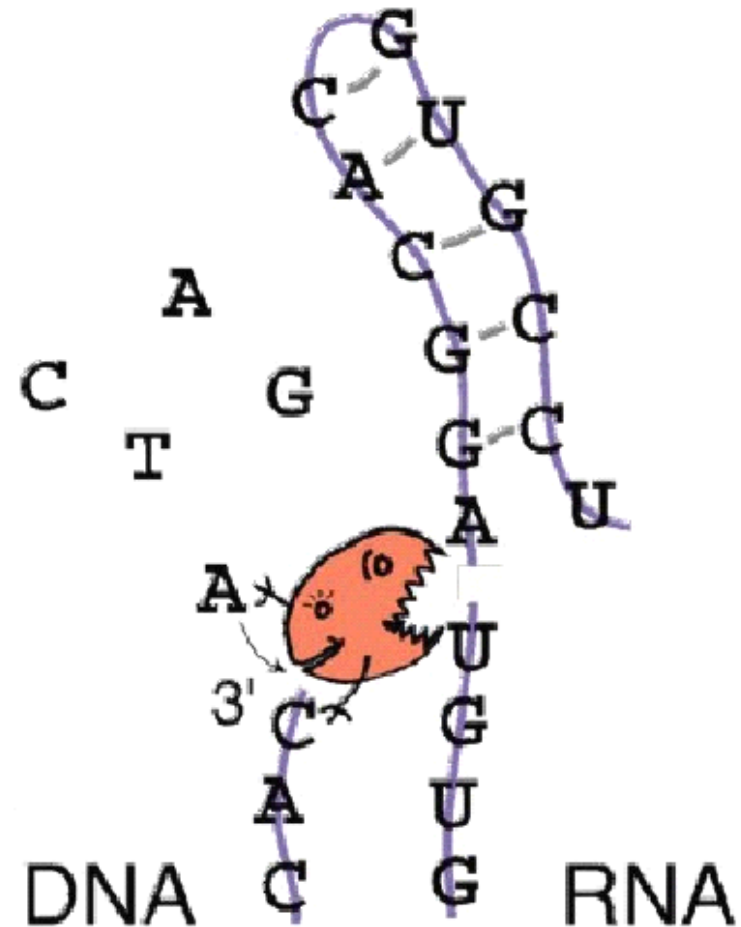
# RNase H activity of wild type RT

Responsible for:

- *removal of DNA:RNA hybrids*
- *generation of PPT primers*
- *removal of tRNA and PPT primers*

Particularly active at sites with RNA secondary structure where RT slows down.

In some commercially available RT RNase H activity is lowered or removed.



Courtesy of the  
TATA Biocenter

# Properties of various RTs

	Temp
MMLV RNase H- Minus (Promega, Germany)	37
M-MLV (Promega)	45
Avian Myeloblastosis Virus (AMV) (Promega)	37
Improm-II (Promega)	45
Omniscript (Qiagen, Germany)	37
cloned AMV (cAMV) (Invitrogen, Germany)	45
ThermoScript RNase H- (Invitrogen)	50
SuperScript III RNase H- (Invitrogen)	50

Sthalberg et al, 2004, Comparison of reverse transcriptases  
In gene expression analysis, Clin Chem. 50(9):1678-80.

**AMV: Avian Leukemia Virus**

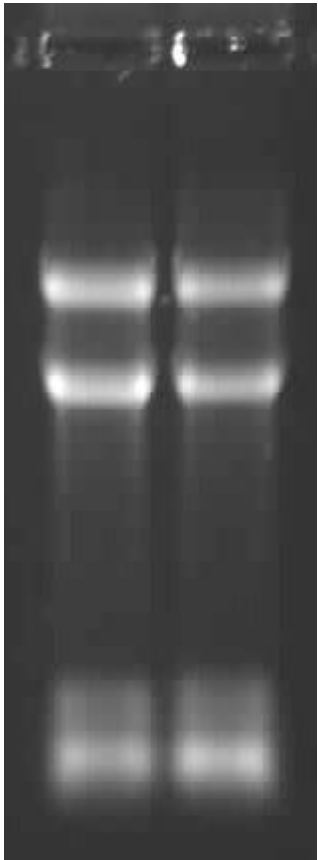
**MMLV: Moloney Mouse Leukemia Virus**

# GOOD LABORATORY PRACTICE AT EVERY STAGE OF WORK WITH RNA!

DNase treatment

1h, 37° C

- +



**RNA quality is  
a key factor for RT!**

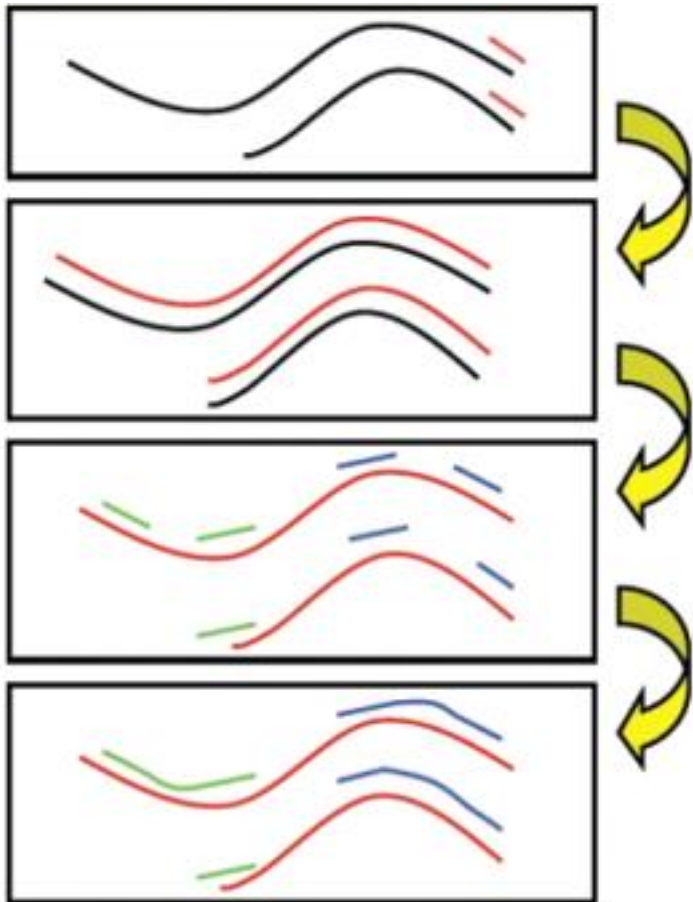
**RNA quality control after isolation  
is always necessary!**

**DNase treatment is always  
necessary!**

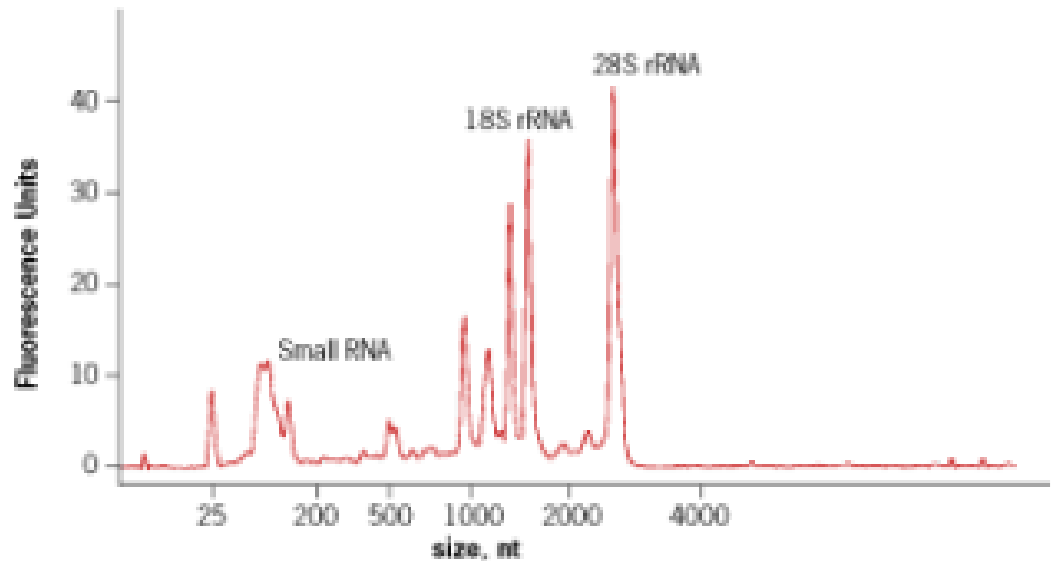
**Recommended use of  
RNase inhibitors!**

# RNA quality assessment

## 5'-3' RNA ends integrity test



## Microfluidic Analyzers



- **Agilent: Bioanalyzer, Tapestation**
- **Biorad: Experion**
- **Capillary electrophoresis**

# Efficiency and specificity of RT

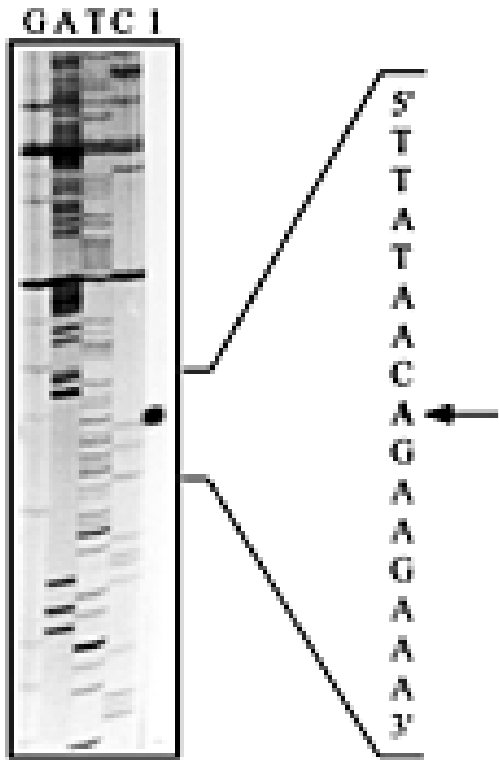
Efficiency and specificity of reverse transcription reaction **STRONGLY** depends on:

- **Methods of priming**
- **Type of reverse transcriptase (features of the enzyme)**
- **The sequence of a particular RNA molecule – it's tendency to form complex secondary structures**
- **Performance differences can be as high as 100X!!!**
- **The optimal RT conditions should always be determined experimentally**

