Transcription termination

Transcription termination

• Transcription initiates pervasively in all organisms.

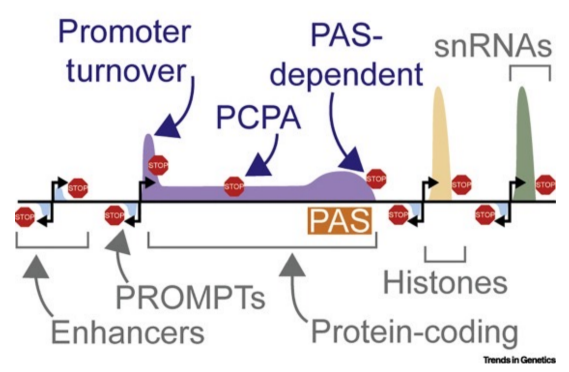
• Termination of transcription is essential for sorting out the functional RNAs from a plethora of transcriptional products that seemingly have no use in the cell.

• Terminating transcription is not that easy, given the high robustness of the elongation process.

Termination Occurs at All Points in the Transcription Cycle

Because Pol II transcribes multiple gene types, its termination occurs in a variety of ways

Where to terminate?



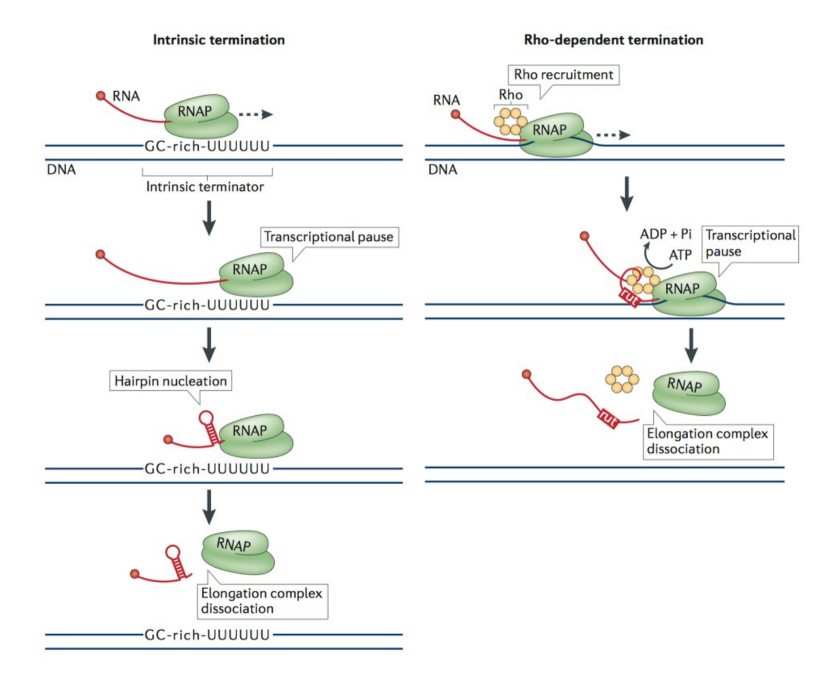
Abbreviations: PAS, polyadenylation signal; PCPA, premature cleavage and polyadenylation; PROMPT, promoter upstream transcript; snRNA, small nuclear RNA.

Transcripts and associated termination, processing and degradation pathways

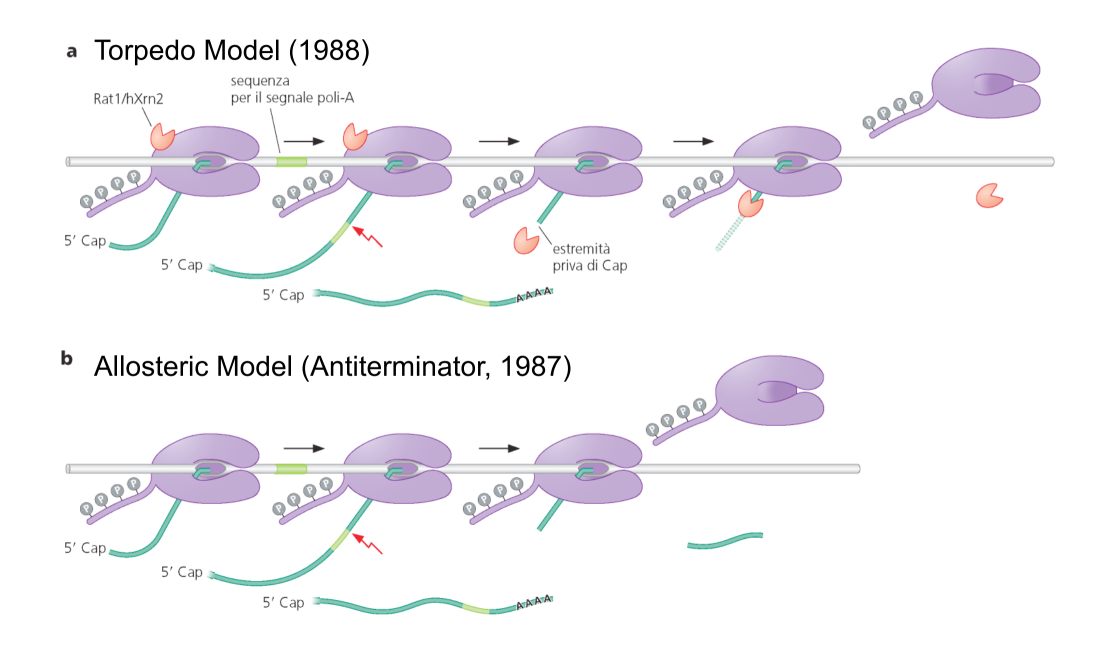
Transcript	Termination pathway	Stability	Degradation factors
Yeast			
mRNA	CPF–CF and possibly Sen1	Stable	None
snRNA and snoRNA	NNS	Stable (3' end processed)	TRAMP, Rrp6, exosome, Rex1 (3' end processing)
CUT	NNS	Unstable	TRAMP, Rrp6, exosome
SUT	CPF–CF and possibly NNS	Partially unstable	Rrp6, exosome, Xrn1 (NMD)
XUT	CPF-CF	Unstable	Xrn1 (NMD)
RUT	Reb1 roadblock	Unstable	TRAMP, Rrp6, exosome
Metazoan			
mRNA	CPSF-CF and SETX	Stable	None
snRNA	Integrator complex, CBC–ARS2, PCF11 and NELF	Stable (3' end processed)	Exosome (3' end processing)
Non productive transcription	Integrator complex	Unstable	Exosome (3' end processing)
mRNAs encoding replication-dependent histones	CBC-ARS2	Stable	None
PROMPT	CPSF-CF and CBC-ARS2	Unstable	NEXT and exosome

CBC, cap-binding complex; CF, cleavage factor; CPSF, cleavage and polyadenylation specificity factor; CUT, cryptic unstable transcript; NELF, negative elongation factor; NEXT, nuclear exosome targeting; NMD, nonsense-mediated decay; NNS, Nrd1–Nab3–Sen1; PROMPT, promoter-proximal transcript; RUT, Reb1-dependent unstable transcript; SETX, senataxin; snoRNA, small nucleolar RNA; snRNA, small nuclear RNA; SUT, stable unannotated transcript; TRAMP, Trf4–Air2–Mtr4; XUT, Xrn1-dependent unstable transcript.

Termination of transcription in bacteria

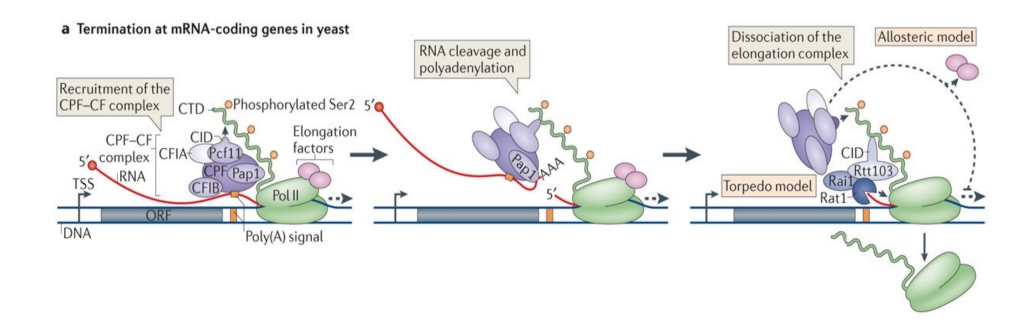


Models for PAS-Dependent Pol II Termination



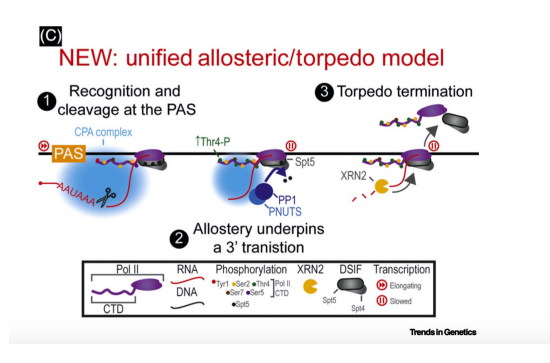
Termination of mRNA-coding genes: the CPF-CF pathway

3'-end processing and termination of mRNAs are triggered by multipartite signals on the nascent RNA, which are recognized by components of the CPF complex that also directly interacts with the polymerase. It is generally accepted that cleavage of the nascent transcripts occurs before release of the enzyme from the DNA template, which occurs further downstream. The biochemical details of the termination reaction are still unclear.



