Cell cycle–regulated phosphorylation of \( p220^{\text{NPAT}} \) by cyclin E/Cdk2 in Cajal bodies promotes histone gene transcription

Tianlin Ma,1,8 Brian A. Van Tine,4,5,8 Yue Wei,2 Michelle D. Garrett,6 David Nelson,7 Peter D. Adams,7 Jin Wang,1,3 Jun Qin,1,3 Louise T. Chow,4 and J. Wade Harper1,2,9

1Department of Biochemistry and Molecular Biology, 2Department of Molecular Physiology and Biophysics, 3Department of Molecular and Cellular Biology, Baylor College of Medicine, Houston, Texas 77030, USA; 4Department of Biochemistry and Molecular Genetics, 5Department of Pathology, University of Alabama, Birmingham, Alabama 35294, USA; 6CRC Centre for Cancer Therapeutics, Institute of Cancer Research, Sutton, SM2 5NG, United Kingdom; 7Fox Chase Cancer Center, Philadelphia, Pennsylvania 19111, USA

Cyclin E/Cdk2 acts at the G1/S-phase transition to promote the E2F transcriptional program and the initiation of DNA synthesis. To explore further how cyclin E/Cdk2 controls S-phase events, we examined the subcellular localization of the cyclin E/Cdk2 interacting protein p220^{NPAT} and its regulation by phosphorylation. p220 is localized to discrete nuclear foci. Diploid fibroblasts in Go and G1 contain two p220 foci, whereas S- and G2-phase cells contain primarily four p220 foci. Cells in metaphase and telophase have no detectable focus. p220 foci contain cyclin E and are coincident with Cajal bodies (CBs), subnuclear organelles that associate with histone gene clusters on chromosomes 1 and 6. Interestingly, p220 foci associate with chromosome 6 throughout the cell cycle and with chromosome 1 during S phase. Five cyclin E/Cdk2 phosphorylation sites in p220 were identified. Phospho-specific antibodies against two of these sites react with p220 within CBs in a cell cycle–specific manner. The timing of p220 phosphorylation correlates with the appearance of cyclin E in CBs at the G1/S boundary, and this phosphorylation is maintained until prophase. Expression of p220 activates transcription of the histone H2B promoter. Importantly, mutation of Cdk2 phosphorylation sites to alanine abrogates the ability of p220 to activate the histone H2B promoter. Collectively, these results strongly suggest that p220^{NPAT} links cyclical cyclin E/Cdk2 kinase activity to replication-dependent histone gene transcription.

[Key Words: Cyclin-dependent kinases; phosphorylation; Cajal [coiled] bodies; histone transcription]

Received June 23, 2000; revised version accepted August 1, 2000.

Cyclin E, an essential regulatory subunit of Cdk2 [Dulic et al. 1992; Koff et al. 1992], plays a central role in coordinating both the onset of S phase and centrosome duplication in multicellular eukaryotes [Sherr 1996; Reed 1997]. Cyclin E/Cdk2 complexes have two major roles in promoting S phase. First, cyclin E/Cdk2 participates, together with cyclin D/Cdk4, in the control of transcriptional processes that are critical to cell cycle progression. The best understood example is control of the E2F/DP transcription factor via phosphorylation of a family of transcriptional repressors (Rb, p130, and p107) [for review, see Reed 1997; Dyson 1998; Nevins 1998]. E2F complexes regulate the S-phase-dependent expression of a number of proteins required for the synthesis of nucleic acids as well as proteins such as Cdc2 and cyclin A that promote subsequent cell cycle transitions. Second, cyclin E/Cdk2 can function in an E2F-independent manner to activate DNA replication. Accumulation of cyclin E is required for S-phase entry, and ectopic cyclin E expression can bypass the requirement for Rb inactivation and E2F activation for S-phase entry [Ohtsubo et al. 1995; Leng et al. 1997; Lukas et al. 1997].

