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The Yeast Cell Cycle Gene *CDC34* Encodes a Ubiquitin-Conjugating Enzyme

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Mutants in the gene *CDC34* of the yeast *Saccharomyces cerevisiae* are defective in the transition from G₁ to the S phase of the cell cycle. This gene was cloned and shown to encode a 295-residue protein that has substantial sequence similarity to the product of the yeast *RAD6* gene. The *RAD6* gene is required for a variety of cellular functions including DNA repair and was recently shown to encode a ubiquitin-conjugating enzyme. When produced in *Escherichia coli*, the *CDC34* gene product catalyzed the covalent attachment of ubiquitin to histones H2A and H2B *in vitro*, demonstrating that the *CDC34* protein is another distinct member of the family of ubiquitin-conjugating enzymes. The cell cycle function of *CDC34* is thus likely to be mediated by the ubiquitin-conjugating activity of its product.

THE CRUCIAL TRANSITION FROM G₁ to the S phase of the cell cycle has been the subject of detailed physiological and genetic analysis in the yeast *Saccharomyces cerevisiae*. When yeast cells reach a critical size during the G₁ phase of the cell cycle, they perform a function known as "start" and become committed to undergoing a cell division (1, 2). The subsequent transition to S phase entails the coordinate initiation of several events, which include bud emergence, spindle morphogenesis, and, finally, DNA replication (1, 3). Many of the genetically controlled functions required during these early phases of the cell

cycle have been identified by characterization of temperature-sensitive (ts) mutations in the cell division cycle (CDC) genes (1). Several of the *cdc* mutants fail to execute the start function under nonpermissive conditions (1, 2). Molecular analysis of the start CDC genes has shown that some of their products regulate pathways of protein phosphorylation (4). The start function is also regulated by yeast analogs of the mammalian guanosine 5'-triphosphate (GTP)-binding proteins G_s-α (5) and RAS (6). After start, functions mediated by the *CDC4* and the *CDC34* gene products are also required before the initiation of chromosomal DNA replication can occur (3, 7, 8). Under nonpermissive conditions, ts mutants in *CDC4* and *CDC34* develop numerous abnormal (elongated) buds, and the spindle pole body duplicates but fails to undergo the separation required for spindle formation (7).

Our molecular analysis of the functions following start was initiated by characterization of the cloned *CDC4* gene. The deduced amino acid sequence of the 779-residue *CDC4* product contains a repeated motif

similar to that found within the β subunits of mammalian GTP-binding proteins, such as G_s and transducin (9). Another segment of the *CDC4* product is similar to the products of the *CDC36* gene and the mammalian *ets* oncogene (10). Here we describe the molecular and functional analysis of another post-start gene, *CDC34*.

Yeast DNA complementing the ts mutation *cdc34-1* in strain G101 was isolated from a library of *S. cerevisiae* genomic DNA (11) constructed in the vector YRp7 (12). The putative *CDC34* gene was mapped within the insert [Fig. 1, (12)], and found by R-loop analysis to encode a polyadenylated RNA ~1.2 kb in length (8). To verify that the cloned DNA sequence contained *CDC34*, we integrated the complementing plasmid into the yeast genome via homologous recombination and mapped the site of integration. Strain G102 :: *CDC34*, which contains a chromosomally integrated copy of plasmid pCDC34-79, was mated to strain G101 (13). Tetrads resulting from the sporulation of this diploid demonstrated tight linkage between *cdc34-1* and the *TRP1* gene present on the integrated plasmid (19/19 parental ditype tetrads). Integration of plasmid pCDC34-79 at the *cdc34* locus indicated that the plasmid contains the *CDC34* gene.

By conventional genetic mapping (Table 1), we located *CDC34* on the right arm of chromosome IV. The map order is *CEN4-trp1-cdc34-mak21-rad55*. Since the restriction map of *CDC34* differs from that of *RRP1*, which also maps in this region, *cdc34* is not allelic to any previously mapped yeast gene.