# **Analysis of 5' and 3' RNA ends**

# Primer extension

## Determination of the transcription start site



**Annealing of radioactively labeled GS primer  
to the RNA 5' end**



**Reverse transcription**



**Separation in polyacrylamide gel together  
with sequencing reactions carried out from  
the same primer**

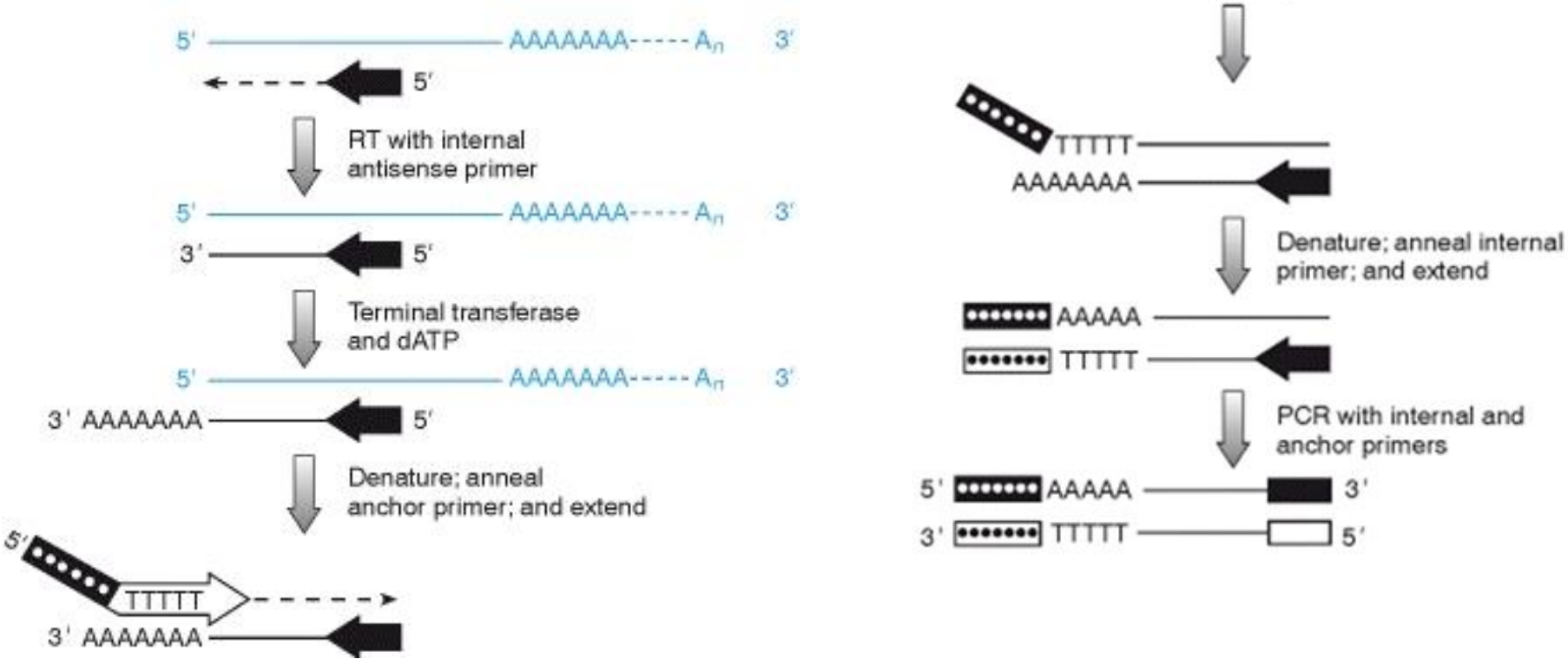
*T.-R. Lee, H.-P. Hsu, and G.-C. Shaw,  
2001, J. Biochem. Vol. 130, pp. 569-574*



# Analysis of 5' RNA ends - 5' RACE

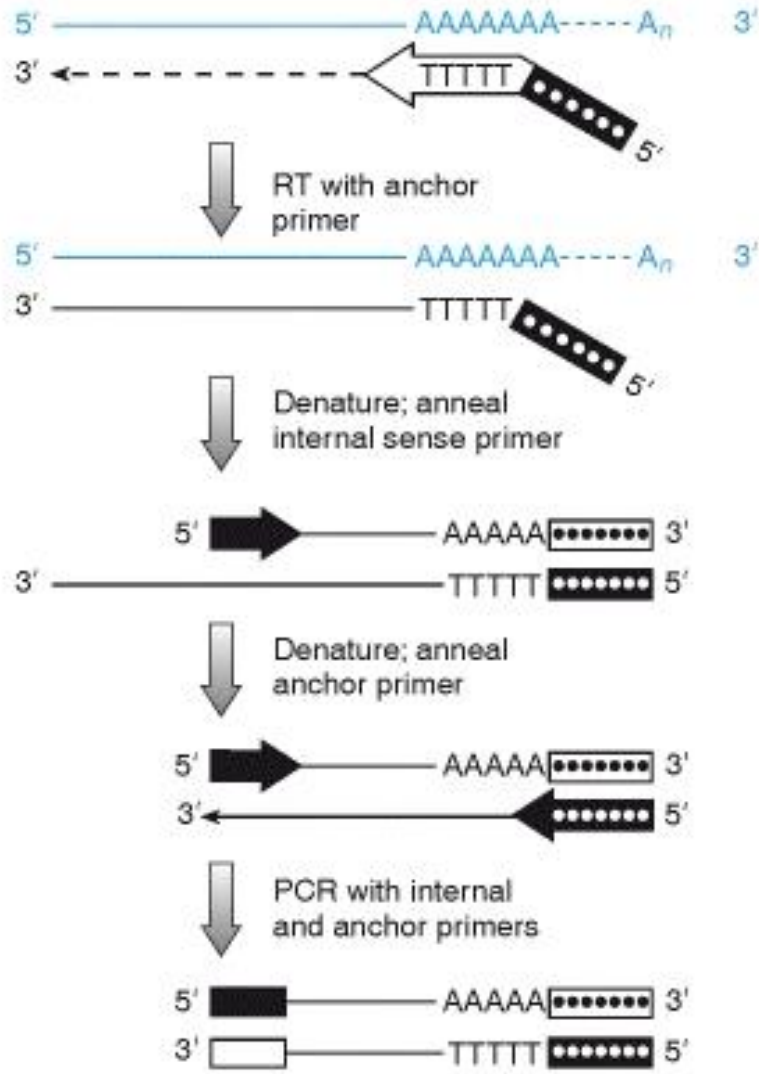
## Rapid Amplification of 5'cDNA Ends

### anchored PCR



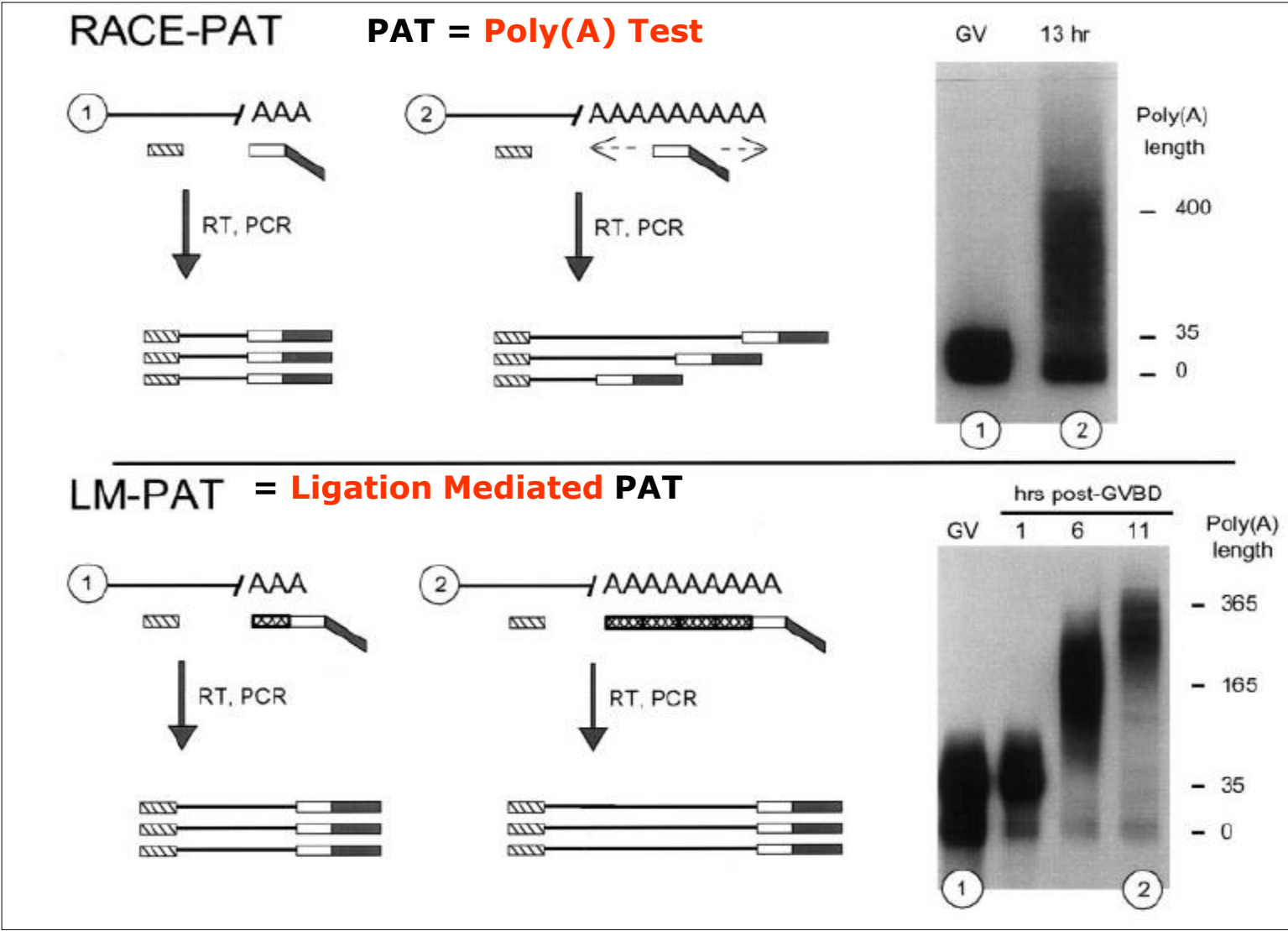
# Analysis of 3' ends of polyadenylated RNAs - 3' RACE

## Rapid Amplification of 3'cDNA Ends





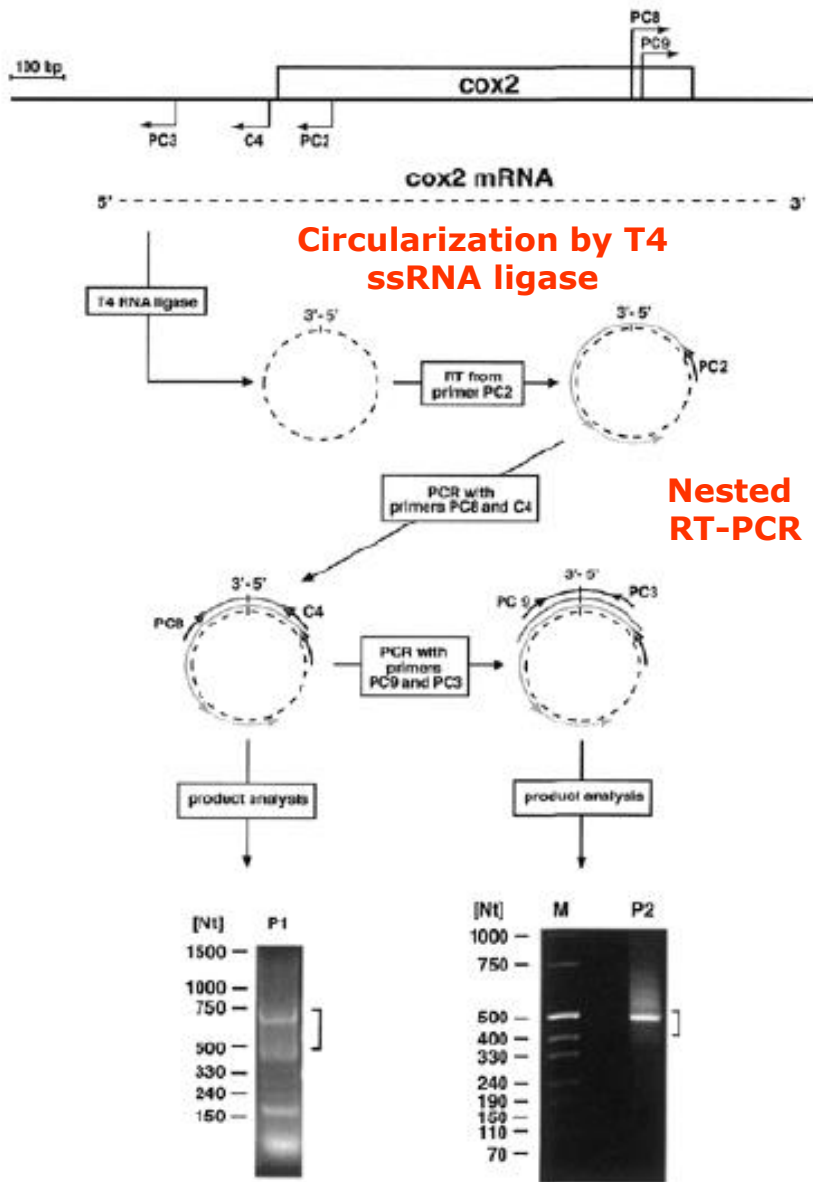
# Analysis of polyA<sup>+</sup> 3' ends



# Circular RT-PCR (cRT-PCR)

## Analysis of 5' and 3' ends of the mitochondrial COX2 transcript in plants

Khun and Binder, NAR, 2002.