An understanding of the role of cyclin E/Cdk2 in promoting S phase requires knowledge of its essential substrates. Insight into Cdk targets has been advanced by the finding that several Cdk substrates bind tightly to the cyclin subunit. In some cases, this interaction involves a motif in the substrate, the RXL motif, and a conserved pocket in the cyclin box [Zhu et al. 1995; Adams et al. 1996; Russo et al. 1996; Schulman et al. 1998; Brown et al. 1999; Ma et al. 1999]. We and others have exploited this property of cyclins to identify relevant cyclin E/Cdk2 substrates with the use of expression cloning [Zhao et al. 1998; Ma et al. 1999]. One of these, p220^{NPAT}, interacts with cyclin E/Cdk2 in extracts from tissue culture cells and accelerates S-phase entry when overexpressed [Zhao et al. 1998]. Moreover, retroviral insertion into the mouse p220^{NPAT} gene leads to embry-
onic lethality at the eight-cell stage, indicating an essential role for p220 in cell division or development (Di Fruscio et al. 1997). However, the precise function of p220 and the role of cyclin E/Cdk2 in its action remain unknown.

Emerging data [Zhao et al. 2000, this paper] suggest that p220 is involved in S-phase-specific histone gene transcription. Histones, components of nucleosomes, have to be supplied on demand during DNA replication. This regulation is attributed to both transcriptional and posttranscriptional control mechanisms [Harris et al. 1991; Heintz 1991], mediated in part by the activation of histone gene–specific transcription factors [Oct-1 in the case of the H2B promoter and H1TF2 in the case of the H1 promoter] through an unknown mechanism [Fletcher et al. 1987; Segil et al. 1991]. Once generated, histone messages are stabilized and processed preferentially in S phase. Implicated in histone gene transcription are Cajal bodies (CBs; sometimes referred to as coiled bodies). CBs were initially described as small nuclear organelles (Cajal 1903), but their function has remained obscure for the better part of the twentieth century. Recent work has led to the hypothesis that CBs are sites of assembly of transcription and splicing complexes [Gall et al. 1999]. The link to histone transcription comes from the finding that a subset of CBs is physically associated with histone gene clusters on chromosomes 1 (1q21) and 6 (6p21) [Frey and Matera 1995] and with histone gene loci in Xenopus lampbrush chromosomes [Abbott et al. 1999]. Moreover, CBs also contain a component of the histone mRNA 3′-end processing machinery SLBP1 [Abbott et al. 1999].

Here, we report that p220NPAT is localized to discrete foci that are coincident with a subset of CBs in normal diploid fibroblasts. The number of p220 foci increased from two in Go and G1 cells in association with chromosome 6 to four in S and G2 phases in association with both chromosomes 6 and 1. Foci are lost during mitosis. Consistent with these observations, Zhao et al. [2000] have found that p220 is associated directly with histone gene clusters and that overexpression of p220 can activate histone 2B and histone 4 transcription. We also demonstrate that cyclin E is contained in p220 foci and that p220 within CBs is phosphorylated on Cdk sites in a cell cycle–dependent manner. Moreover, mutation of cyclin E/Cdk2 phosphorylation sites in p220 reduces its ability to activate expression from histone H2B reporter constructs in transiently transfected cells. These data, together with those of Zhao et al. [2000], suggest that cyclin E/Cdk2 functions in conjunction with p220 to coordinate S-phase-dependent histone gene transcription; they also demonstrate a role for CBs in cell cycle–regulated transcriptional control.

Results

p220 is localized in cell cycle–regulated nuclear foci

We previously identified a C-terminal fragment of NPAT (residues 1054–1397) in a cyclin E/Cdk3 interaction screen [Ma et al. 1999]. Affinity-purified anti-NPAT an-tibodies generated against this C-terminal fragment recognize a closely spaced protein doublet of 220 kD in molecular mass in nuclear extracts from HeLa and 293 cells, as determined by either immunoblotting or immunoprecipitation (Fig. 1a, lane 2; data not shown). The identity of the p220 protein obtained by immunoprecipitation was confirmed by mass spectral analysis of tryptic peptides [see below].