To determine whether the distinctive multibudded morphology of arrested *cdc34* cells results from the loss of *CDC34* function under nonpermissive conditions, we inserted the yeast *HIS3* gene within the open reading frame of one copy of *CDC34* in two diploid strains (14). When these diploids were sporulated and dissected, 60

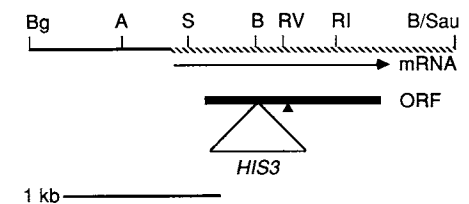


Fig. 1. Physical map of the *CDC34* locus. Indicated is part of plasmid pCDC34-79 showing positions of the sequenced yeast genomic DNA insert (striped segment), the *CDC34* mRNA (arrow), the *CDC34* open reading frame (ORF), the insertion used to generate a truncated *cdc34-3* ORF (19) (small triangle), and the site of a *HIS3* insertion used to disrupt *CDC34* (14). Designations: A, Apa I; B, Bam HI; Bg, Bgl II; RI, Eco RI; RV, Eco RV; S, Sca I; and B/Sau, Bam HI/Sau 3 AI restriction site junction.

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Table 1. Chromosomal mapping of *cdc34* [see (27) for details].

| Gene pair | Segregation (no. of tetrads)* | | | Map distance† (cM) |
|--------------------|----------------------------------|-----|-----|-----------------------|
| | PD | NDP | TT | |
| <i>aro1-cdc34</i> | 12 | 1 | 23 | 42.5 |
| <i>trp1-rad55</i> | 31 | 7 | 108 | 56.9 |
| <i>trp1-cdc34</i> | 43 | 0 | 100 | 44.6 |
| <i>cdc34-rad55</i> | 206 | 0 | 47 | 9.2 |
| <i>cdc34-mak21</i> | 104 | 0 | 3 | 1.4 |
| <i>mak21-rad55</i> | 91 | 0 | 16 | 7.5 |

*PD, parental ditype; NDP, nonparental ditype; TT, tetraptype. †Value (x') in centimorgans (cM) as described in (28).

tetrads each contained two viable spores (which were His⁻) and two inviable spores. The inviable spores germinated without dividing, forming multiple, elongated buds (8). The striking similarity of this probable null *cdc34* phenotype to that caused by the *cdc34-1* ts mutation indicates that the defect in *cdc34-1* mutants results from loss of *CDC34* function under nonpermissive conditions.

When a 2.2-kb Apa I–Sau 3AI DNA fragment containing *CDC34* (Fig. 1) was hybridized to electrophoretically fractionated *S. cerevisiae* DNA that had been digested with various restriction endonucleases, several cross-hybridizing bands were detected (Fig. 2A). A computer-assisted search of the sequenced *CDC34* DNA (see below, Fig. 3) revealed no sequence similarities between the probe and known yeast repetitive elements. However, the DNA encoding a highly acidic region of the *CDC34* product consists largely of aspartic acid-coding triplets GAC and GAT in tandem array, and could possibly cause cross-hybridization to other genes encoding tracts of aspartic acid residues. To explore this possibility, the Apa I–Sau 3AI fragment (Fig. 1) was digested with Eco RV to yield two fragments, one of which contained all of the repeated (GACT)_n sequences (Fig. 3). Both of these fragments hybridized to *CDC34* sequences (Fig. 2B), whereas only the fragment containing the (GACT)_n repeats also hybridized to multiple DNA sequences (8). We conclude that although a number of yeast genes may encode tracts rich in aspartic acid, there are no close homologs of *CDC34* in the *S. cerevisiae* genome.

The nucleotide sequence of a 1820-bp genomic DNA segment from pCDC34-79 that includes *CDC34* is shown in Fig. 3. The only large open reading frame encodes a protein of 295 residues if the first methionine codon of the frame serves as the initiation codon. We conclude that this open reading frame encodes the *CDC34* gene product, which would have a molecular

mass of 34.1 kD. A computer-assisted search (15) for similarities between the deduced amino acid sequence of the *CDC34* product and known proteins revealed substantial similarity only to the yeast *RAD6* protein (Fig. 3). The *RAD6* gene is required for a variety of functions including DNA repair, induced mutagenesis, and sporulation (16). Amino acid sequence similarities between the *RAD6* and *CDC34* products are distributed throughout their lengths, and are especially pronounced in the central region of the *CDC34* product as well as in the COOH-terminal regions where aspartic acid-rich tracts occur in both proteins. Overall, 38% of the amino acids in the *RAD6* product are identical to those in the *CDC34* product. With the inclusion of conservative replacements (17), 55% of the *RAD6* product is conserved within the *CDC34* product. Recently, Jentsch *et al.* (18) found that *RAD6* is a member of a family of genes encoding related but distinct ubiquitin-conjugating enzymes (E2 enzymes) that catalyze the covalent attachment of ubiquitin, a highly conserved 76-residue protein, to specific protein substrates (18). The striking amino acid sequence similarity between the *RAD6* and *CDC34* gene products (Fig. 3) suggested that the latter protein may also be a ubiquitin-conjugating enzyme.