1	TCGCAATAGCGAGCCTACTTATCGCTTTTTT	+300	-277	AGAAAGAA...
3	TCGCAATAGCGAGCCTACTTATCGCTTTTTT	+299	-280	GATAGAAA...
1	TCGCAATAGCGAGCCTACTTATCGCTTTTTT	+299	-278	TAGAAAGA...
1	TCGCAATAGCGAGCCTACTTATCGCTTTTTT	+299	-277	AGAAAGAA...
1	TCGCAATAGCGAGCCTACTTATCGCTTTTTT	+299	-273	AGAATCGT...
1	TCGCAATAGCGAGCCTACTTATCGCTTTTTT	+299	-270	ATCGTTCA...
1	TCGCAATAGCGAGCCTACTTATCGCTTTTTT	+299	-269	TCGTTCAG...
1	TCGCAATAGCGAGCCTACTTATCGCTTTTTT	+298	-291	ATCACGTA...
2	TCGCAATAGCGAGCCTACTTATCGCTTTTTT	+298	-289	TCGTTCAG...
2	TCGCAATAGCGAGCCTACTTATCGCTTTTTT	+298	-261	TAGGCTAA...
1	TCGCAATAGCGAGCCTACTTATCGCTTTTTT	+297	-262	GTAGGCTA...

Σ 15

# Circular RT-PCR (cRT-PCR) cont.

- RNA ligase ligates **RNA** molecules **lacking a cap** at the 5' end
- When testing molecules with a cap, **the cap must be removed** to detect RNA
- The cap can be removed by enzymes that hydrolyze P-P bonds, e.g. bacterial RNA 5'-pyrophosphohydrolase RppH (NEB) or plant acid pyrophosphatase Cap-Clip (CellScript) (formerly **acid pyrophosphatase from tobacco TAP**, now not produced) or **by digesting RNA-DNA duplexes with RNase H** (makes sense when the RNA is highly expressed, exact 5' end of the transcript is known and we are not interested in alternative 5' ends)

<https://www.neb.com/en/tools-and-resources/selection-charts/mrna-decapping-selection-chart>

# Techniques for studying transcription

# Transcription Run ON technique



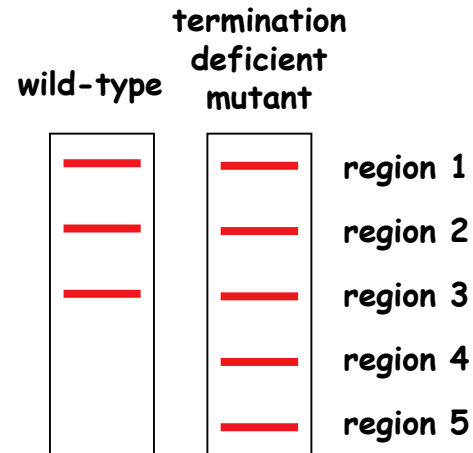
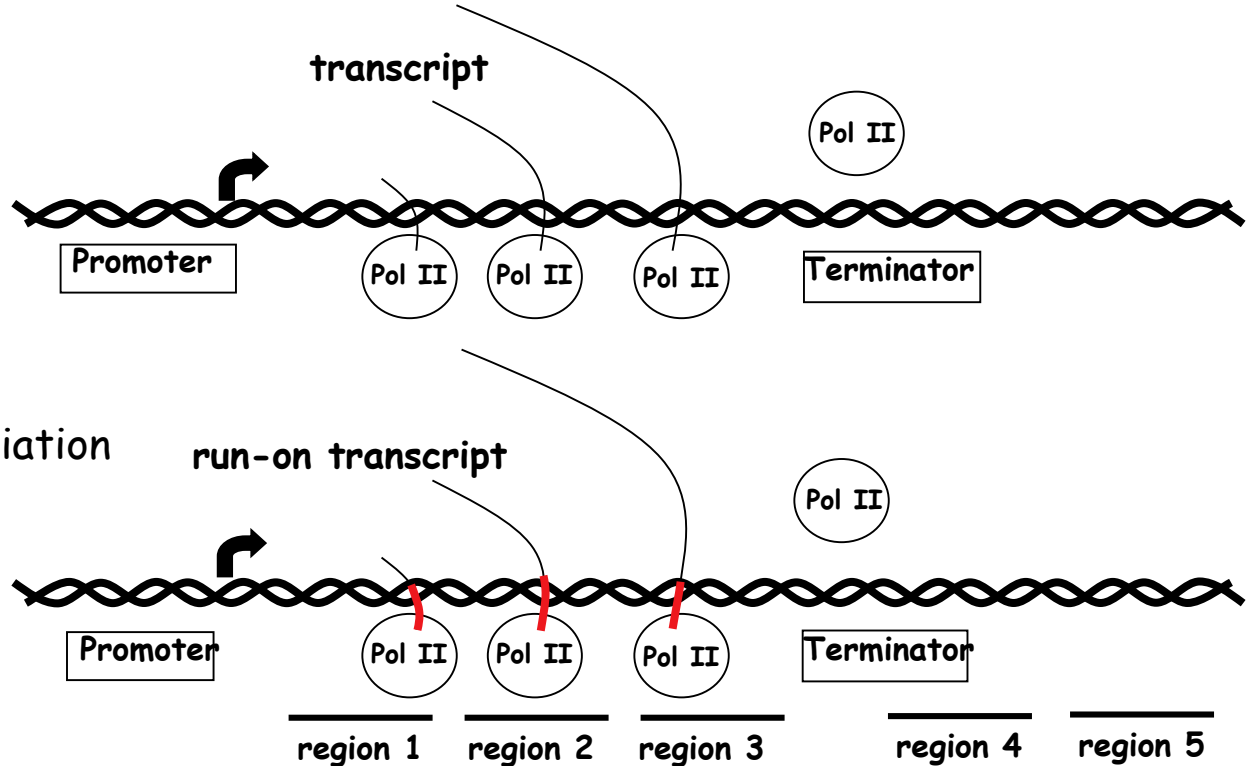
# Transcription Run-On (TRO)

early log-phase yeast culture  
↓  
sarkosyl treatment  
- permeabilize cells  
- inhibit transcription initiation

transcription reaction  
with P-UTP<sup>32</sup>

RNA extraction

Hybridize RNA to single  
strand DNA probes relative  
to different regions on the  
gene of interest



Courtesy of  
B. Dichtl

# Molecular probes used in TRO

---

**Antisense** probes that are approx. 70 to 400 nt in length:

- **ssDNA** obtained from M13-derived phagmids
- **Oligonucleotides** usually 70-85 nt
- *in vitro* transcribed **RNA** (e.g., using T7 promoter)
- **PCR** products
- **dsDNA plasmids**

# Blotting TRO probes on filters

Standard **nylon membranes** for nucleic acids, e.g. *Amersham Hybond N*.

Vacuum application, e.g. *Bio-Rad slot/dot blotter*.

The amounts of probes applied depends on the expression levels of the transcripts under study, e.g. for ACT1, 25s or 18s rRNA - **5  $\mu$ g oligonucleotides** 75 nt. or **5  $\mu$ g ssDNA M13 probes** (inserts about 250 nt).



# Performing TRO in yeast - *in vivo* transcription

Modified from Birse et al, Science 280 (1998):

**Culture cells to  $OD_{600} \sim 0.2-0.3$  (0.1-0.4).**

**Permeabilization with sarcosyl - sarcosyl breaks down complexes - "rips off" proteins that interact with the polymerase - any induction before cell harvesting!**

**Transcription with a mixture of AGC and  $^{32}P$ -UTP for 5 min - longer makes no sense because the polymerase "falls off" after a few tens/hundreds of sec.**

# Performing TRO in yeast - isolation of labeled RNA and hybridization

---

**Isolation of RNA** by any method, e.g. "hot phenol", GTC/phenol/zirconia beads.

**Hydrolysis of RNA in NaOH solution** before application to filters. Depending on the duration of hydrolysis, different sizes of fragments obtained.

Hybridization, washes and exposure similar to the **northern-blot** technique.

# TRO applications and variants

## Quantitative applications:

- Testing the transcriptional activity (power) of promoters
- Determination of transcription rates for individual genes
- Differences in transcriptional activity between mutants or upon external conditions.

## Qualitative applications:

- Study of transcription termination defects (*read-through*)
- Distinguishing between RNA Pol presence and RNA synthesis

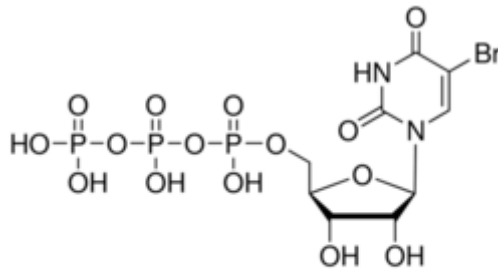
## TRO Variants:

- Nuclear Run-On, on isolated cell nuclei: Gariglio (1981) and Brown (1984)
- Mitochondrial Run-On, on isolated mitochondria