To examine the subcellular localization of p220, we performed immunofluorescence by using normal diploid fibroblasts [Fig. 1b]. The majority of cells (>80%) in an asynchronous culture contained either two or four nuclear foci staining for p220, whereas the remaining cells contained one or three obvious p220 foci. This immunoreactivity was blocked by competition with antigen [Fig. 1b]. The variation of p220 staining patterns suggested that p220 localization might be cell cycle regulated. To test this possibility, we examined p220 localization in several asynchronous growing fibroblast lines [normal dermal fibroblasts, WI38 fibroblasts, and bjtERT fibroblasts] labeled with BrdU to mark S-phase cells. Similar results were observed, and the data for bjtERT cells are shown in Figure 1c,d. The vast majority (70%) of cells lacking BrdU staining contained two p220 foci, whereas 87% of BrdU-positive cells contained four p220 foci [in green] [Fig. 1d].

Consistent with the hypothesis of cell cycle–regulated foci formation, quantitative analysis of relative 4′,6-di-
loss of p220 foci during this short period would explain
the small number of cells with one or no p220 focus (Fig.
2a). Taken together, these data demonstrate that the
number of p220 foci is cell cycle regulated and that S
phase is accompanied by the generation of two addi-
tional p220 foci not seen in G1 or Go cells.

p220 foci are associated with CBs and with chromosomes 1 and 6

The size and number of p220 foci observed in S-phase
cells are reminiscent of those displayed by CBs, as de-
ected by antibodies against a component p80coilin. CBs are present in variable numbers in tissue culture cell
lines [three to eight CBs/cell] [Frey and Matera 1995;
Almeida et al. 1998]. Because CBs typically are difficult
to detect in nontransformed cells, we used antigen re-
tieval to examine whether p220 might be associated
with CBs in normal human dermal fibroblasts. As shown
in Figure 3a, p220 foci coincide with coilin-containing
CBs. In contrast to the colocalization of p220 and coilin
throughout most of the cell cycle observed with three
lines of fibroblasts [diploid dermal fibroblasts, WI38 lung
fibroblasts, and BjTERT fibroblasts], transformed cells,
including HeLa, Caski, SiHa, and MCF7 cells, displayed
a larger and more variable number of p220 foci, ranging
from three to >12 [data not shown]. In HeLa cells, most
if not all of the p220 foci are associated with CBs, but
only a subset of CBs is associated with p220 foci [Fig. 3c].

Because CBs have previously been demonstrated to as-
sociate with histone gene clusters on chromosomes 1
and 6, it follows that one or more p220 foci may be ex-
pected to associate with these chromosomal domains. By
using interphase chromosome painting in normal dermal
fibroblasts, we found that chromosome 6 signals were
closely associated with both p220 foci in 100% of cells
containing two p220 foci. In 87% of cells containing four
foci, the signals were associated with two foci, whereas
the remaining 13% had more than two associated foci
[Fig. 3d; Table 1]. In contrast, chromosome 1 signals typi-
cally were not associated with p220 foci in cells contain-
ing two foci, but 93% of cells with four foci had two
associated foci (Fig. 3e; Table 1). In our experience, false
positive association occurs at a frequency of ∼10% or
lower. Because a subset of CBs is physically associated
with an snRNA U2 gene loci at 17q21 [Frey and Matera
1995], we painted chromosome 17 as well as chromo-
somes 5 and Y as additional controls. These chromosom-
al domains displayed only rare association with p220 foci
[Fig. 3e]. For example, out of 400 cells, two were found to
have one p220 focus associated with chromosome Y.
Taken together, these data indicate that p220 foci are
intimately linked with chromosome 1- and chromo-
some 6-associated CBs and that the chromosome 6 do-
main is associated with p220 foci throughout the cell
cycle, whereas association with chromosome 1 occurs
during S phase and coincides with the increase in p220
foci from two to four. These data also imply the exis-