A Unified Model

Recent experiments provide strong evidence for a termination mechanism that unifies both original models. This is initiated by an allosteric switch that decelerates Pol II beyond the PAS – a step that was first uncovered in fission yeast where the protein phosphatase 1 (PP1) enzyme, Dis2, dephosphorylates the Spt5 elongation factor. Dis2-catalyzed dephosphorylation of Spt5 occurs after the PAS and is predicted to reduce Pol II speed, thus facilitating its termination by XRN2. Human PP1 enzyme and its nuclear targeting factor, PNUTS, are present in CPA complexes providing a connection between 3'-end processing and this termination mechanism.

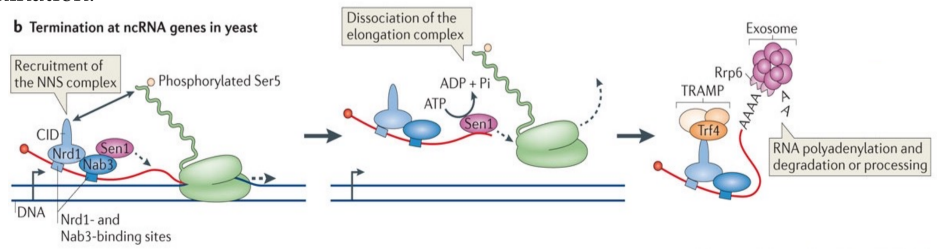


Termination of ncRNAs: the NNS-dependent pathway

In *S. cerevisiae*, the **Nrd1–Nab3–Sen1 (NNS) complex** is responsible for transcription termination at genes encoding **snRNAs** and **snoRNAs** and at **cryptic unstable transcripts (CUTs**). The essential NNS complex contains two RNA-binding proteins, **Nrd1 and Nab3**, and the conserved superfamily I RNA and DNA helicase **Sen1**. Cleavage of the primary transcript has never been demonstrated for this termination pathway, and release of the polymerase occurs by a mechanism that strictly requires the action of the helicase Sen1 (similar to bacterial Rho-dependent termination).

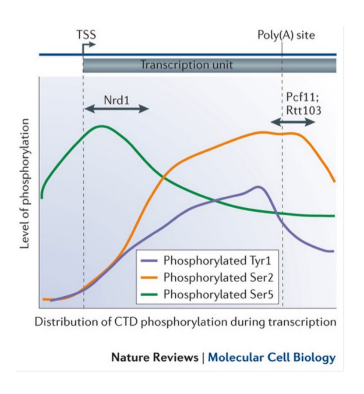
An important and distinctive trait of the transcripts produced by NNS-dependent termination is that <u>they are targeted by the nuclear exosome rapidly after their release.</u>

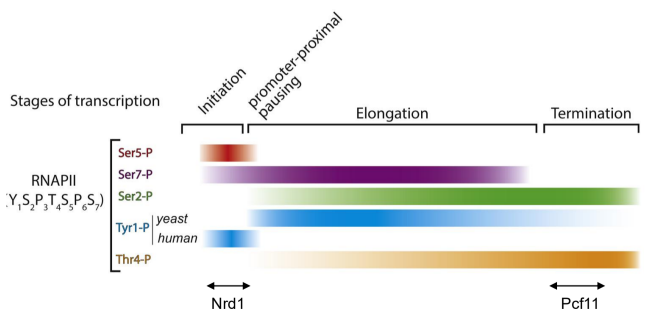
The presence of short sequence motifs on the nascent RNA that are recognized by Nrd1 and Nab3 (GUAA/G and UCUUG, respectively) has been shown to be a crucial specificity determinant of NNS-dependent termination. These motifs are often clustered and associated with AU-rich sequences, which contribute to the efficiency of termination.



The CTD of RNA Pol II and termination

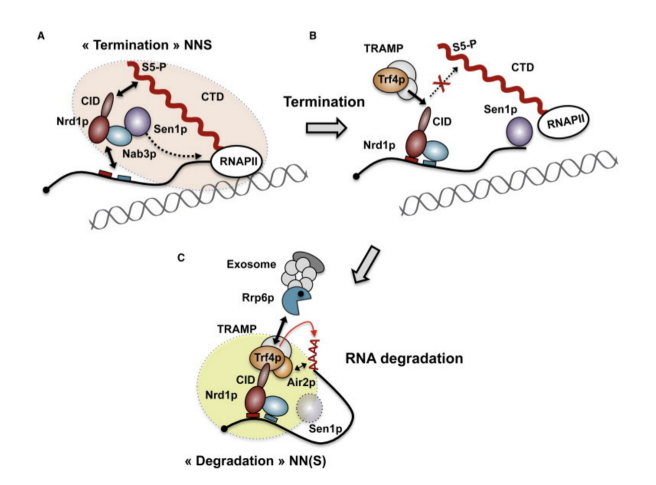
The most relevant modifications for transcription termination are <u>phosphorylation at Tyr1, Ser2, Ser5 and Ser7</u>, which alter the binding specificity of termination factors. For instance, in yeast the CPF complex component Pcf11 interact preferentially with the Ser2P form of the CTD only when Tyr1 is dephosphorylated. Conversely, Nrd1 recognizes the Ser5P form of the CTD, which predominates early in transcription, but only before the phosphorylation of Tyr1, which helps to restrict the recruitment of the NNS complex to the early stages of transcription.



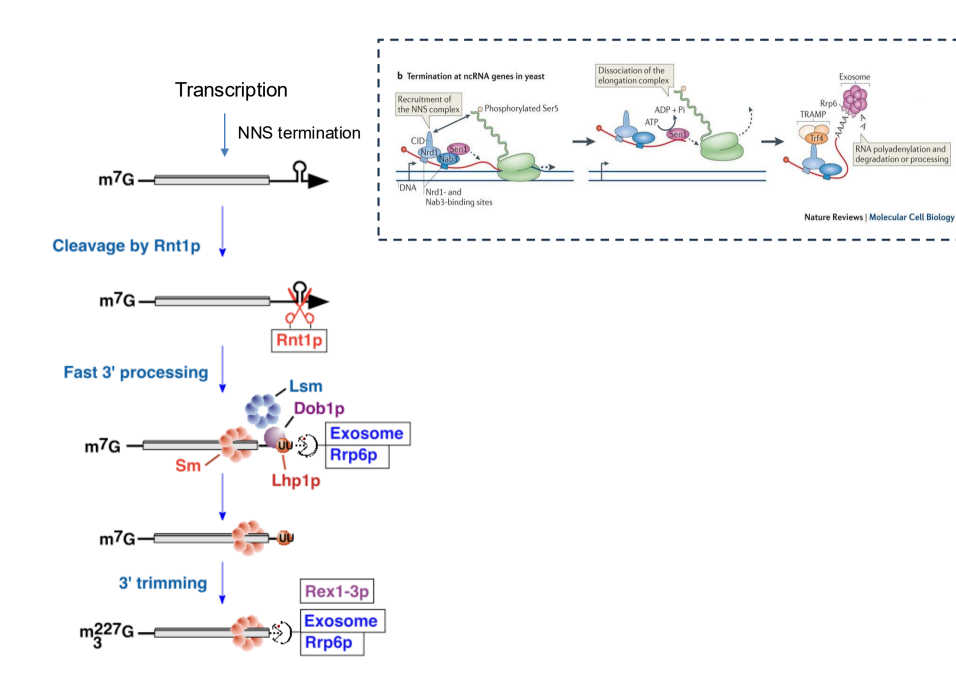


Model for the Coordination of Transcription Termination with RNA Degradation at NNS

After interacting with the Ser5P CTD for the termination step, Nrd1 recruits TRAMP through the direct recognition of a CTD mimic — known as the **Nrd1-interacting motif (NIM)** — in the TRAMP component Trf4. The sequential (and mutually exclusive) interaction of Nrd1 with the CTD and Trf4 contributes to the temporal coordination of termination with degradation.



Synthesis of the RNA Pol II-snRNAs

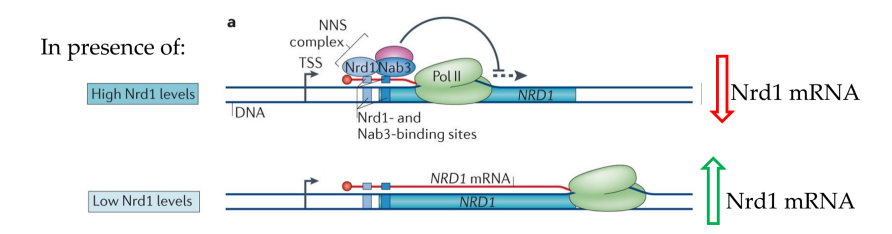


Regulation of gene expression by transcription termination

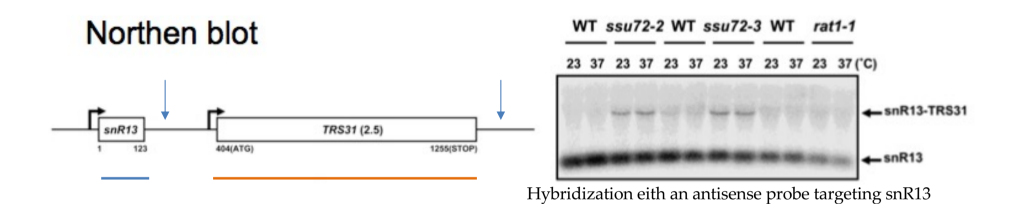
The occurrence of premature termination or termination that is associated with degradation of the transcript effectively prevents or limits gene expression. This can lead to bona fide regulation of gene expression, for example, when the occurrence of premature termination is modulated in response to an external stimulus or a physiological condition.