# GRO-seq: Global (Genomic) Run-On

RNA labeling: Run-On in the presence of BrU or 4tU

## 5-Bromouridine (BrU)

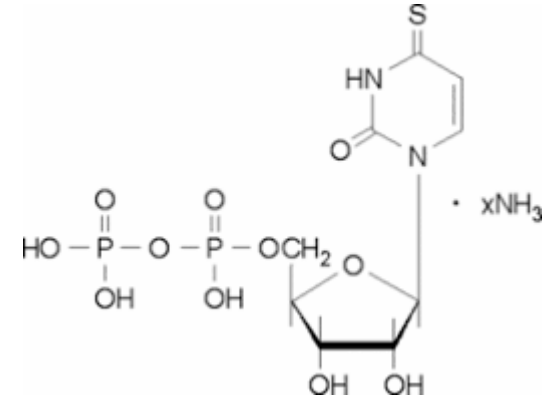


• xNa



RNA fragmentation and immunoprecipitation on agarose-anti BrU resin

## 4-thiouridine (4tU)



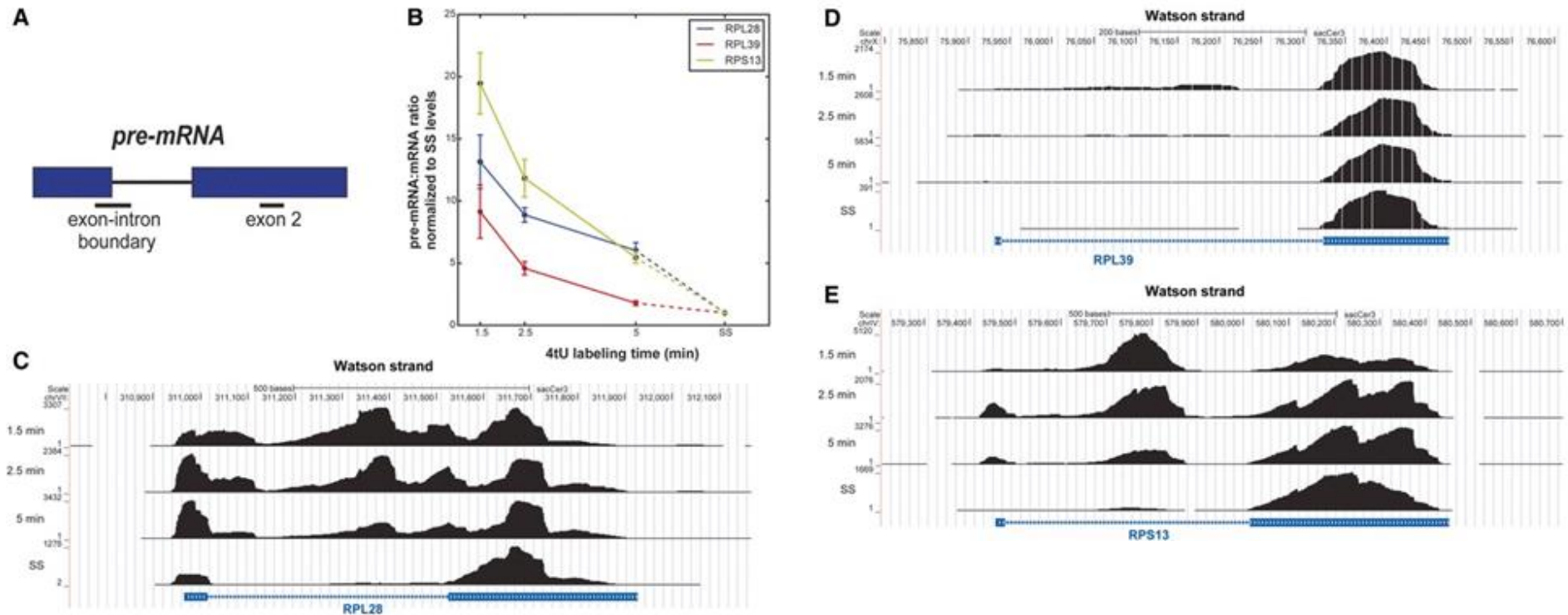
Biotinylation of the thiol group and binding to the streptoavidin complex



Reverse transcription  
-> RNAseq

# 4tU-seq

## Transcriptome-wide RNA processing kinetics revealed using extremely short 4tU labeling

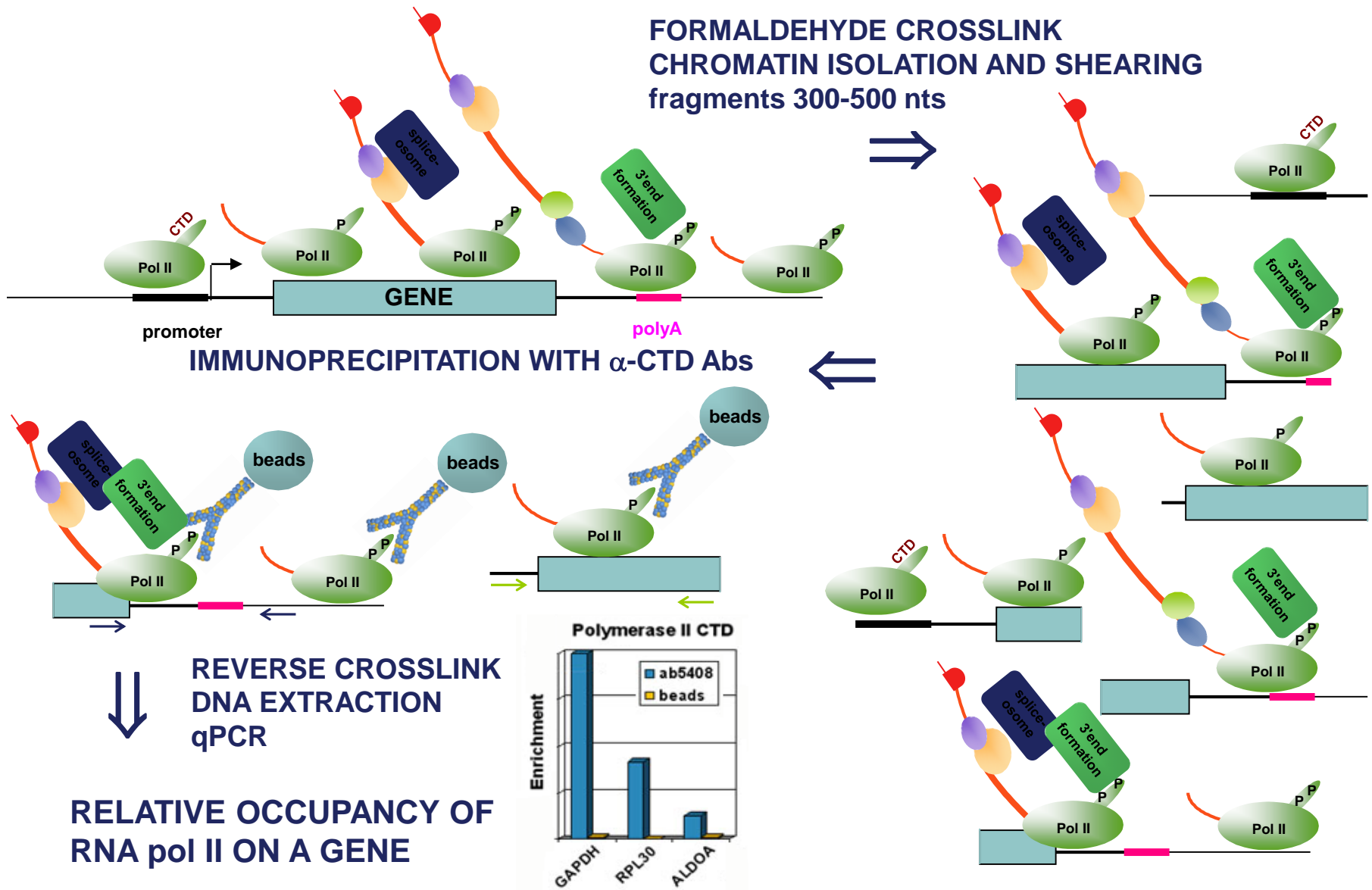


(Barras et al, Genome Biol 2015)



# ChIP – Chromatin Immunoprecipitation

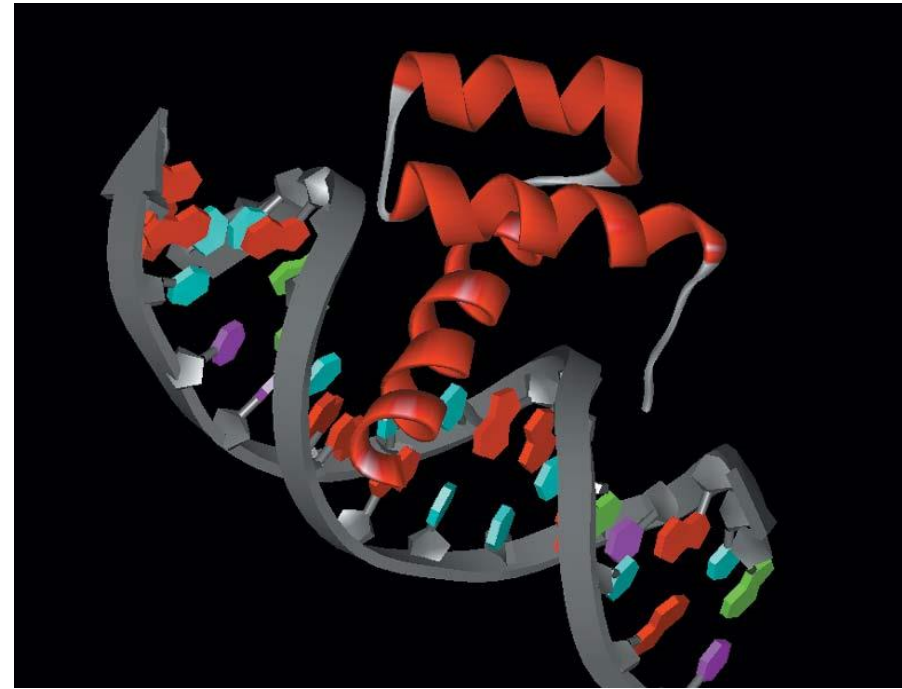
# Chromatin immunoprecipitation - ChIP

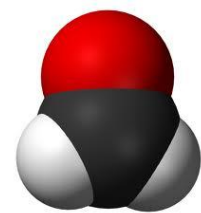


# ChIP applications

## Studying protein-DNA interactions *in vivo*!

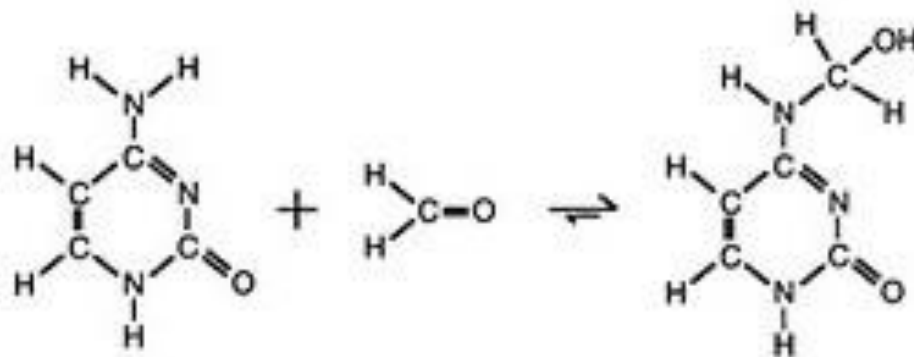
- Transcription factor binding sites
- Presence (distribution) of RNA Pol along chromatin (genes)
- Histone modifications
- Chromatin remodeling factors