To test this possibility, we carried out assays for ubiquitin-conjugating activity in extracts of *E. coli* expressing *CDC34*, as well as in control extracts of *E. coli* harboring the same vector but with *CDC34* inserted in the opposite (presumably non-expressing) orientation (19, 20). The conjugation of ubiquitin to acceptor proteins begins with an

adenosine 5'-triphosphate (ATP)-requiring step in which the COOH-terminal glycine residue of ubiquitin is joined, through a high-energy thiolester bond, to a cysteine residue of the ubiquitin-activating enzyme, E1 (21–23). The activated ubiquitin is then transferred to a specific cysteine residue within a ubiquitin-conjugating (E2) enzyme, which in turn catalyzes the formation of a branched isopeptide bond-mediated conjugate between the COOH terminus of ubiquitin and the ε-amino group of a lysine residue in the acceptor protein (21, 23). When supplemented with ATP and purified yeast ubiquitin-activating enzyme, E1, only extracts of *CDC34*-expressing *E. coli* cells mediated the covalent conjugation of ¹²⁵I-labeled ubiquitin to histones H2B (Fig. 4A) and H2A (24). The formation of a metastable intermediate, in which an E2 enzyme is joined to ubiquitin by a thiolester bond in a reaction dependent on both ATP and ubiquitin-activating (E1) enzyme, could also be demonstrated with the extract of *E. coli* expressing the *CDC34* gene but not with an otherwise identical extract lacking the *CDC34* protein (Fig. 4, B and C). The remarkable degree of substrate specificity of the *CDC34* ubiquitin-conjugating enzyme is indicated by its apparent failure to ubiquitinate endogenous proteins in the *E. coli* extract (Fig. 4A).

The demonstration of ubiquitin-histone conjugating activity in bacterial cells expressing *CDC34* and the ability of the *CDC34* protein to form a thiolester bond-mediated covalent complex with ubiquitin directly identifies the *CDC34* gene as yet another member of the gene family encoding related but distinct E2 enzymes. In

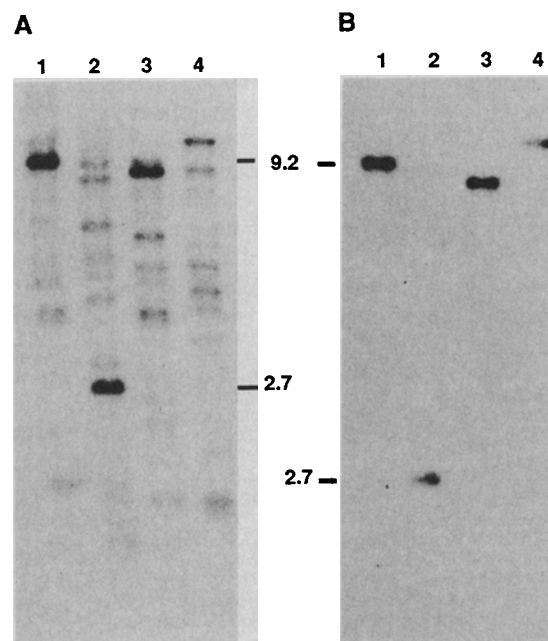


Fig. 2. DNA hybridization analysis of yeast genomic DNA with *CDC34* probes. *Saccharomyces cerevisiae* DNA was digested with the following restriction endonucleases before electrophoresis in a 1.0% agarose gel and DNA hybridization with ³²P-labeled probes (29, 30): lanes 1, Bam HI; lanes 2, Bgl II; lanes 3, Cla I; and lanes 4, Eco RI. (A) The hybridization probe was a 2.2-kb Apa I–Sau 3AI fragment (see Fig. 1) containing the entire *CDC34* gene. (B) The hybridization probe was a 1.0-kb Apa I–Eco RV fragment containing the portion of *CDC34* that encodes an NH₂-terminal part of the *CDC34* protein (see Figs. 1 and 3).