Nrd1:

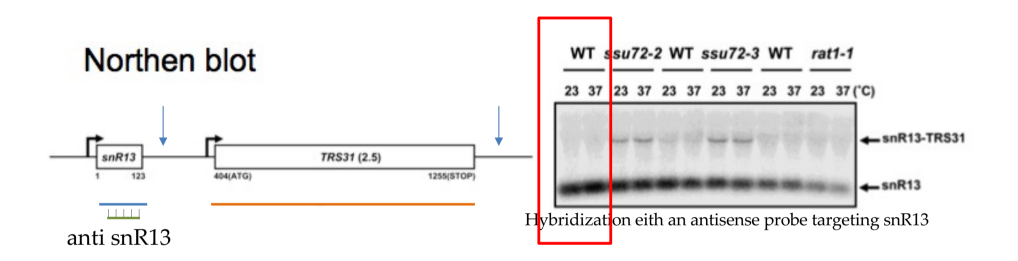
In yeast, the *NRD1* locus contains Nrd1- and Nab3-binding sites in the 5' region of the gene, and NNS-dependent termination occurs with a suboptimal efficiency that depends on the levels of the Nrd1 protein, thus establishing a negative feedback loop



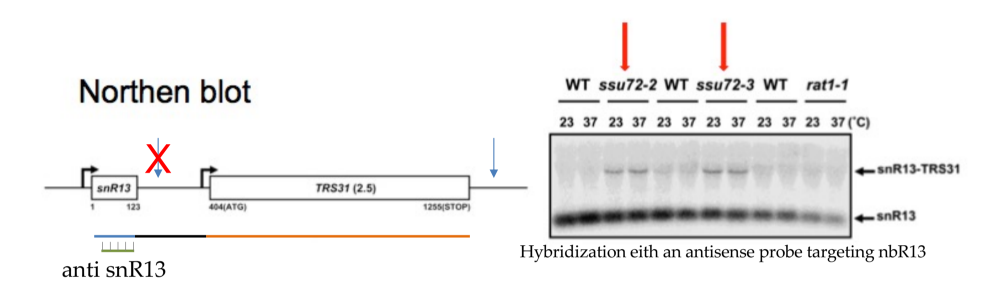
Experimental methodologies for transcription termination



Experimental methodologies for transcription termination

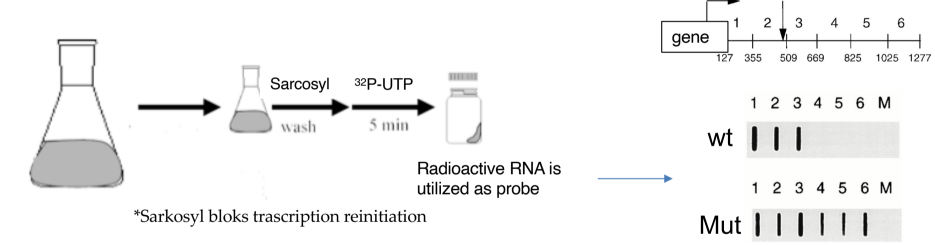


Experimental methodologies for transcription termination



Experimental methodologies for transcription termination

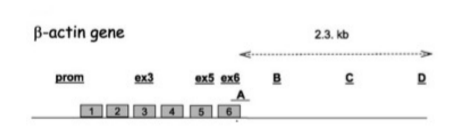
Transcriptional Run ON

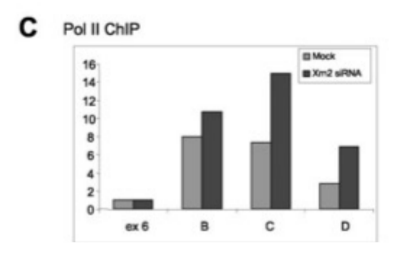


Antisense DNA oligos (1-6) are spotted on a membrane

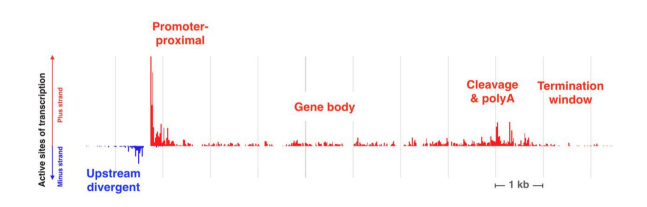
Experimental methodologies for termination

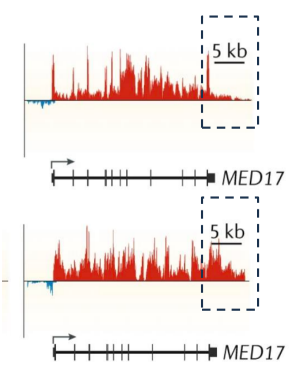
RNA Pol II ChIP





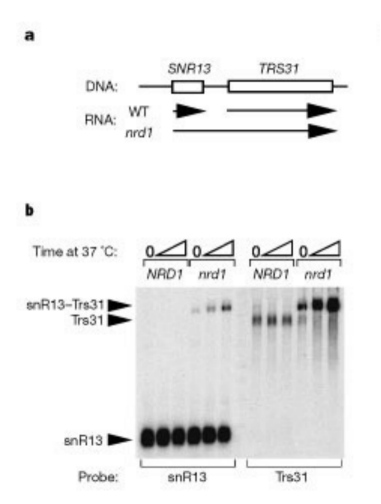
Nascent RNA-seq





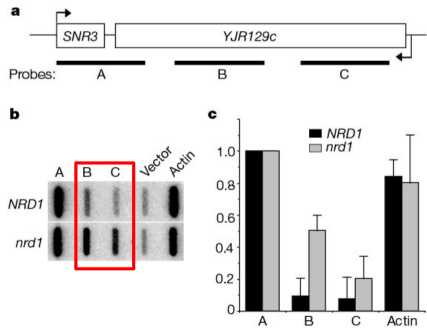
RNA-binding protein Nrd1 directs poly(A)-independent 3-end formation of RNA Pol II transcripts

• Direct evidence that ncRNAs might be natural targets for **Nrd1** was first obtained from expression profiling of poly(A)+ RNA (microarray) derived from temperature-sensitive *nrd1* yeast strains as compared with wild type. Several open reading frames (ORFs) exhibiting increased expression in the *nrd1* mutant strain are located downstream of snoRNA genes in the yeast genome.



RNA-binding protein Nrd1 directs poly(A)-independent 3-end formation of RNA Pol II transcripts

A significant proportion of polymerase molecules that terminate transcription downstream of the *snoRNA* coding region in the wild-type strain fail to do so in the *nrd1* mutant



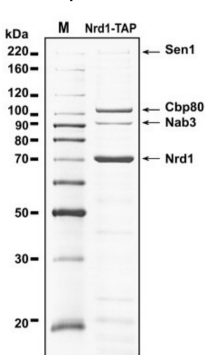
In contrast to SNR13. the downstream ORF is convergent orientation. therefore single-stranded probes for SNR3 read-through will transcripts not **mRNA** hvbridize with from downstream **ORF**

transcription run-on analysis

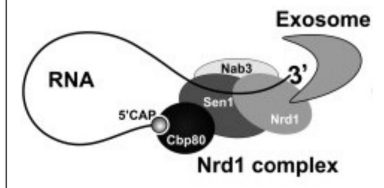
The Sen1-Nab3-Nrd1 complex

The alternative Pol II termination pathway for non-coding RNAs in yeast is composed by the RNA-binding proteins **Nrd1** and **Nab3**, and the RNA and DNA helicase **Sen1**.



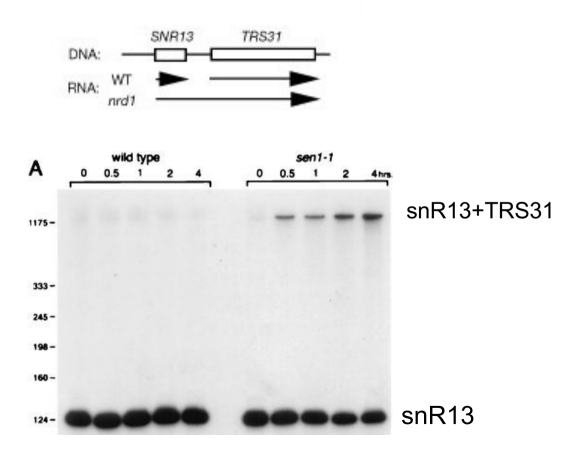


	Protein	Number of peptides	Protein function
Nrd1 complex	Sen1	57	RNA helicase, sn/snoRNA processing
Ĭ,	Nab3	42	sn/snoRNA termination
2	Cbp80	32	Cap-binding
N P	Cbp20	5	Cap-binding
	Nrd1	27	sn/snoRNA termination
RNA pol II complex	Spt5	24	transcription factor
	Rpb1	24	RNApol II subunit
	Rpb2	19	RNApol II subunit
	Rpb3	9	RNApol II subunit
	Rpb5	3	RNApol II subunit
	Rpb4	2	RNApol II subunit
RNA processing complex	Rrp6	15	nuclear exosome
	Rrp44	5	exosome core
	Rrp4, Csl4, Rrp45, Rrp		exosome core
	Air2	9	Trf4 cofactor
	Trf4	7	poly(A) polymerase
	Pab1	5	poly(A) binding
	Rnt1	3	sno/snRNA processing, endonuclease