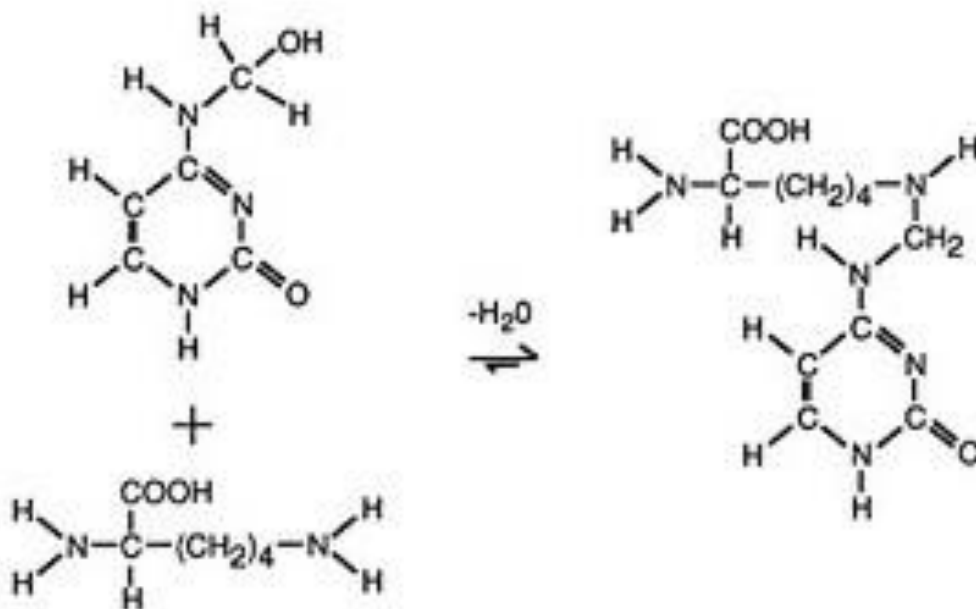


# Crosslinking agent: formaldehyde

**Reaction 1 (Shiff base is formed)**



**Reaction 2**



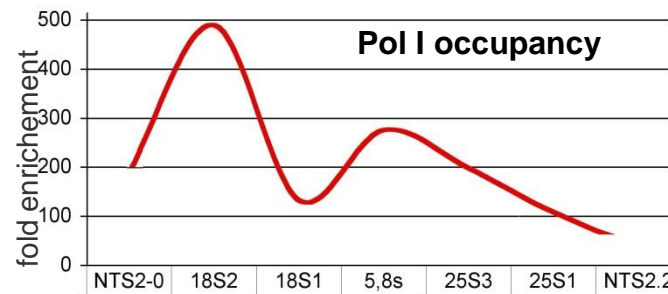
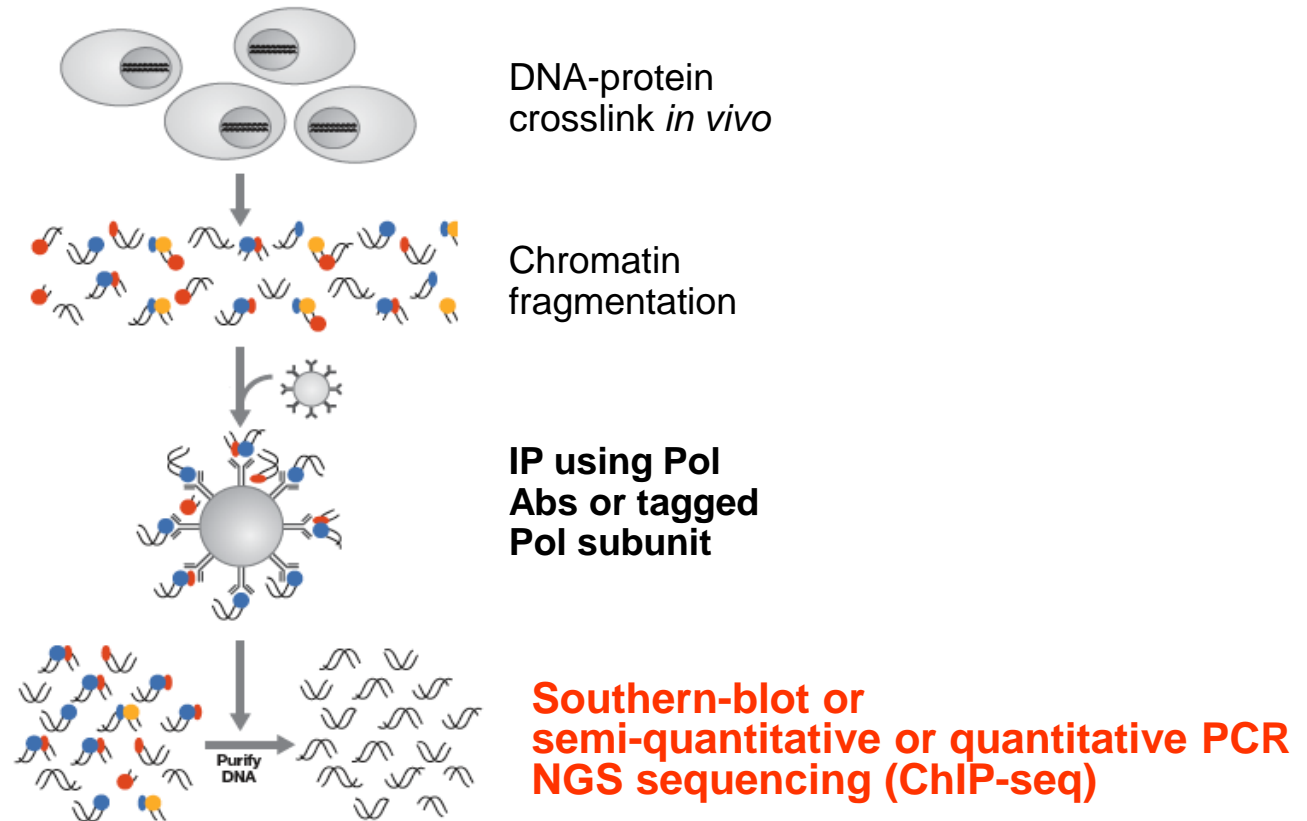
# Resolution vs. specificity of crosslinking

---

- ***In vivo* crosslinking, high reactivity of formaldehyde**
- **High resolution: 2Å**
- **Problem with crosslinking of weaker interacting proteins with DNA (greater distance between atoms in residues)**
- **Other cross-linking agents (e.g., DTBP)**

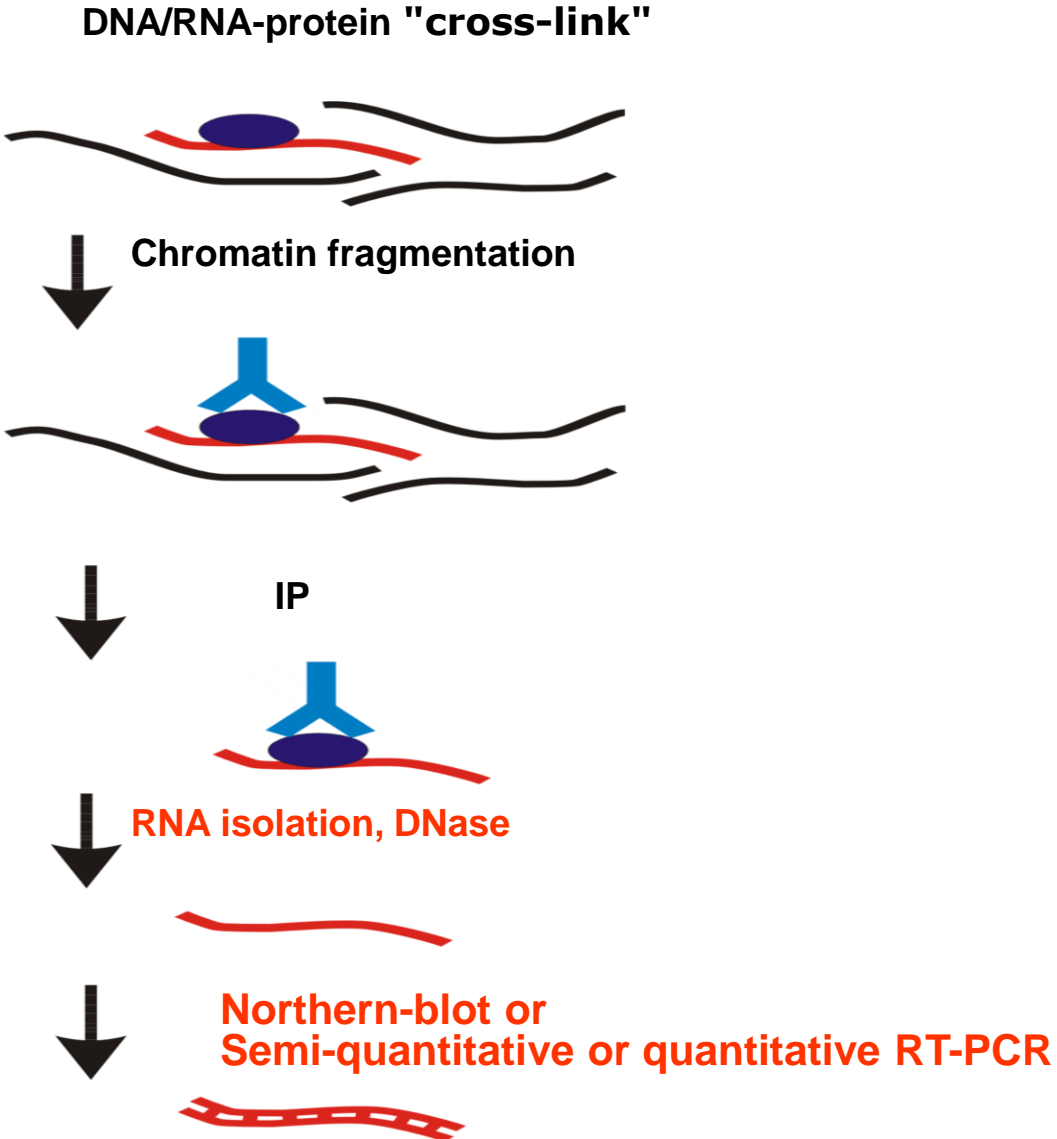
# ChIP procedure

## Chromatin Immunoprecipitation (ChIP)



# RNA Immunoprecipitation - RIP

# RNA immunoprecipitation is similar to ChIP





# Transcription termination

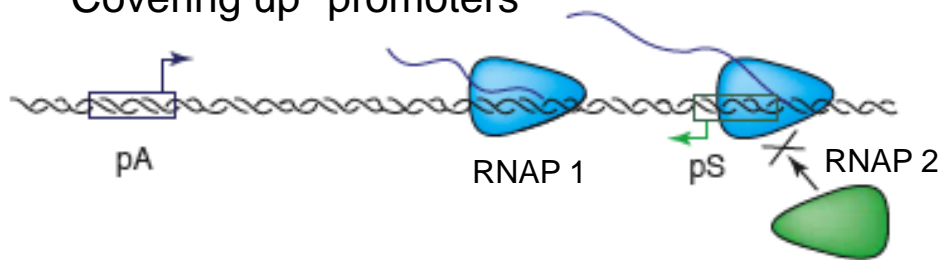
# Why to terminate transcription?

In eukaryotes, few genes are organized in polycistrons (exceptions: ncRNA, mtRNA)

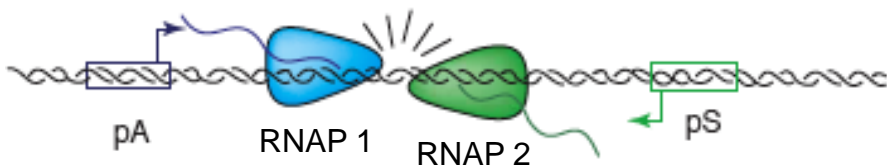
**Incorrect transcription termination affects gene expression**