Figure 1. p220 is located in cell cycle-regu-
lated nuclear foci. (a) Affinity-purified poly-
clonal antibodies against p220 immunopre-
cipitate a closely spaced doublet of proteins
220 kD in molecular mass from tissue cul-
ture cells. For a large-scale immunoprecipi-
tation, nuclear extracts from 293T cells [44
mg in 9 mL] were immunoprecipitated with
20 µg of anti-p220 antibodies or pre-im-
immune IgG bound to 80 µL of protein
A–Sepharose. Washed immunoprecipitates
were separated using SDS–PAGE, and the
gel was stained with Coomassie blue (top).
A small fraction of this immune complex
was immunoblotted with anti-p220 anti-
obodies [bottom]. [M] Molecular mass mark-
ers with masses indicated at left, [NRS] nor-
mal rabbit sera, [IP] immunoprecipitate. (b)
p220 is localized in discrete nuclear foci.
WI38 fibroblasts were subjected to indirect
immunofluorescence using anti-p220 anti-
obodies in the presence (right) or absence
(left) of 0.5 µg of antigen. (red) p220, (blue)
nuclei stained with 4′,6-diamidino-2-phe-
nylindole (DAPI). (c) Cells with four p220
foci accumulate during S phase. Asynchro-
nous bjTERT fibroblasts were pulse-labeled
with BrdU for 60 min and then stained for
p220 and BrdU. The number of p220 foci in
BrdU-positive and BrdU-negative cells was determined from a minimum of 100 cells. (d) An example of BrdU-positive [green] cells
displaying three or four p220 foci [red], whereas a BrdU-negative cell had two p220 foci. DAPI staining of nuclei is in blue.
tence of mechanisms that restrict association of p220 with particular chromosomes to particular points in the cycle. The increased number of p220 foci observed in some transformed cells (data not shown) likely reflects at least in part an increased ploidy in chromosomes 1 and 6.

p220 is specifically phosphorylated by cyclin E/Cdk2 on sites near the cyclin E interaction domain

Cyclin E and p220 co-immunoprecipitate from cell extracts, and cyclin E/Cdk2 can phosphorylate associated p220 (Zhao et al. 1998). To elucidate the role of cyclin E/Cdk2 in p220 regulation, we sought to determine the specificity of phosphorylation. Initially we examined the ability of p220 to bind to various cyclin/Cdk complexes. Flag-tagged p220 was expressed in insect cells and cell lysates used in binding assays with immobilized cyclin/Cdk complexes (Fig. 4). Although p220 associated efficiently with the cyclin E/Cdk2 complex (lane 16), it did not associate with the cyclin D1/Cdk4, cyclin A/Cdk1, or cyclin B/Cdk1 complex (lanes 4, 10, and 13, respectively) and bound only weakly with cyclin A/Cdk2 (lane 7). Thus, p220 displays specificity for cyclin E/Cdk2.

As expected, p220 was readily phosphorylated by the associated cyclin E/Cdk2 complex, and this phosphorylation was accompanied by reduced mobility of p220 (Fig. 4, lanes 16 and 17). Although p220 bound weakly to cyclin A/Cdk2, the associated protein underwent a similar mobility shift in the presence of ATP (lanes 7 and 8), suggesting that cyclin A/Cdk2 can also phosphorylate p220 when bound. We note that control reactions employing control insect cell lysates revealed the presence of a cyclin E/Cdk2–associated substrate (indicated by the asterisk) that migrated slightly faster than did p220 (Fig. 4, lanes 9 and 18). It can be distinguished from the human p220, because it was also phosphorylated to similar levels by cyclin A/Cdk2.