The Sen1-Nab3-Nrd1 complex

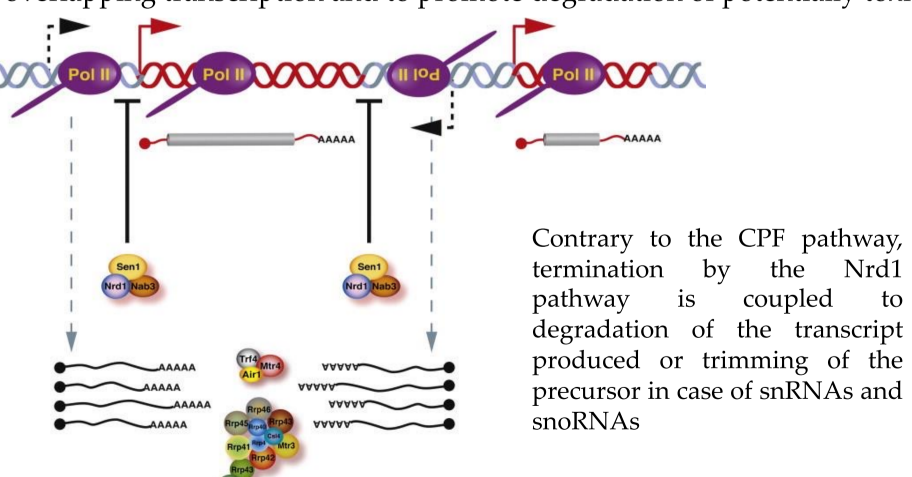
The alternative Pol II termination pathway for non-coding RNAs in yeast is composed by the RNA-binding proteins **Nrd1** and **Nab3**, and the RNA and DNA helicase **Sen1**



Directing transcription to the right way

Yeast

Transcription from regions upstream of or antisense to mRNA-coding genes (red DNA) is terminated by the Nrd1p complex and transcript are rapidly degraded. Quality control by the Nrd1p complex operates to avoid overlapping transcription and to promote degradation of potentially toxic.

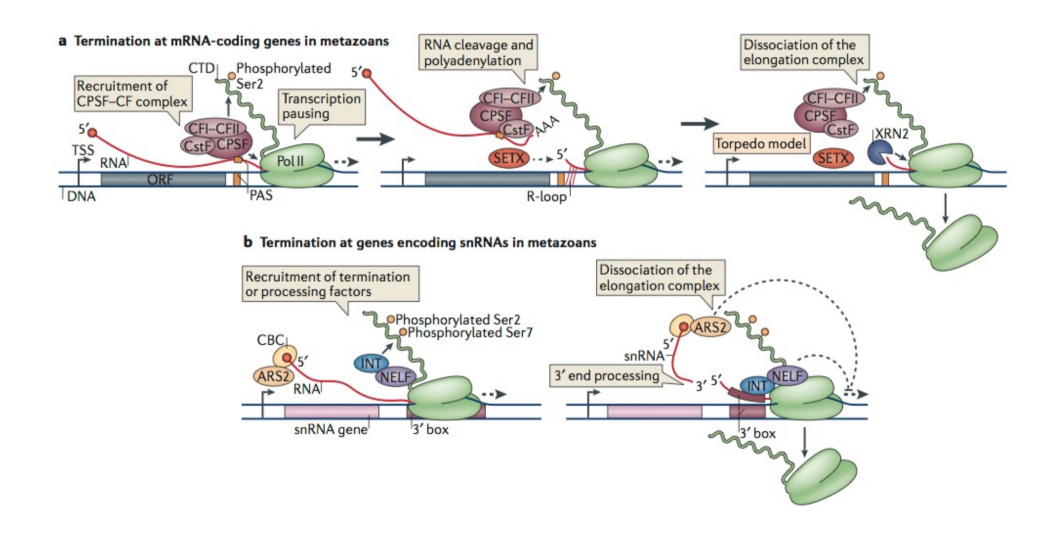


Transcription termination in metazoans

Three pathways of Pol II transcription termination have been described in metazoans, generating:

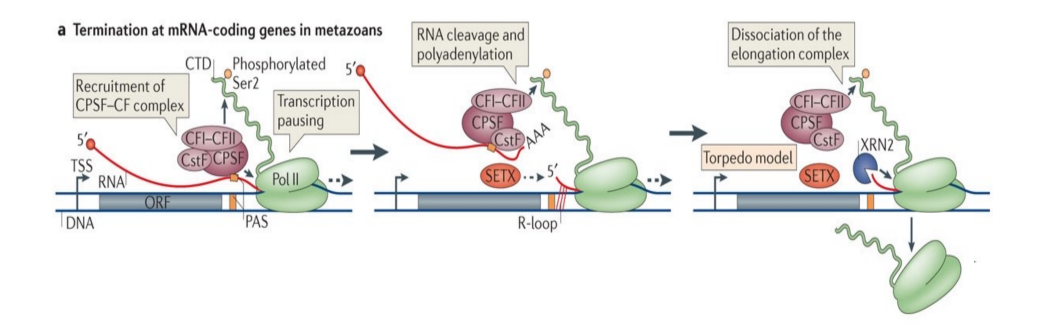
- 1. mRNAs,
- 2. snRNAs
- 3. transcripts encoding replication-dependent histones.

Transcription termination in metazoans

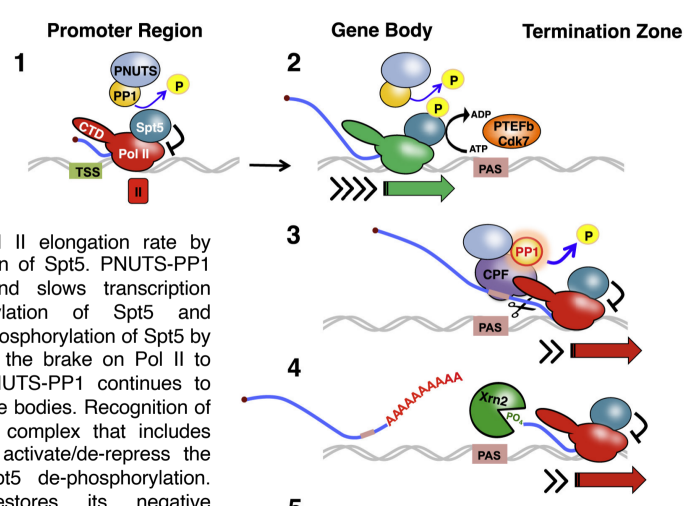


General termination of mRNAs in metazoans

Cleavage of the nascent transcripts by the CPSF component CPSF73 (also known as CPSF3) occurs 18–30 nucleotides downstream of a polyadenylation signal (PAS; AAUAAA). It is commonly accepted that the PAS is required to trigger termination; A role for **senataxin** (**SETX**) in transcription termination of mRNAs has been proposed in several reports.



The Sitting Duck Torpedo Model of Pol II Transcription Termination

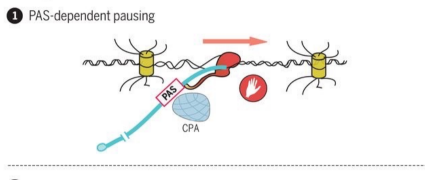


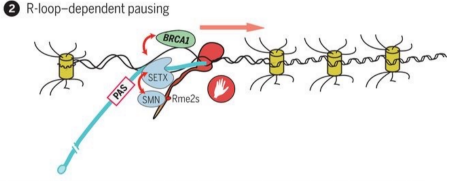
Model for the control of Pol II elongation rate by PNUTS-PP1 dephosphorylation of Spt5. PNUTS-PP1 is recruited to 50 ends and slows transcription elongation by dephosphorylation of Spt5 and potentially other substrates. Phosphorylation of Spt5 by PTEFb and/or Cdk7 releases the brake on Pol II to accelerate elongation but PNUTS-PP1 continues to limit elongation rate within gene bodies. Recognition of the poly(A) site (PAS) by a complex that includes PNUTS-PP1 is proposed to activate/de-repress the phosphatase resulting in Spt5 de-phosphorylation. dephosphorylation restores Spt5 its negative elongation activity that causes Pol II to decelerate. Slow elongating Pol II is an easy target or "sitting duck" for the torpedo exonuclease Xrn2 that chases down and helps dissociate Pol II from the DNA template.

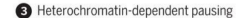
General termination of mRNAs in metazoans

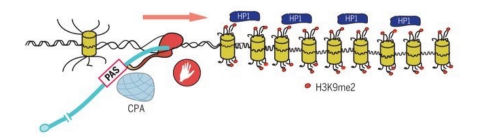
Transcription termination in metazoans is thought to be associated with Pol II pausing.