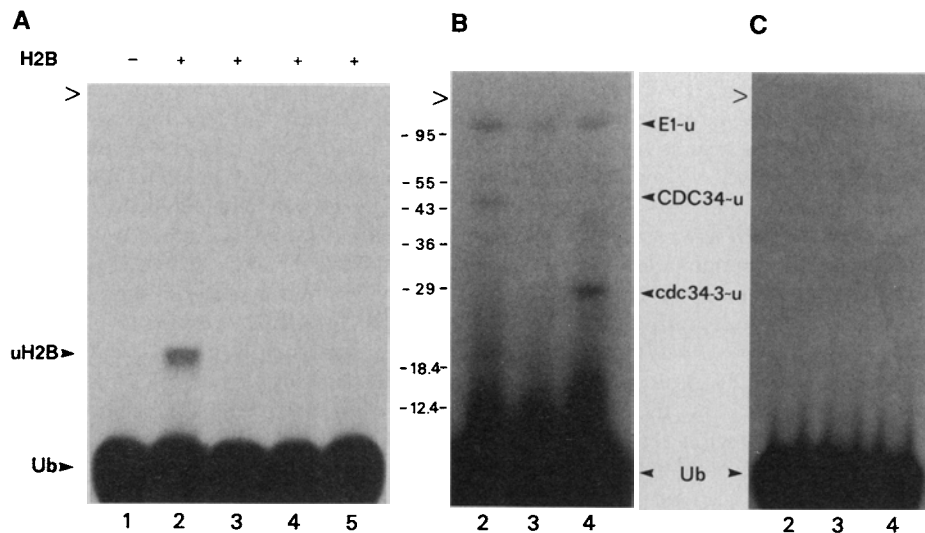


Fig. 4. The *CDC34* gene product is a ubiquitin-conjugating enzyme. Extracts of induced *E. coli* cells expressing the *CDC34* gene (lanes 1, 2, and 5), extracts of cells containing the *CDC34* gene in an opposite (non-expressing) orientation within the vector (lanes 3), and extracts of cells expressing the mutant (truncated) *cdc34-3* gene (lanes 4) were assayed for ubiquitin conjugation activity (20). Products of the assay were separated by electrophoresis in an 18% polyacrylamide-SDS gel and detected by autoradiography (20). (A) Extracts were assayed for ubiquitin-histone H2B conjugation in either the absence (lane 5) or presence (lanes 1 to 4) of the purified ubiquitin-activating enzyme, E1, and in the presence of ATP, ¹²⁵I-labeled ubiquitin, and histone H2B (20). The extract of *CDC34*-expressing cells could form a monoubiquitinated species of histone H2B in an E1-dependent reaction (lanes 2 and 5), whereas neither the extract of non-expressing cells (lane 3) nor the extract of cells expressing the mutant (truncated) *cdc34-3* gene (lane 4) possessed this activity. (B) When assayed for ubiquitin-CDC34 protein thiolester formation (21–23), complexes with the properties of thioesters were detected between ubiquitin and added E1 (E1~u; lanes 2 to 4), *CDC34* protein (*CDC34*~u, ~45 kD; lane 2), and *cdc34-3* protein (*cdc34-3*~u, ~28 kD; lane 4). (C) Same as in (B) but the samples were heated under reducing conditions (20) before electrophoresis to selectively cleave thiolester-mediated complexes. Designations: Ub, ubiquitin; uH2B, monoubiquitinated histone H2B. Molecular masses of size markers are in kilodaltons. Arrowheads indicate gel origins.

CDC34 (UBC3) ubiquitin-conjugating enzyme in mediating the G₁ to S transition.