We next sought to determine the sites of p220 phosphorylation in vitro by using mass spectrometry (Zhang et al. 1998). p220 contains 18 potential Cdk phosphorylation sites [Thr/Ser followed by Pro]. Four tryptic p220 phosphopeptides containing five phosphorylation sites were identified in recombinant p220 phosphorylated by
Figure 3. Association of p220 foci with Cajal bodies (CBs) and with domains of chromosomes 1 and 6. In all panels, 4′,6-diamidino-2-phenylindole (DAPI) stained nuclear DNA blue. [a] Normal dermal fibroblasts were subjected to immunofluorescence by using anti-p220 (green) and anti-p80 coilin monoclonal antibodies [red] known to stain CBs. Co-localization is demonstrated in the merged image. [b] p220 foci are present in prophase (left, right, cell on top), but are no longer detectable in metaphase (middle) and telophase (right, cell at bottom). Prophase cells with four p220 foci were also observed (not shown). [c] HeLa cells contain CBs devoid of p220 foci. In a–c, p220 is green and DAPI is blue, in a and c, coilin is red. [d] and [e] p220 foci are associated with chromosomes 1 and 6 but not with other chromosomes. Normal dermal fibroblasts were stained for p220 and for the indicated chromosomal domains by using chromosome paints. For chromosomes 6, 17, 5, and Y, p220 is green and chromosome paint is red. For chromosome 1, p220 is red and chromosome paint green. [Chr.] Chromosome; [M] metaphase; [P] prophase; [T] telophase.
associated cyclin E/Cdk2 in vitro (Fig. 5a,b, Table 2). Three singly phosphorylated peptides were sequenced by liquid chromatography/mass spectrometry/mass spectrometry (LC/MS/MS) to identify the phosphorylation sites as S1100 (site 3), T1270 (site 4), and T1350 (site 5), respectively (Table 2). The sequence of a doubly phosphorylated peptide encompassing residues 742–788 could not be determined because of its large size (m/z 5077.5, average mass; see Fig. 5a). However, this peptide contains two consensus Cdk substrates at S775 and S779 (sites 1 and 2) (Table 2), allowing a tentative assignment as sites of modification by cyclin E/Cdk2. Indeed, we found that p220 mutated in both of these serine residues was resistant to a cyclin E/Cdk2–induced shift in mobility (Fig. 5d). In contrast, p220 proteins mutated in one or more of the other identified phosphorylation sites still underwent a mobility shift in response to cyclin E/Cdk2 treatment (Fig. 5d). These data are consistent with the assignment of S775 and S779 as in vitro substrates and indicate that they are primarily responsible for the mobility shift observed upon phosphorylation by cyclin E/Cdk2.

To examine phosphorylation in vivo, we purified p220 from a 293T cell nuclear lysate by using immunoprecipitation [Fig. 1a] and subjected it to mass spectral analysis. p220 is present at low levels in this cell line, and from 44 mg of nuclear extract, we obtained 200 ng of p220. We were able to identify one peptide with the mass expected for a doubly phosphorylated peptide spanning Val742–Lys788 whose quantity was consistent with the level of protein available for analysis (Fig. 5b, Table 2). Importantly, this peptide is absent from spectra after treatment with a phosphatase (Fig. 5b), indicating that S775 and S779 are phosphorylated in vivo. Signals for the three singly phosphorylated peptides observed in vitro were not evident, possibly because of the small amounts of material available for analysis.

Table 1.  Association of p220 foci with chromosomes 6 and 1

| No. of foci associated | Chromosome 6 (%) | Chromosome 1 (%) |
|------------------------|------------------|------------------|
| Cells with 2 foci       |                  |                  |
| 0                      | 0                | 91               |
| 1                      | 0                | 9                |
| 2                      | 100              | 0                |
| Cells with 4 foci       |                  |                  |
| 0                      | 0                | 0                |
| 1                      | 0                | 1                |
| 2                      | 87               | 93               |
| 3                      | 11               | 6                |
| 4                      | 2                | 0                |

One hundred cells with 2 and 4 foci, respectively, were counted for association with the indicated chromosomes. With chromosomes Y, 5, and 17, association was rare; for example, 2 of 400 cells displayed one p220 foci associated with chromosome Y.