A Pausing









Three different types of **Pol II pausing:**

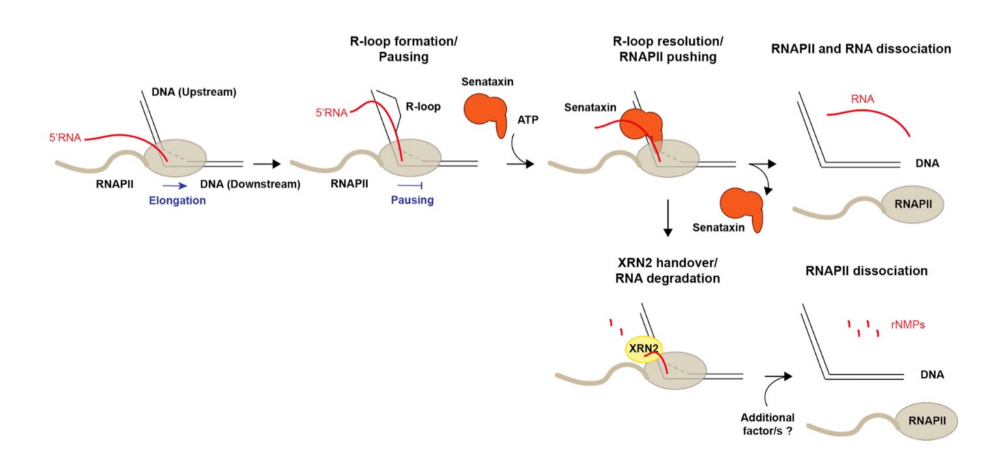
- 1. induced by CPA recognition of the PAS
- 2. R-loop formation
- 3. heterochromatin patches

Elongating Pol II (red) is shown transcribing the DNA template, with extruded, capped RNA transcript (blue) indicated. Nucleosomes are depicted by yellow barrels, with histone N-terminal tails indicated. Pol II CTD is shown as an extended tail. Red dots on the CTD and histone tails denote methylation. The hand denotes Pol II pausing.



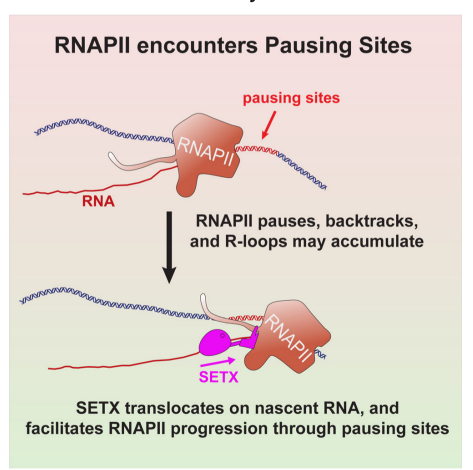
Senataxin is involved in RNA-Pol II transcription termination

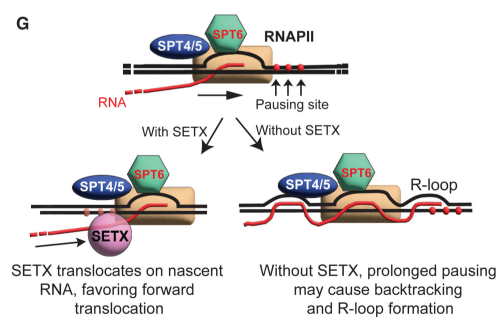
Nascent transcripts form **RNA/DNA hybrid structures (R-loops)** behind elongating Pol II and are especially prevalent over G-rich pause sites positioned downstream of gene poly(A) signals. **Senataxin**, a helicase protein associated with neurodegenerative disorders (*Ataxia-Oculomotor Apraxia 2 and amyotrophic lateral sclerosis 4*) acts to resolve these R-loop structures thus, facilitating termination.



SETX facilitates RNAPII progression through pausing sites

Recently experiments employing a degradation tag system for acute SETX depletion demonstrated that SETX loss perturbs RNAP II elongation but does not markedly influence transcription termination at the end of genes.





R-loops arising more slowly over several hours after losing SETX, likely as a secondary effect explaining previously reported effects of SETX depletion on termination

R loops in human disease

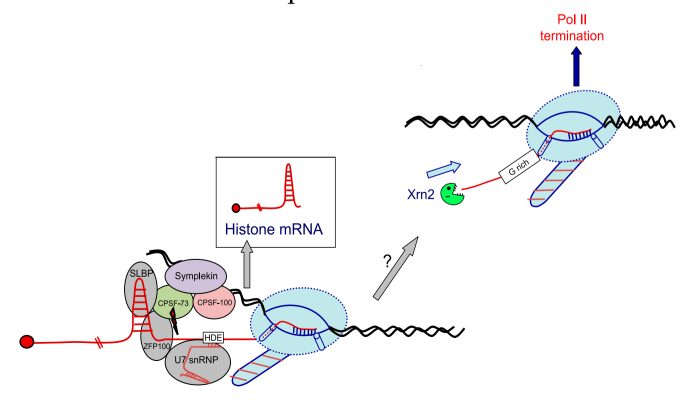
rable 2 Genes retated to K-toop metabolism that can cause number diseases in dystanctional	Table 2 Genes related to	R-loop metabolism that can cause h	numan diseases if dysfunctional
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Gene	Disease	Cause	Refs			
SETX	Ataxia-ocular apraxia type 2 (AOA2) and amyotrophic lateral sclerosis type 4 (ALS4)	Mutations in the RNA-DNA helicase SETX	102,103			
FXN	Friedreich ataxia (FRDA)	Expansion of GAA repeats in FXN gene promotes R-loop formation, H3K9me2 and decreased FXN expression	93,98			
FMR1	Fragile X syndrome (FXS) and fragile X-associated tremor/ataxia syndrome (FXTAS)	Expansion of CGG repeats in FMR1 gene promotes R-loop formation, H3K9me2 and decreased FMR1 expression	93,99, 100			
C9orf72	Amyotrophic lateral sclerosis (ALS) and frontotemporal dementia (FTD)	Expansion of GGGCC repeats causes R-loop formation and accumulation of aborted transcripts	101			
BRCA1	Cancer	Genome instability caused by R-loop accumulation in BRCA1-deficient cells	80,95, 108			
BRCA2	Cancer and Fanconi anaemia (FA)	Genome instability caused by R-loop accumulation in BRCA2-deficient cells	80			
FIP1L1	Cancer	Genome instability caused by R-loop accumulation in FIP1L1-deficient cells inferred by the yeast mutant $\mathit{fip1}\Delta$	21			
BRE1	Cancer	Genome instability caused by R-loop accumulation in BRE1-deficient cells	111			
SRSF1	Cancer	Deregulation of cancer-associated genes due to SRSF1 overexpression	112			
ORF57	Kaposi sarcoma-associated herpesvirus (KSHV)	Sequestration of human TREX complex by ORF57 causes R-loop formation and DNA damage	113			

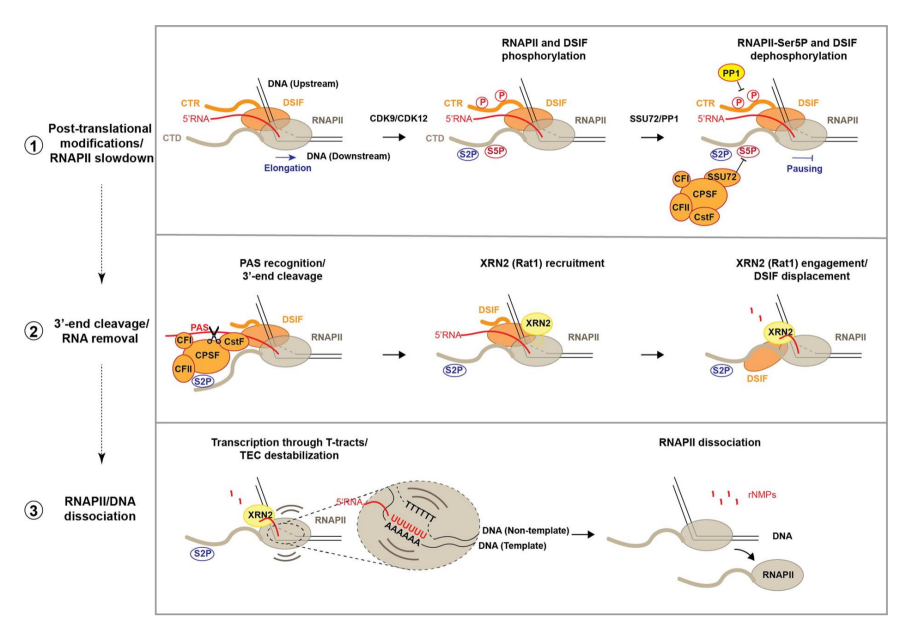
C9orf72, chromosome 9 open reading frame 72; FIP1L1, factor interacting with PAPOLA and CPSF1; FMR1, fragile X mental retardation 1; FXN, frataxin; H3K9me2, histone H3 lysine 9 dimethylation; SETX, senataxin; SRSF1, serine/arginine-rich splicing factor 1.

Transcription termination at replication-dependent histones mRNAs

In contrast to many mRNAs, transcripts encoding replication-dependent histones are not polyadenylated but rather undergo cleavage at a particular stem-loop structure. U7 small nuclear ribonucleoprotein (snRNP), CBC, NELF, ARS2 and CPSF factors, including the CPSF73 endonuclease, have been shown to be involved in termination. A torpedo model has been proposed in which the Exonuclease (Xrn2) entry site is produced by CPSF-73 recruited by the U7/SLBP/ZFP100 complex.