Although the *RAD6* (*UBC2*) and *CDC34* (*UBC3*) products are clearly distinct both structurally and functionally, their enzymatic specificities appear to be quite similar in vitro: both enzymes apparently fail to ubiquitinate endogenous proteins in an *E. coli* extract but efficiently ubiquitinate histones H2A and H2B in E3-independent reactions (Fig. 4) (18). If the *CDC34* (*UBC3*) and *RAD6* (*UBC2*) enzymes share functionally relevant in vivo substrates, the distinct functions of these enzymes must be derived from differences in their specific modes of action. One possibility is that ubiquitination of histones or other substrates by the *CDC34* (*UBC3*) and *RAD6* (*UBC2*) enzymes might be confined to different specific regions of chromatin, such as sites involved in the initiation of DNA replication in the case of the *CDC34* (*UBC3*) enzyme or sites of DNA damage in the case of the *RAD6* (*UBC2*) enzyme. Alternatively, the functionally relevant in vivo protein substrates of the *CDC34* (*UBC3*) and *RAD6* (*UBC2*) enzymes may include factors involved in the regulation of genes that control, respectively, post-start events required

for the initiation of DNA replication and events necessary for DNA repair or sporulation. All of the above possibilities are consistent with immunofluorescence evidence (8) that the *CDC34* (*UBC3*) enzyme is a nuclear protein.

That progress through the cell cycle may involve specific ubiquitination events has been suggested by earlier work on the mutant mammalian cell line, ts85, whose preferential G₂ arrest under nonpermissive conditions results from thermolability of the ubiquitin-activating enzyme in these cells (22). The discovery that a regulatory step in the G₁ to S transition is controlled in yeast by a specific ubiquitin-conjugating enzyme underscores the remarkable functional diversity of the ubiquitin system and opens up a new direction for studies of the cell cycle.

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12. The vector YRp7, which consists of the *S. cerevisiae* *TRP1-ARS1* sequence inserted into the Eco RI site of pBR322, is able to replicate in both *S. cerevisiae* and *E. coli* [K. Struhl *et al.*, *Proc. Natl. Acad. Sci. U.S.A.* **76**, 1035 (1979)]. All plasmids were constructed by standard techniques (31). The plasmids were maintained in the *E. coli* strains DH-1 or RRI. The yeast strains used were G101 (*MATα cdc34-1 trp1 ura1 ade2 his*) and G102 (*MATα cdc34-1 trp1 ura1 leu2 lys2*). Plasmid DNA was isolated from *E. coli* by alkaline lysis [H. C. Birnboim and J. Doly, *Nucleic Acids Res.* **7**, 1513 (1979)] and from *S. cerevisiae* as described (11). Transformations of *E. coli* and *S. cerevisiae* were performed by standard techniques [M. Mandel and A. Higa, *J. Mol. Biol.* **53**, 159 (1970); and (32)]. Although the *cdc34-1* mutant is viable at 37°C on medium containing 1M sorbitol, it grows more slowly than *CDC34* strains. Thus, *cdc34-1* cells transformed with *CDC34* could be initially identified by their ability to grow more rapidly under the above conditions (8).
13. The plasmid pCDC34-79 was transformed into the strain G102 (32), and a stable ts⁺ (non-ts) transformant was isolated and designated G102::*CDC34* [after confirmation by DNA hybridization analysis (29, 30) that the plasmid integration site was within the homologous genomic sequences]. Standard genetic techniques were used to construct and sporulate diploids for tetrad analysis (32).
14. To construct a disruption of *CDC34* [R. Rothstein, *Methods in Enzymol.* **101**, 202 (1983)], the plasmid Cmp170 was digested with Bam HI, liberating a 1.7-kb fragment containing the yeast *HIS3* gene [K. Struhl, *Nucleic Acids Res.* **13**, 8587 (1985)]. This fragment was inserted, by standard techniques (31), into the Bam HI site of pGEM34 *H/S* (33). The resulting plasmid was digested with Apa I-Eco RI (Fig. 1) and transformed (32) into two diploid strains. DNA hybridization analysis of DNA from His⁺ transformants and from the spore-derived colonies of one tetrad from each transformant confirmed that His⁺ diploids contained both the wild-type and insertional alleles of *CDC34*, whereas the viable spore colonies contained only the wild-type allele of *CDC34*.
15. The nucleotide sequence of *CDC34* was determined by the chain termination method [F. Sanger *et al.*, *Proc. Natl. Acad. Sci. U.S.A.* **74**, 5463 (1977)]. The plasmids pGEM34 B/S and pGEM34 H/S (33) were used to generate a series of overlapping deletions extending into either side of *CDC34* by recloning fragments produced by controlled exonuclease III digestion [S. Henikoff, *Gene* **28**, 351 (1984)]. This approach allowed both strands to be sequenced completely. The similarity between the deduced amino acid sequences of *CDC34* and *RAD6* proteins was found by searching a database of amino acid sequences of yeast proteins (collected by M. Goebel) by means of the GENEPRO program (Riverside Scientific Enterprises, Seattle). Alignment of the two gene products utilized the algorithm of W. J. Wilbur and D. Lipman [*Proc. Natl. Acad. Sci. U.S.A.* **80**, 726 (1983)].
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20. *Escherichia coli* cells harboring the pK34-1, pK34-2, or pK34-3 plasmid (19) were induced (19) and lysed by freeze-thawing in 5 volumes of 50 mM tris-HCl (pH 7.5), 1 mM EDTA, and 1 mM dithiothreitol (DTT). The assay for ubiquitin-histone conjugation was carried out in 50 μ l of 50 mM tris-HCl (pH 7.5), 2 mM ATP, 5 mM MgCl₂, and 0.2 mM DTT, in the presence of 50 pmol of ¹²⁵I-labeled ubiquitin (~1 \times 10⁵ cpm) (21, 22), 0.2 μ g of bovine histone H2B (Boehringer Mannheim), and 4 pmol of the purified yeast E1 enzyme (18). The assay for ubiquitin thioesters was carried out as above except that histone H2B was omitted, and the reaction products were incubated for 5 min at room temperature in an SDS-containing electrophoretic sample buffer lacking reducing agents before a polyacrylamide-SDS gel electrophoresis at 4°C (22). Reduction of thioesters was performed by heating the samples prior to electrophoresis at 100°C for 10 min in an SDS sample buffer containing 4M 2-mercaptoethanol (18, 21).
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Physical Analysis of Transcription Preinitiation Complex Assembly on a Class II Gene Promoter