## 1. co-transcriptionally

"Covering up" promoters

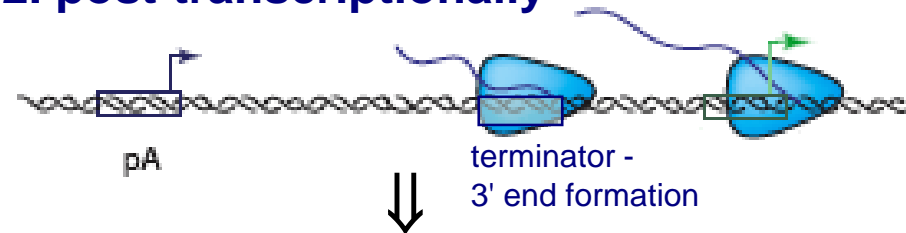


Polymerase collisions

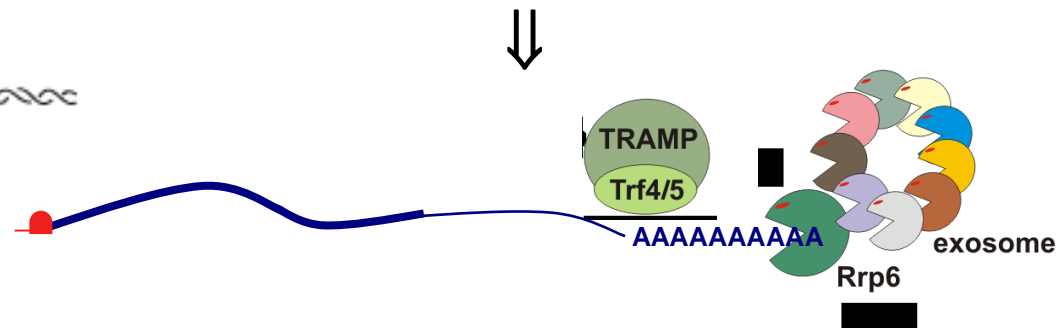


Shearwin et al, *TiG.*, 2005

## 2. post-transcriptionally

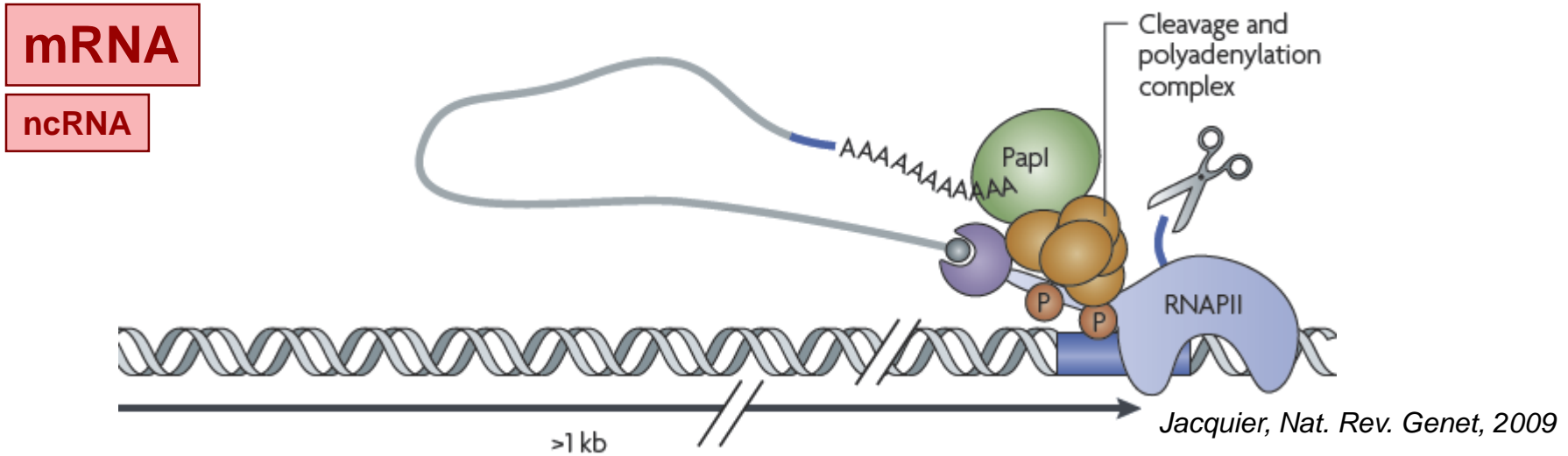


"Readthrough" - Transcription Readthrough (RT).



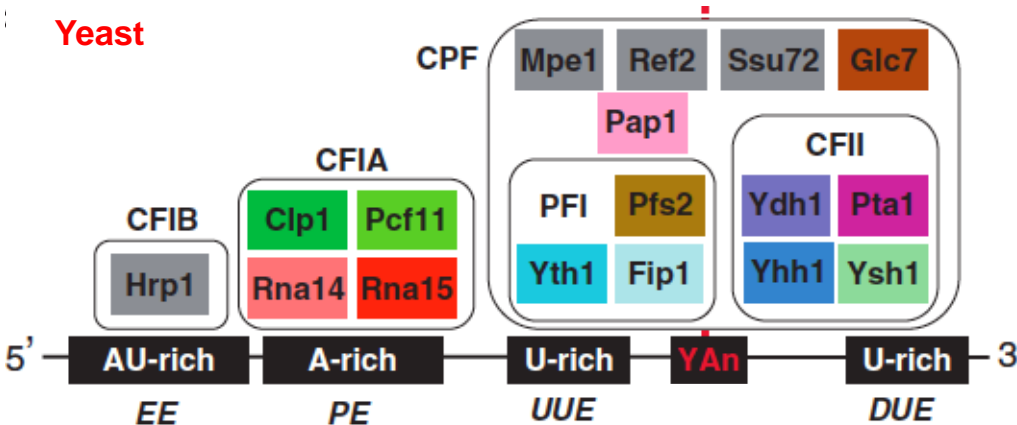
Removal of defectively elongated transcripts (nuclear RNA surveillance)

# RNA polymerase II - cleavage and polyadenylation

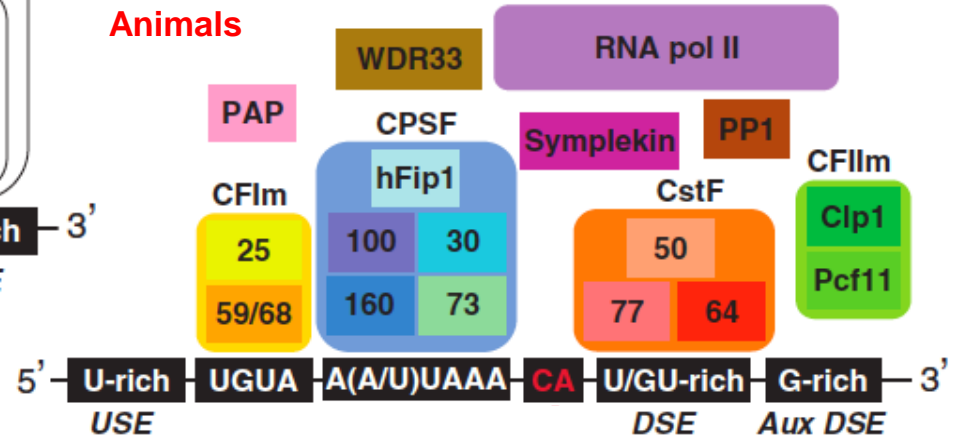


Cleavage and polyadenylation complex (CP) (recruitment via Ser2-P CTD)

Yeast



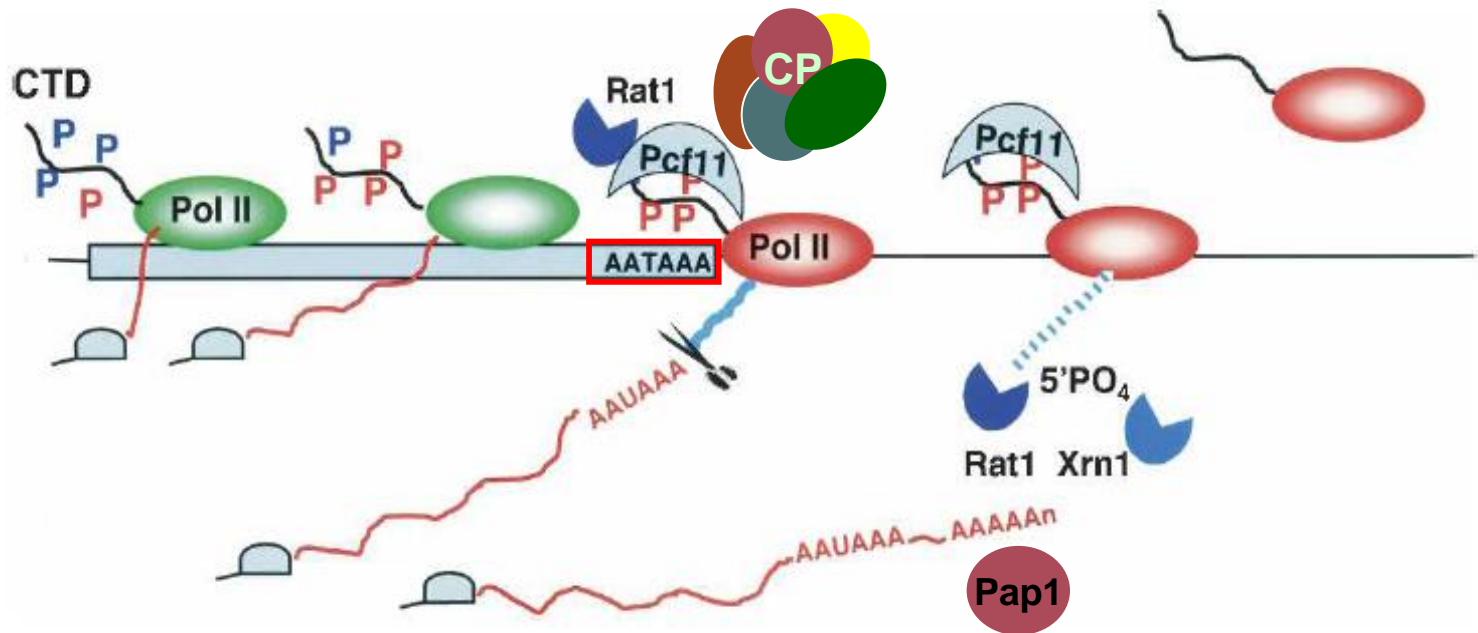
Animals



In yeast, cleavage by Brr5/Ysh1 and in humans by CPSF-73

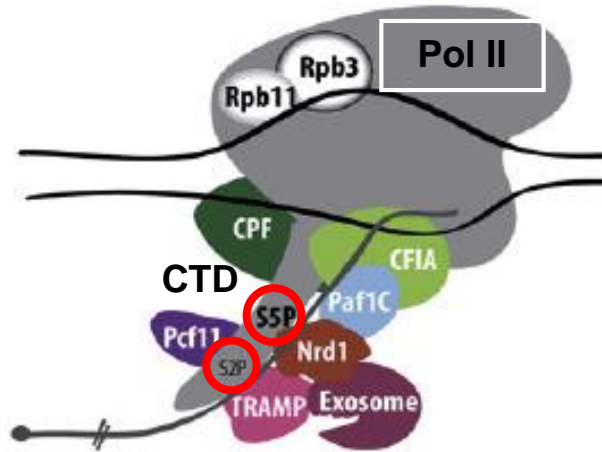
Millevoi and Vagner, NAR, 2008

# Hybrid - allosteric-torpedo - model of RNA polymerase II transcription termination



Luo and Bentley, *Gene Dev*, 2006

# Nrd1/Nab3/Sen1 dependent termination of pol II



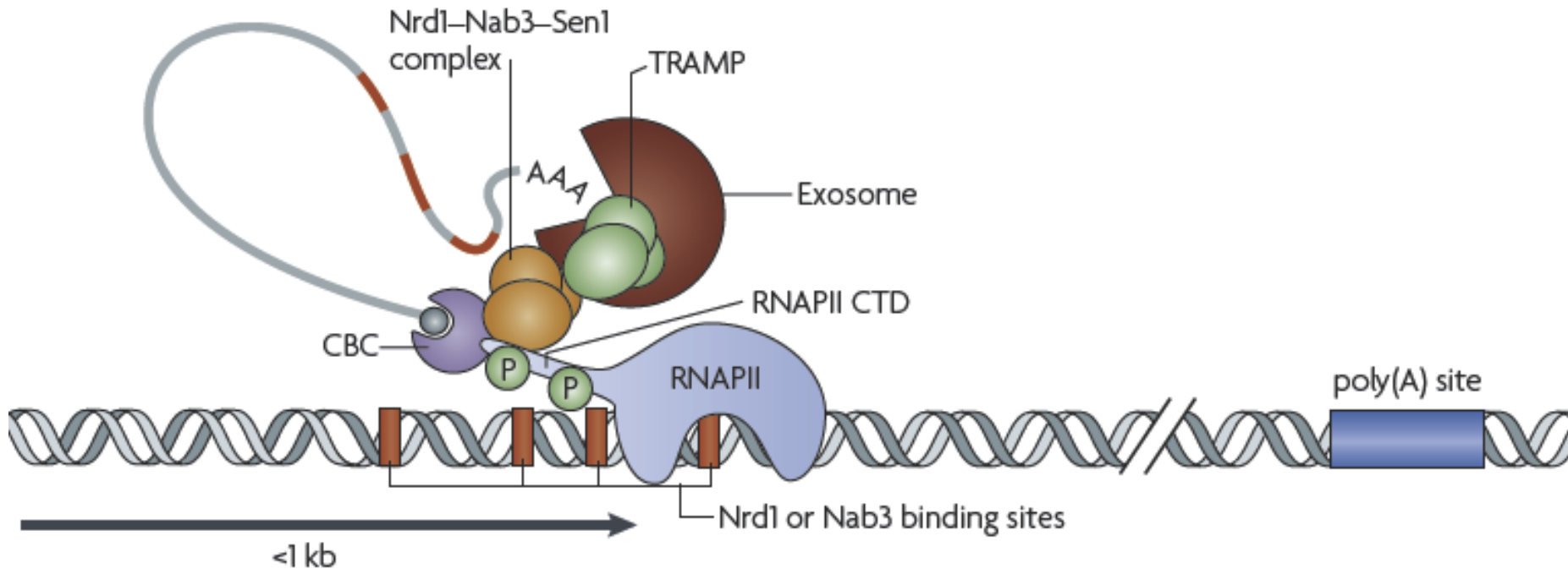
Nrd1/Nab3/Sen1 - **NNS complex**

- sn/snoRNAs
- CUTs
- short mRNAs (< 600 nt)