Model of RNAPII 3'-end termination



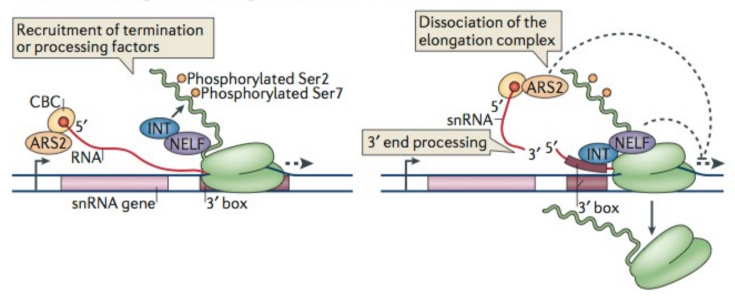
Legend in the next slide

Model of RNAPII 3'-end termination

- 1. During elongation, both the RNAPII-CTD and SPT5-CTR (part of DSIF with SPT4) are phosphorylated by transcriptional CDKs to ensure the processivity of the elongation complex. This process is reversed by specific phosphatases, such as **PP1** and **SSU72** during termination to slowdown RNAPII and generate Ser2-P RNAPII-CTD.
- 2. Ser2-P on RNAPII-CTD and RNA sequence signals, such as the PAS, are specifically recognized by the 3'-cleavage machinery. Following 3'-cleavage, a free 5'-end is generated in the nascent pre-mRNA associated with transcribing RNAPII, which can be targeted by XRN2. XRN2 is first recruited to RNAPII through interactions with RPB2 and then is fully engaged through contacts with the RNAPII stalk and the RNA exit channel.
- 3. While docked on RNAPII, XRN2 can degrade and pull the RNA out from RNAPII. Finally, transcription through specific terminator DNA regions rich in T-tracts generate rU/dA hybrids (R-loops) inside of RNAPII, which together with downstream DNA sequences weaken the RNAPII/DNA interaction, destabilizing the TEC (transcription elongation complex) and leading to efficient RNAPII dissociation from the DNA.

Transcription termination in metazoans

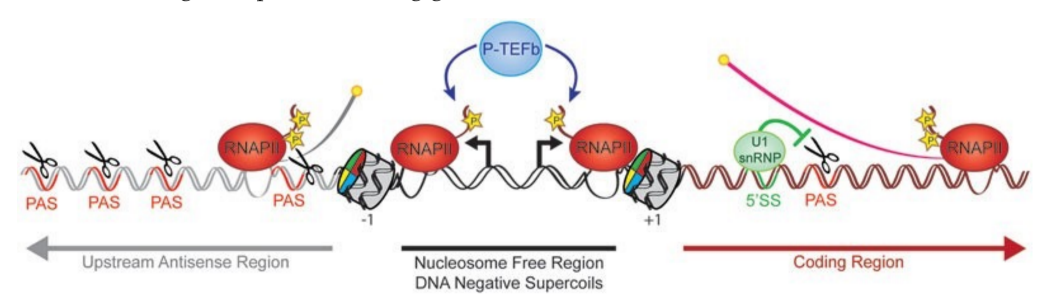
b Termination at genes encoding snRNAs in metazoans



Directing transcription to the right way

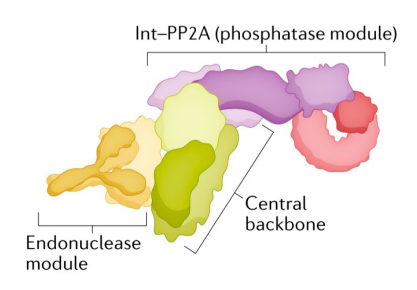
Mammals

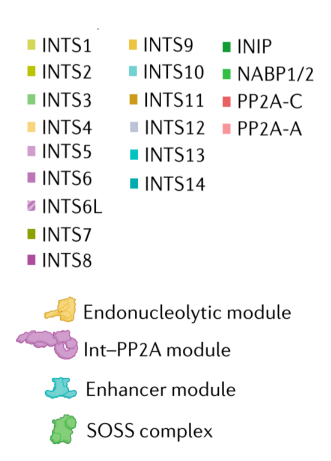
RNAPIIs are initiated in two opposite orientations from an active divergent promoter, and paused at the–1 and +1 nucleosomes, respectively. The paused RNAPIIs reinforce the boundaries of NFR and maintain the accessibility of promoter to transcription factors. P-TEFb is recruited to both upstream and downstream paused RNAPIIs and releases them from the pause sites. The elongation by RNAPIIs produces negative supercoils in the promoter region, facilitating further rounds of initiation. Shortly after released from the pause site, the upstream RNAPII transcribes through the PAS, and the transcription is terminated by the canonical 3'-end-processing factors (Left; upstream antisense region). In the downstream coding region (Right), the PAS is suppressed by the U1 snRNP recruited by 5'SS, and therefore productive elongation occurs through the protein-coding gene.



The integrator (Int) complex

Integrator was discovered in 2005 as a new multisubunit complex in human cells capable of binding the Pol II CTD. Orthologues of all subunits were identified throughout metazoans, but not in yeast, suggesting that the complex is unique to multicellular eukaryotes. Sequence homology revealed that Integrator complex subunit 9 (INTS9) and INTS11 are highly homologous to cleavage and polyadenylation specificity factor subunit 73 (CPSF73) and CPSF100, providing the first hint that these INT subunits may be endowed with RNA endonuclease activity. Depletion of either the largest subunit (INTS1) or the putative catalytic core (INTS11) of Integrator resulted in specific accumulation of unprocessed, precursor snRNAs.





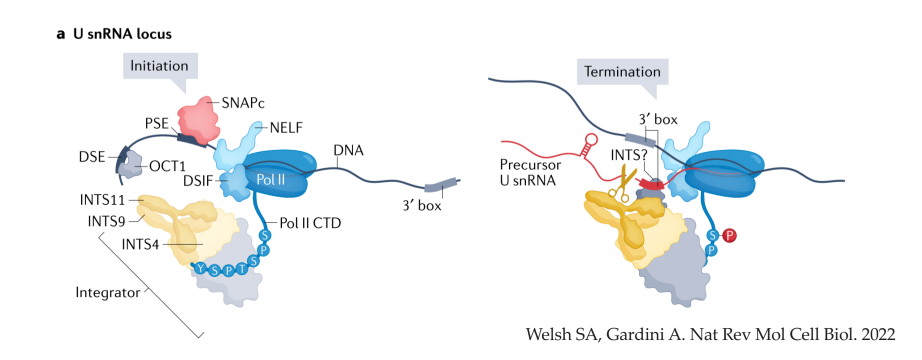
The functions of integrator at different non-coding RNAs

RNA species	Function of Integrator	Cell type, organism	Refs.
Short RNAs (<0.2 kb)			
U snRNAs	3' box-mediated cleavage	All cell types, various metazoans	21,23, 25,59,63
Viral miRNAs	Transcript release and maturation	Infected lymphocytes, marmosets	83,84
piRNAs	3' cleavage (unknown motif)	Germ cells, Caenorhabditis elegans	87,88
Long RNAs (>0.2 kb)			
eRNAs	3' cleavage (at unknown motif)	Multiple cell types, humans	58,81,82
TERC	3' cleavage (at unknown motif)	Human cell lines	92
NEAT1	Support of early transcription termination	Human cell lines	94
Other IncRNAs	Support of early transcription termination	Human cell lines	82

eRNA, enhancer RNA; IncRNA, long non-coding RNA; miRNA, microRNA; NEAT1, nuclear paraspeckle assembly transcript 1; piRNA, PIWI-interacting RNA; snRNA, small nuclear RNA; TERC, telomerase RNA template component.

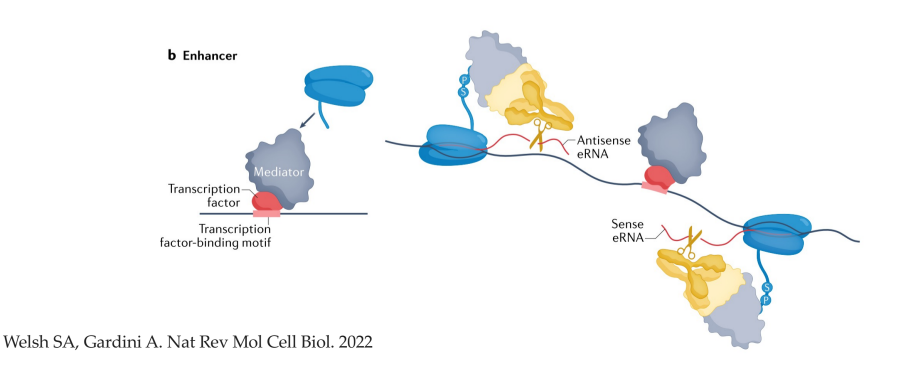
Integrator terminates transcription of snRNAs

snRNA loci have distinctive promoter elements that recruit the transcription initiation-competent RNA polymerase II (Pol II) holoenzyme including DSIF, NELF and the **Integrator complex**. Shortly after transcribing through the 3' box (13-16 nts), which is a highly conserved motif at the termination site of all U snRNAs, Integrator cleaves the nascent small RNA, triggered by phosphorylation (P) of Ser7 of the carboxy-terminal domain (CTD) of Pol II's largest subunit (RBP1). A 3' stem–loop in the precursor U snRNA and recognition of the ensuing 3' box RNA sequence by a set of Integrator accessory subunits (INTS?) may support an efficient cleavage process



Integrator terminates transcription of eRNAs

Enhancer loci are activated by sequence-specific transcription factors that recruit the co-activator Mediator complex. Upon Mediator recruitment and assembly of the transcription pre-initiation complex, bidirectional transcription of the enhancer locus occurs, producing long (>200-bp) sense and antisense transcripts called 'enhancer RNAs' (eRNAs). Both sense- transcribing and antisense-transcribing Pol II holoenzymes recruit the Integrator complex to terminate transcription and release eRNAs without eliciting their polyadenylation.