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Transcription of protein-encoding genes by human RNA polymerase II requires multiple ancillary proteins (transcription factors). Interactions between these proteins and the promoter DNA of a viral class II gene (the major late transcription unit of adenovirus) were investigated by enzymatic and chemical footprinting. The experiments indicated that the assembly of functionally active RNA polymerase II-containing transcription preinitiation complexes requires a complete set of transcription factors, and that both specific protein-DNA and protein-protein interactions are involved. This allows individual steps along the transcription reaction pathway to be tested directly, thus providing a basis for understanding basic transcription initiation mechanisms as well as the regulatory processes that act on them.

GENE ACTIVITY IN EUKARYOTES IS frequently regulated at the level of DNA transcription. Understanding the molecular mechanism behind this transcriptional control of gene expression has become one of the major goals of modern biology. Mutational analysis has revealed that the activity of protein-encoding genes is governed by several kinds of promoter elements. A basal level of transcription is usually observed when all but a small region of the promoter DNA has been deleted. This minimum or core promoter is often centered on a TATA box sequence (1). Two classes of cis-acting DNA elements are involved in modulating the TATA box-driven transcription. Upstream promoter elements act at short distance, while enhancer elements can be located up to several thousand base pairs from the transcription initiation site (2). The various components of the human class II transcription machinery are being elucidated by the biochemical dissection of crude cell-free systems (3, 4). This approach has revealed that specific transcription initiation by RNA polymerase II requires the coordinated action of multiple protein factors. These can be classified into two categories: general transcription factors and upstream element-binding proteins (2). Through both kinetic analysis and the use of

various inhibitors, several steps have been defined within the transcription mechanism (5-7). These include: (i) a commitment of the template after binding of a subset of the transcription factors, (ii) formation of an activated state through the action of other transcription factors and RNA polymerase II, (iii) fulfillment of an energy requirement (hydrolysis of the β - γ phosphate bond of either ATP or dATP), (iv) initiation of transcription (formation of the first phosphodiester bond), and (v) transcription elongation, with some transcription components remaining committed to the template. Within these various steps, the exact roles of the different protein factors, their assembly into active transcription complexes, and the various transitions that lead to the initiation event, remain for the most part unknown.

The major late (ML) promoter of adenovirus provides a useful model system for in vitro analysis of specific transcription by human RNA polymerase II. This promoter is quite simple structurally, with only two essential DNA elements: a TATA box at position -28 and a single upstream element at position -58 (8-11). At least three transcription factors (designated TFIIB, TFIID,

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