ncRNA

mRNA

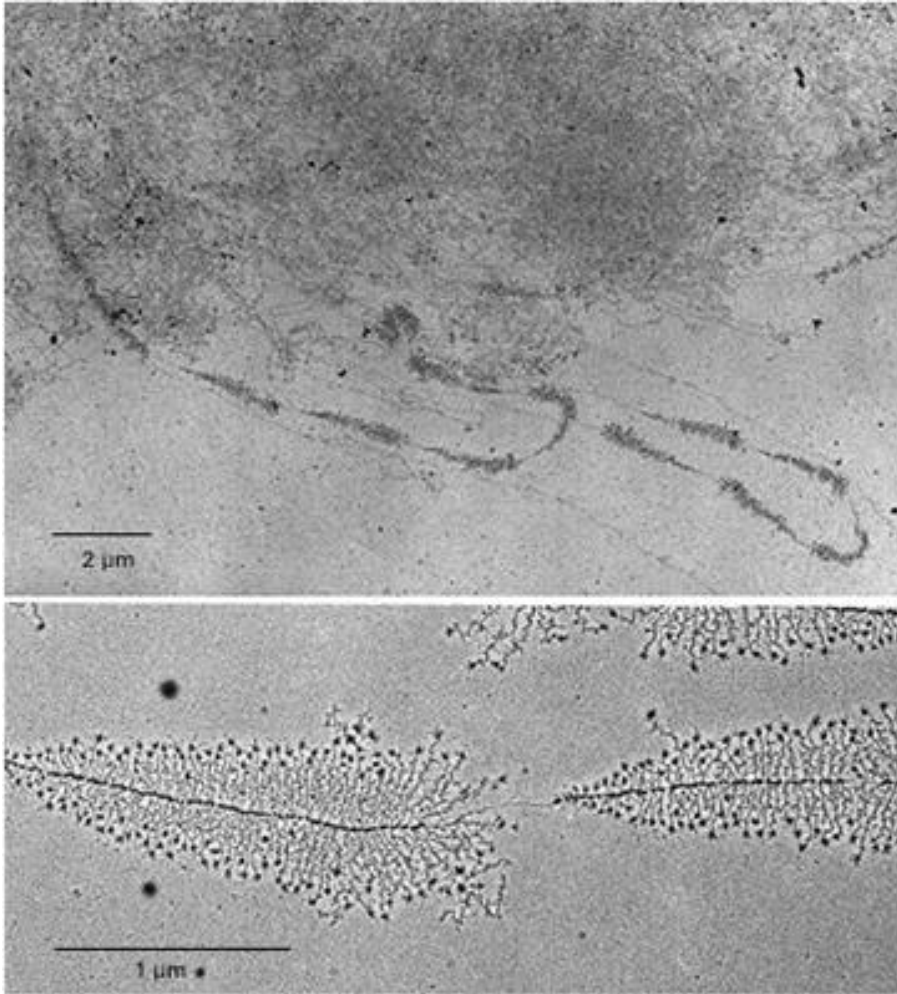
(Recruitment through Ser5-P CTD)



# **Transcription termination of RNA polymerase I**

# rDNA genes in the nucleus

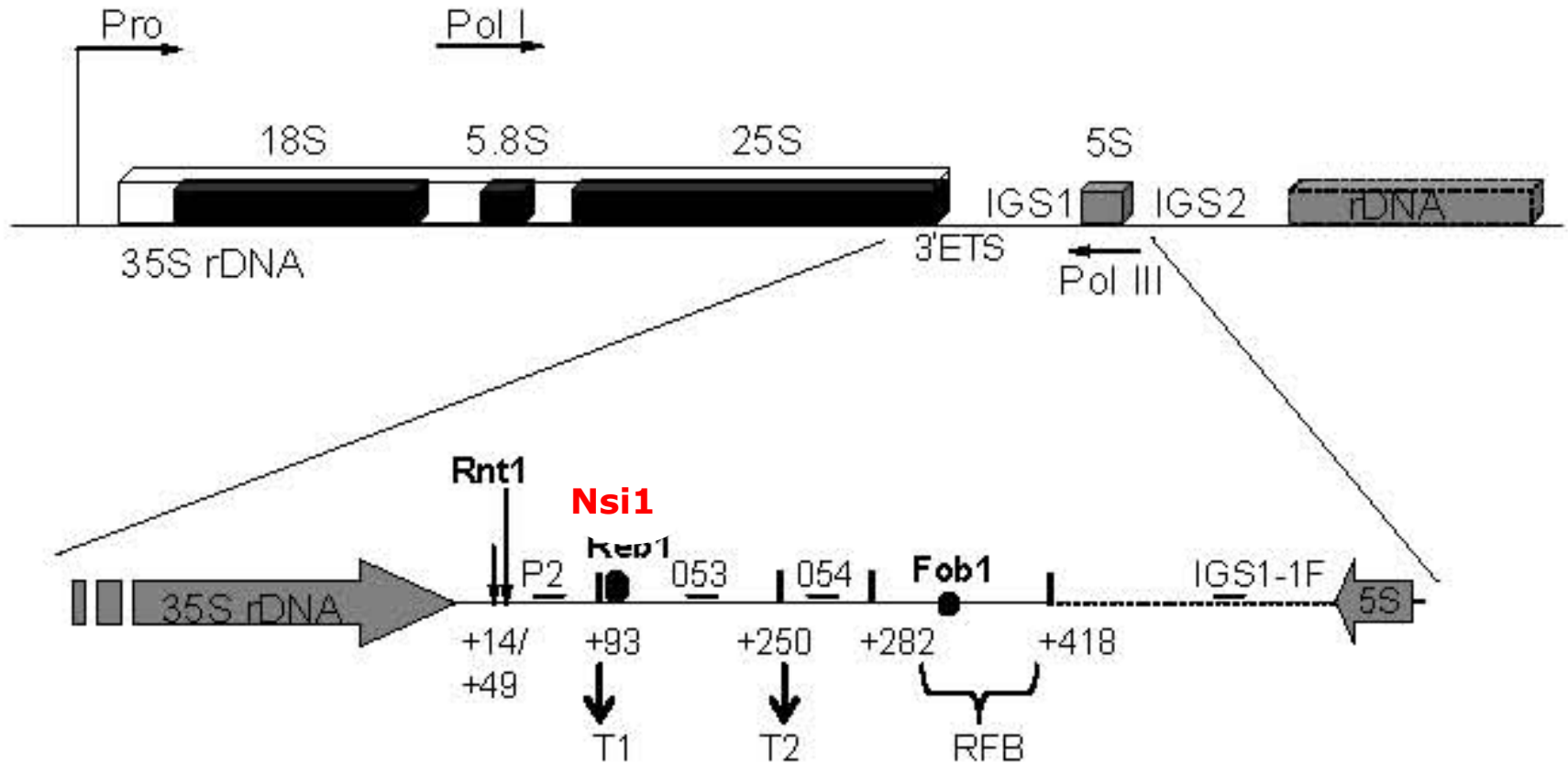
## Miller's spreads - "Christmans tree".



Top: V.E. Foe, Cold Spring Harbor Symp. Quant. Biol. 42:723-740, 1978; bottom: Ulrich Scheer, MBC on NCBI.

- Pol I transcribes ribosomal RNA (**35S rRNA**)
- Pol I RNA – up to **80%** of all transcriptional activity of the cell nucleus!!!
- In yeast, rDNA genes are organized in **150-200** tandem repeats

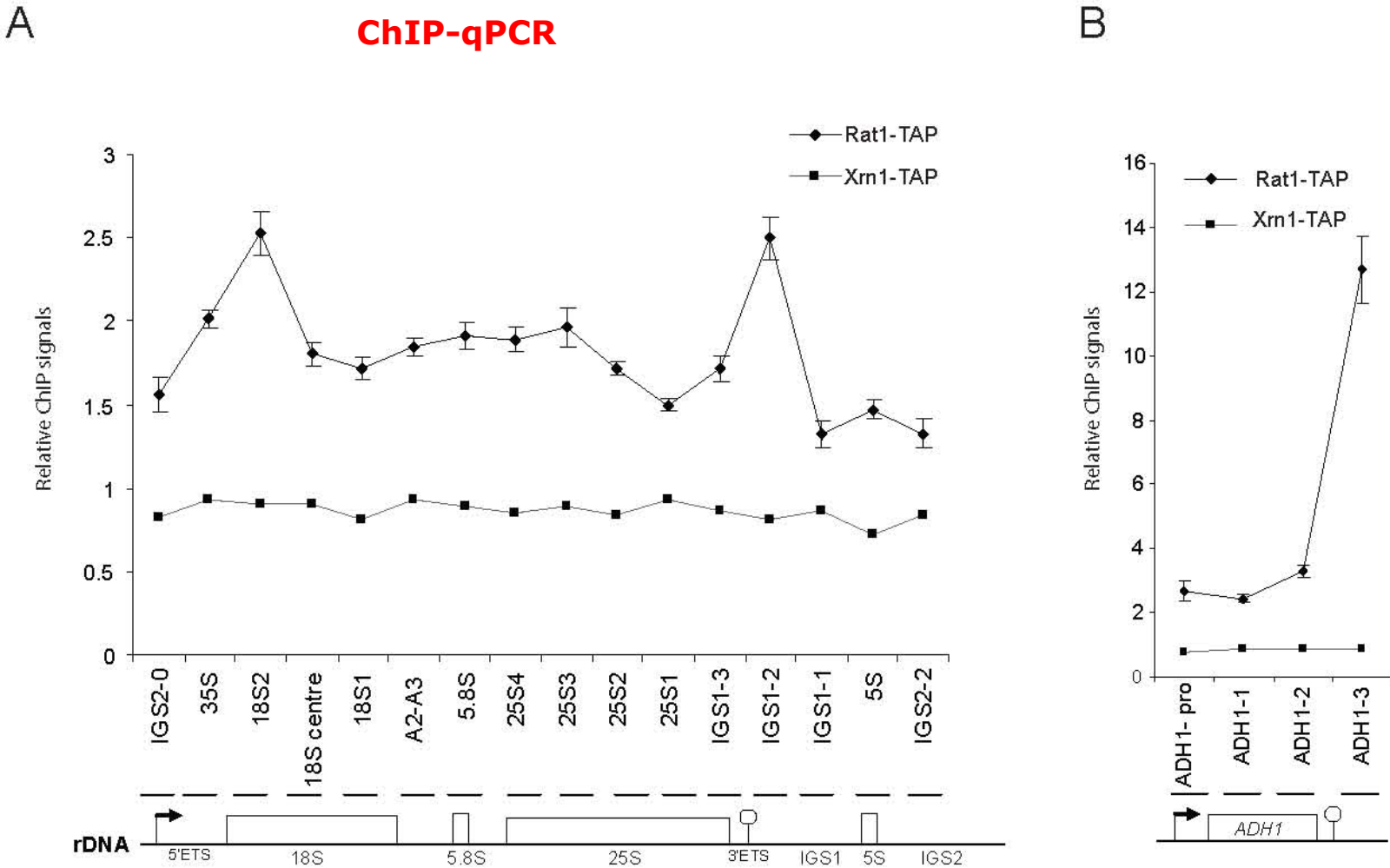
# Schematic of an rDNA repeat in *S. cerevisiae* with detailed transcription termination region