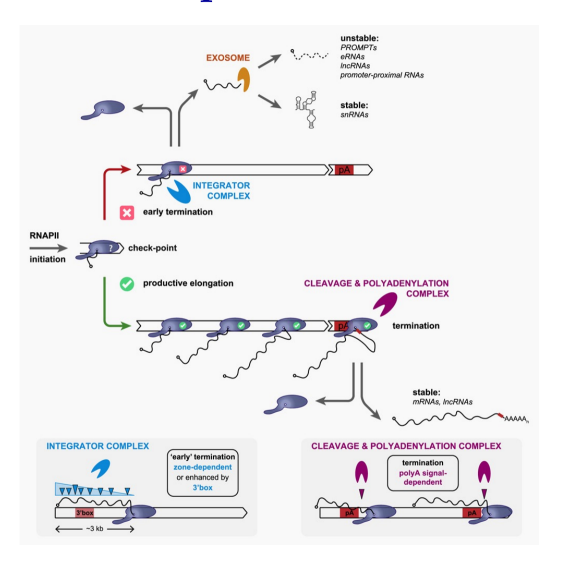


Integrator (INT) is a genome-wide attenuator of non-productive transcription

Two functions of genome-wide INT activity:

- 1. it dampens transcriptional output from weak promoters,
- 2. it provides quality control of RNAPII complexes that are unfavorably configured for transcriptional elongation.

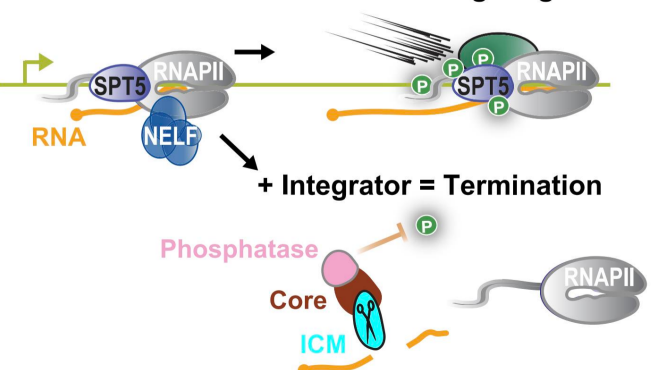
The function of INT in stable snRNA production is an exception from its general cellular role, the attenuation of non-productive transcription.



The integrator (Int) complex

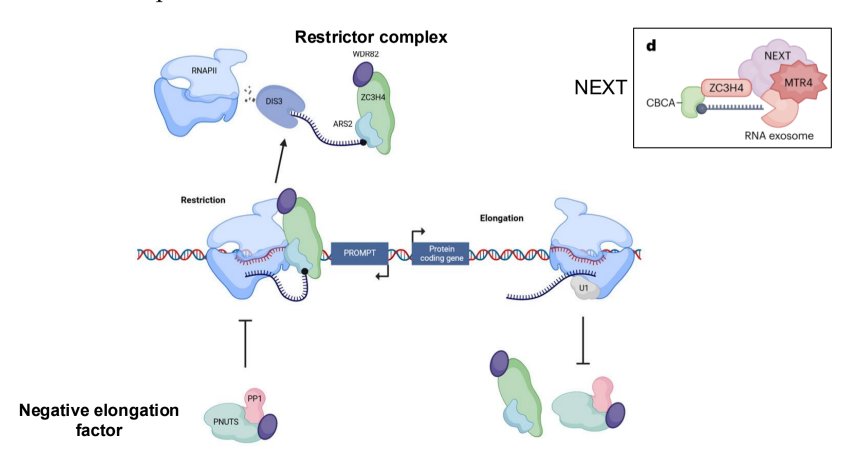
In addition to the endonuclease module, Integrator contains a module with phosphatase activity, the Int-PP2A phosphatase module. This module was shown to antagonize transcriptional kinases (CDK7) to suppress pause release and transcription elongation. Given the intimate contacts of Integrator subunits with paused RNAPII, it is not surprising that all Integrator subunits analyzed to date by ChIP-seq display enrichment just downstream of transcription start sites (TSSs). The Integrator complex is a major effector of promoter-proximal termination.

Paused RNAPII + P-TEFb = Elongating RNAPII



RESTRICTOR complex is a genome-wide attenuator of non-productive transcription

The restrictor ZC3H4/WDR82, an additional mediator of transcriptional quality control, binds to the CBCA complex and terminates antisense transcription from bidirectional promoters. It associates with the PP1-PNUTS complex, which induces RNA Pol II CTD Ser5 dephosphorylation. This promote termination by increasing RNA Pol II pausing. Longer protein-coding transcription is supported by U1 snRNA, which shields transcripts from restrictor and PNUTS.



RNAPII must evade early termination to transcribe long mammalian proteincoding genes.

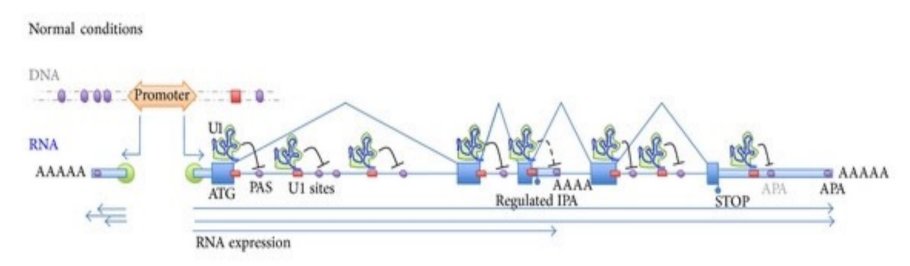
RNAPII escape from the promoter-proximal pause requires CDK9, and this antagonizes Integrator-dependent attenuation. Beyond this pause, U1 small nuclear ribonucleoprotein (snRNP) complexes play a major role in promoting elongation through long protein-coding genes.

This so-called "**telescripting**" function was first shown to suppress premature cleavage and polyadenylation (PCPA) within introns, thereby preventing early poly(A) signal (PAS)-dependent termination.

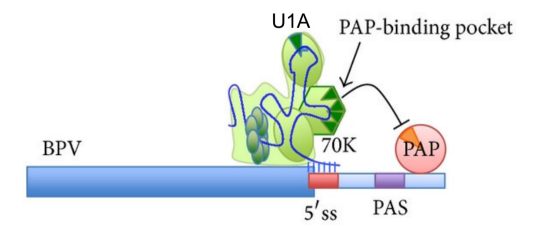
Unlike rapidly terminated ncRNAs, which typically lack 5' splice sites, the near-universal presence of introns in protein-coding RNAs likely explains why U1 selectively promotes their elongation. The mechanism by which U1 snRNP promotes telescripting/elongation remains incompletely understood. However, the U1 snRNP subunits U1A and U1-70K inhibit cleavage and polyadenylation (CPA). Notably, both factors cross-link to PCPA sites *in vivo*, along with CPA machinery components. This suggests that U1 recruitment to 5' splice sites positions U1A and U1-70K close to CPA factors, thereby preventing PCPA.

U1 is also suggested to prevent RNA cleavage by Integrator, and it opposes Restrictor at some protein-coding genes.

U1 snRNP suppression of cleavage and polyadenylation safeguards transcriptome integrity

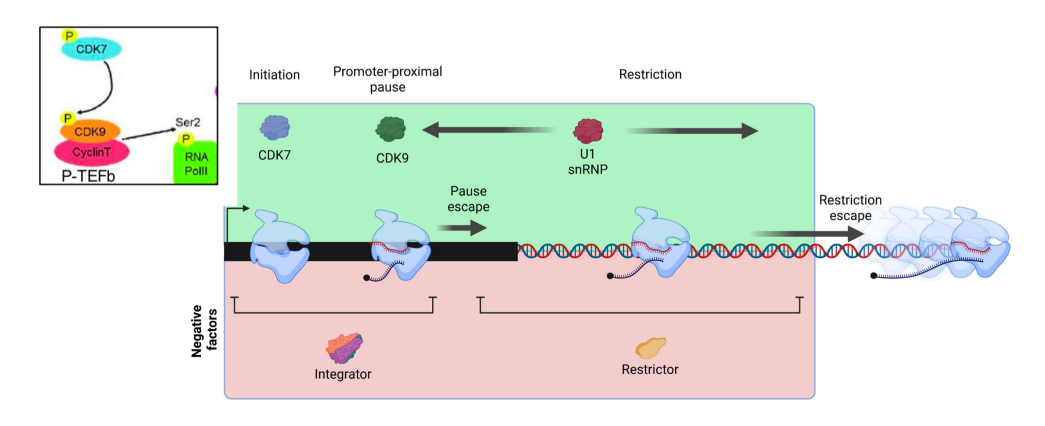


The U1 interacting protein U1-70K and in some context also U1A inhibit the PAS usage throught direct PAP inhibition

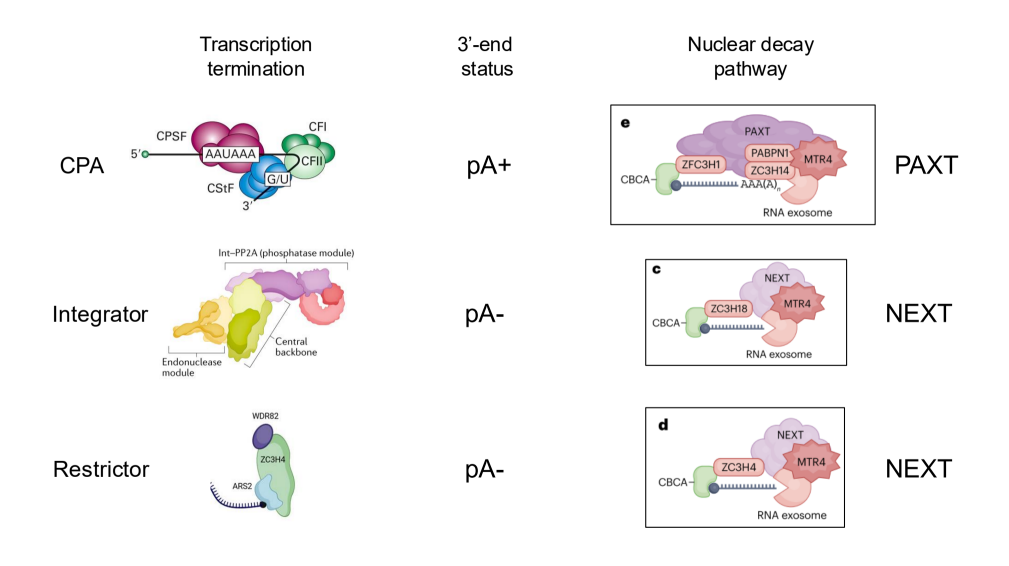


Integrator and Restrictor complexes are implicated in promoter-proximal termination

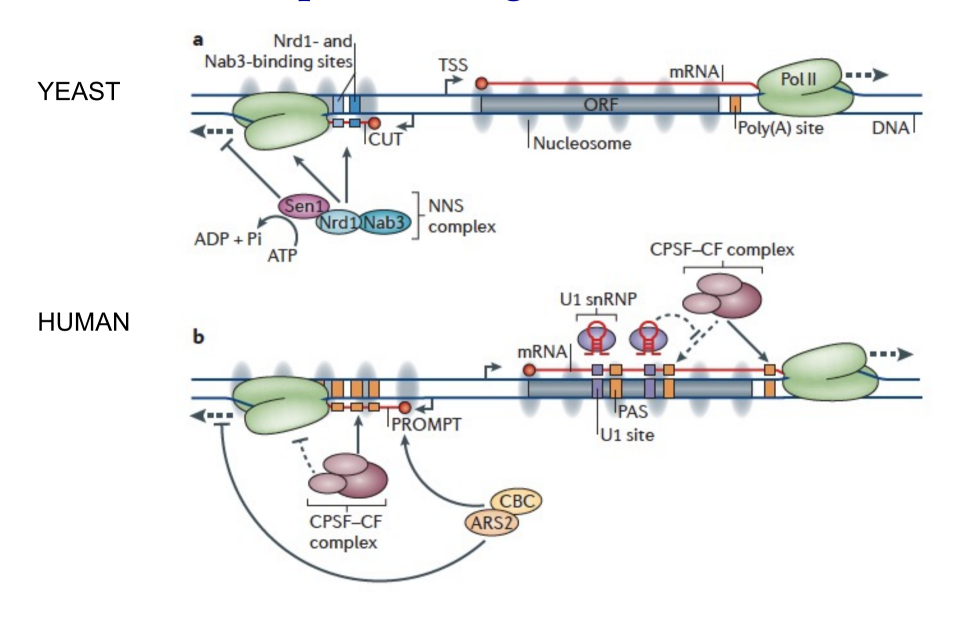
Integrator and Restrictor act sequentially to monitor distinct stages of transcription. Integrator predominantly engages with promoter-proximally paused RNAPII to trigger premature termination, which is prevented by the P-TEFb components cyclin-dependent kinase 9 (CDK9) and CDK7, which activates CDK9. After pause release, RNAPII enters a "restriction zone", universally imposed by Restrictor. Unproductive RNAPII terminates within this zone, while progression through it is promoted by U1 snRNPs, which antagonize Integrator and Restrictor.



Early Transcription termination and Nuclear decay



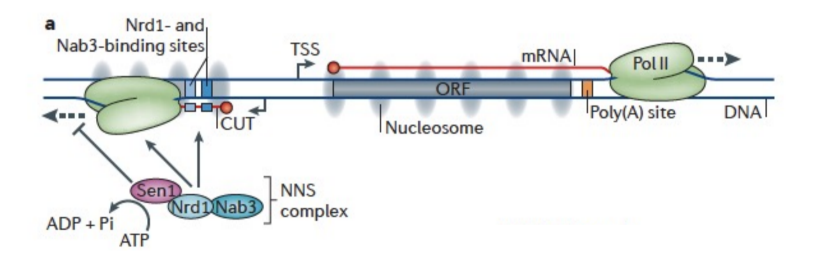
Transcription termination in the control of pervasive transcription throughout evolution



Directing transcription to the right way

Yeast

Transcription from regions upstream of or antisense to mRNA-coding genes (red DNA) is terminated by the Nrd1p complex and transcript are rapidly degraded. Quality control by the Nrd1p complex operates to avoid overlapping transcription and to promote degradation of potentially toxic.

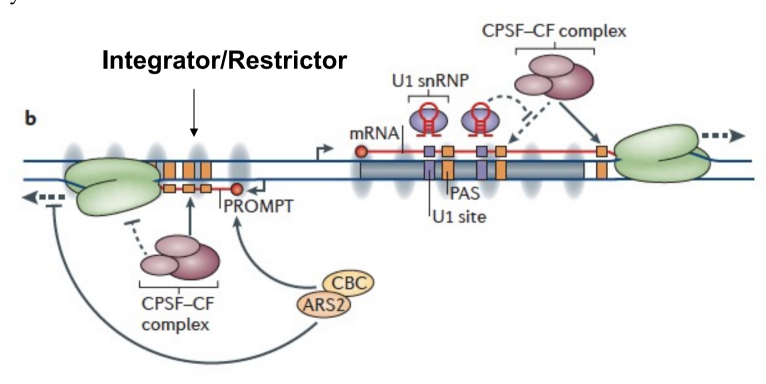


Contrary to the CPF pathway, termination by the Nrd1 pathway is coupled to degradation of the transcript produced or trimming of the precursor in case of snRNAs and snoRNAs

Directing transcription to the right way

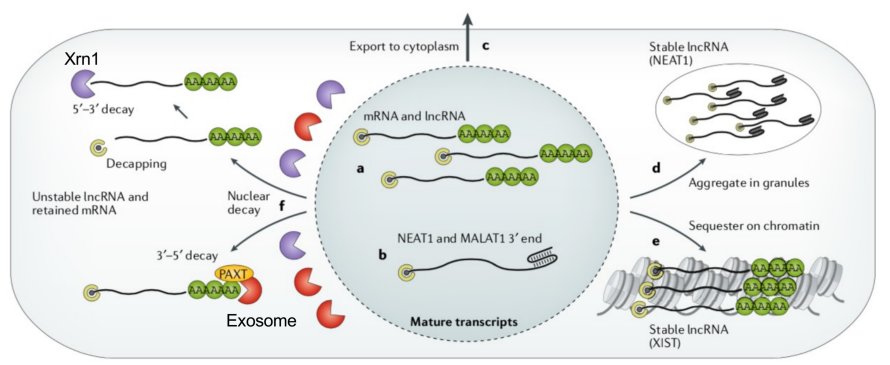
Mammals

RNAPIIs are initiated in two opposite orientations from an active divergent promoter, and paused at the -1 and +1 nucleosomes, respectively. The paused RNAPIIs reinforce the boundaries of Nucleosome Free Regions and maintain the accessibility of promoter to transcription factors. **The** CPSF–CF pathway recognizes the PASs that are present more frequently in the non-functional transcript and induces promoter proximal termination. Interestingly, when present in the mRNA-coding direction, these termination signals are suppressed by the presence of antagonistic U1 snRNP-binding sites that have been shown to inhibit polyadenylation and termination.



Escaping nuclear decay

RNA survival in the degradative environment of the cell nucleus requires end-protective features.. To achieve this, more specialized RNA structures, such as the **triple-helical 3'-ends** of the lncRNAs NEAT1 and MALAT1 (part **b**) can be established. Long-term RNA survival is further provided by its **export to the cytoplasm** (part **c**) or its **sequestration in ribonucleoprotein (RNP) granules** such as paraspeckles (part **d**), as exemplified by the NEAT1 lncRNA, or **on chromatin** (part **e**), as exemplified by the accumulation of XIST on the inactive X chromosome. RNAs that lack such features get degraded by XRN2 or the nuclear RNA exosome, assisted by their respective decapping and poly(A) RNA exosome targeting (PAXT) cofactors (part **f**), composed by Trf4 and Zfc3h1.



Pervasive transcription

- Only 10% of the elongating Pol II molecules in the yeast *Saccharomyces cerevisiae* are engaged in transcription that initiates from conventional promoters. The remaining 90% of the elongating Pol II molecules represent transcriptional noise. Of these, 60% are hyperphosphorylated on the C-terminal domain and associated with chromatin in a salt-stable manner, indicating that they are in the act of transcriptional elongation.
- Most RNAs initiated at inappropriate positions are <u>unstable and rapidly</u> <u>degraded</u> (exosome and nonsense-mediated decay) but a high proportions of eukaryotic genomes produce numerous stable noncoding and antisense RNAs.
- The relative proportions of biologically significant noncoding RNAs and transcriptional noise are unknown.
- Pervasive transcription provides the opportunity for the evolution of new genes.