El Hage et al, Genes Dev., 2008,15;22(8)



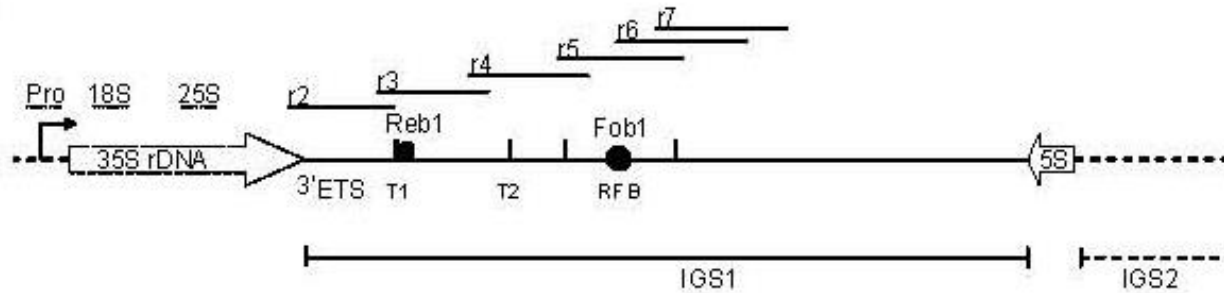
# Rat1 associates with the 3'-ETS region on the rDNA



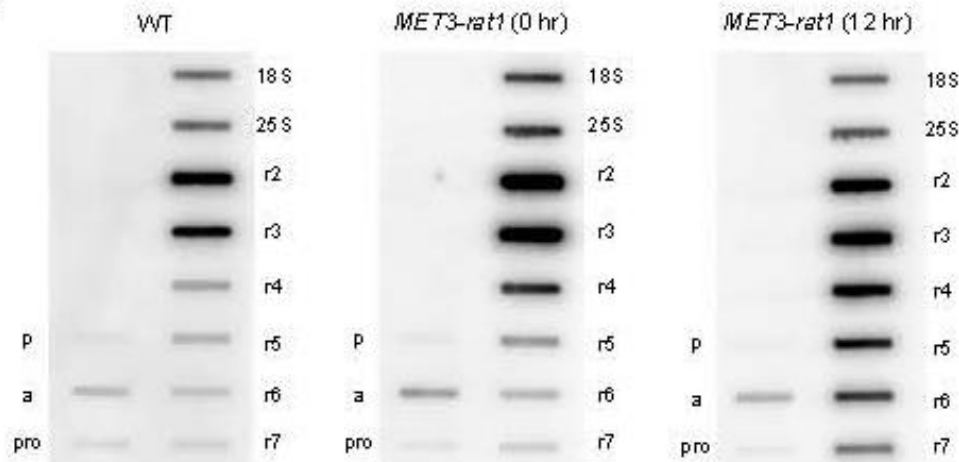
El Hage et al, Genes Dev., 2008,15;22(8)

# Transcription Run-On shows read-through of the major Pol I terminator in strains lacking Rat1 or Rai1

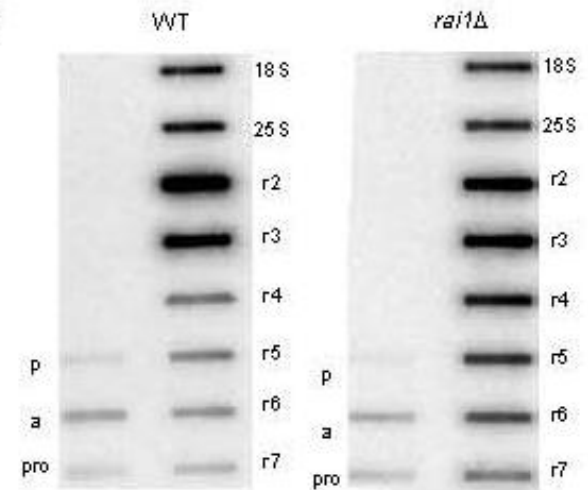
A



B



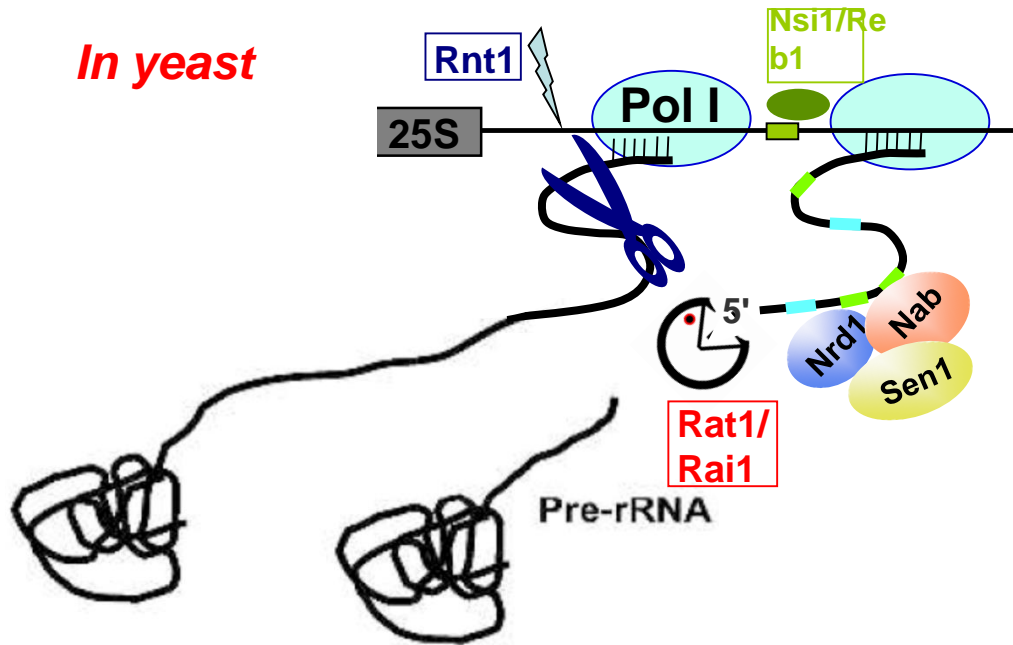
D



El Hage et al, Genes Dev., 2008,15;22(8)

# RNA polymerase I transcription termination

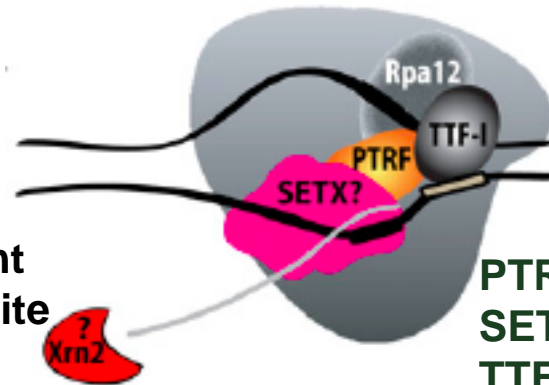
*In yeast*



Pol I termination factors:

- DNA-binding protein Nsi1
- Pol I subunit Rpa12
- endonuclease Rnt1
- RFB binding protein Fob1
- 5'-3' exonuclease Rat1/Rai1 (*torpedo mechanism*)
- RNA helicase Sen1
- Nrd1/Nab3 complex (??)

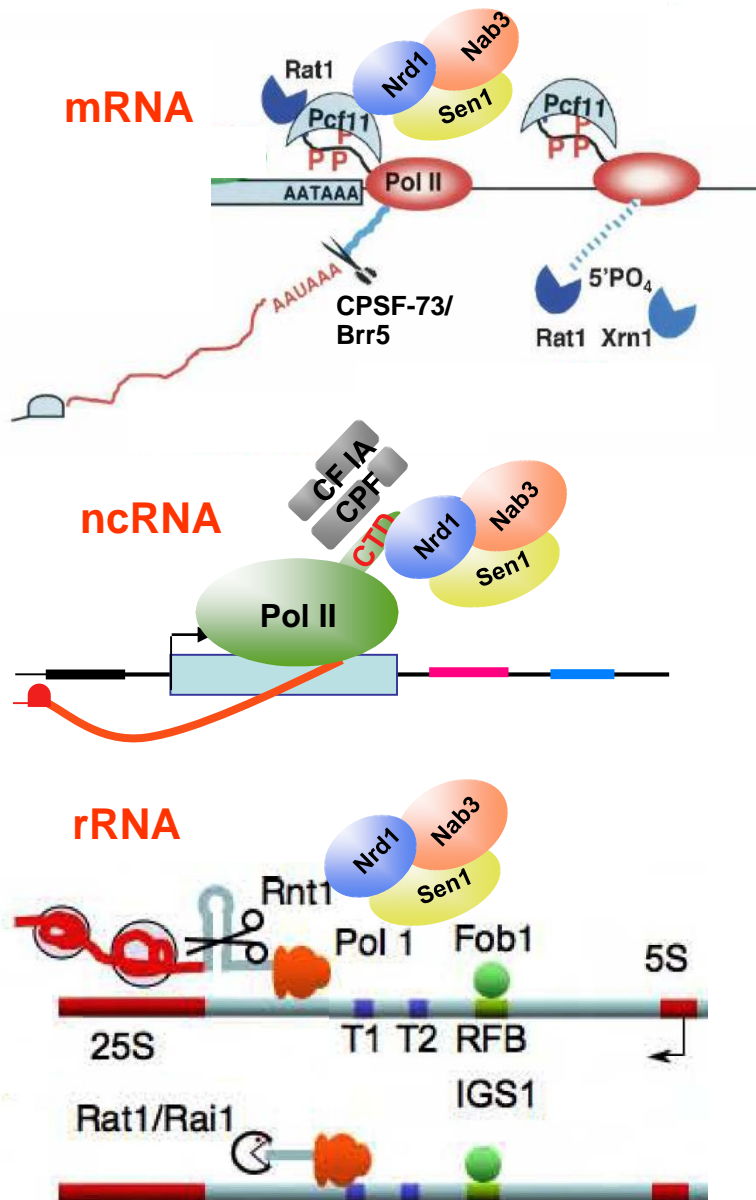
*In mammalian cells*



transcript release element  
T-stretch + TTF-I pause site

PTRF - release factor  
SETX - helicase, Sen1 homolog  
TTF-I - transcription termination factor I

# Conclusion - RNA pol I and RNA pol II „share” termination pathways



Pathways for transcription termination by different polymerases, Pol I and II, utilize overlapping factors and mechanisms

# SUMMARY OF TECHNIQUES

The level of RNA in a cell is the result of its synthesis and degradation.

The fact that RNA polymerase is bound to DNA does not mean that it transcribes RNA.

To study a transcript at the time of its formation, various molecular techniques should be used:

**TRO allows to study qualitatively and quantitatively the „ongoing“ transcription (to detect nascent transcripts).**

**ChIP makes it possible to study *in vivo* the interaction of proteins with DNA.**

**RIP allows to study *in vivo* the interaction of proteins with RNA.**

**THANK YOU  
FOR  
YOUR ATTENTION**