Figure 4. p220 preferentially associates with and is phosphorylated by cyclin E/Cdk2 in vitro. Immobilized cyclin/Cdk complexes were incubated with control insect cell lysates or insect cell lysates containing Flag-p220, as described in Materials and Methods. Complexes were washed with lysis buffer followed by 10 mM MgCl2 and 20 mM Tris-HCl. Some samples were supplemented with γ-[32P]ATP for 20 min before SDS–PAGE and visualization of proteins by immunoblotting or autoradiography. Flag-p220 was detected by anti-flag antibodies. The quantities of GST–cyclin/Cdk complexes were similar, as determined by immunoblotting with GST antibodies. Cdk2 complexes associated with an insect cell protein migrating slightly faster than p220 that was also a substrate for the kinase (indicated by an asterisk). An anti-Flag immunoprecipitate of Flag-p220 (lane 19) was included as a control.
immunoprecipitation from ATP-containing reticulocyte lysates, because larger amounts of recombinant p220 purified from insect cells did not react with the phosphopeptide-specific antibodies in the absence of phosphorylation by cyclin E/Cdk2 (lanes 5 and 6). We also note that a general phosphothreonine-proline antibody gave similar results, indicating that other TP sequences in p220/H9004 Cdk are not phosphorylated by bound cyclin E/Cdk2 in vitro. We found that both phosphopeptide antibodies reacted with p220 immunoprecipitated from cyclin 293T cells (Fig. 5f), indicating that these sites are indeed phosphorylated in vivo. On the basis of the comparison with p220 in 293T cells examined in parallel [Fig. 5f], the more slowly migrating p220 protein

Table 2. Mass spectral identification of p220 phosphorylation sites

| Sites          | Peptides                                      | Molecular mass [measured/calculated] | No. of PO3 group |
|---------------|-----------------------------------------------|--------------------------------------|------------------|
| In vitro      |                                               |                                      |                  |
| S775, S779    | V I I S D D P F V S S D T E L T S A V S S I N G E N L P T I I L S S T K S P T K N A E L V K<sup>1</sup> | 4918/4916.5 2                      |                  |
| S1106         | N A V S F P N L D S P N V S S T L K P S N A I K<sup>2</sup> | 2712/2713.0 1                      |                  |
| T1270         | L A D S D L P V P P T G S A G E K<sup>3</sup> | 1708/1707.9 1                      |                  |
| T1350         | T T S A T P L K D N T Q F R<sup>4</sup> | 1954/1954.1 1                      |                  |
| In vivo       |                                               |                                      |                  |
| S775, S779    | V I I S D D P F V S S D T E L T S A V S S I N G E N L P T I I L S S T K S P T K N A E L V K<sup>1</sup> | 4918/4916.5 2                      |                  |

Average molecular masses of dephosphorylated peptides are shown.
was preferentially detected by the phosphopeptide antibodies.

**CB-associated p220 co-localizes with cyclin E and is phosphorylated on Cdk sites in a cell cycle-dependent manner**

The data described thus far suggest that p220 is targeted to CBs and is a substrate of cyclin E/Cdk2. However, it is unclear whether p220 is phosphorylated while in CBs. To examine whether CB-associated p220 is phosphorylated on Cdk2 sites, we performed immunofluorescence by using antibodies against phospho-T1270 and phospho-T1350. Both recognized nuclear foci similar to the antibody against p220 (Fig. 6a; data not shown). To demonstrate that the foci coincided with those found with anti-p220, we performed double immunofluorescence staining with antibodies against phospho-T1270 and coilin in cycling fibroblasts (Fig. 6). From 500 cells scored, 30.2% displayed primarily two foci reactive toward both antibodies, whereas 40.4% had primarily four co-localized foci (Fig. 6a), including cells in prophase (Fig. 6b). The remaining 29.4% of the cells lacked staining with the phospho-T1270 antibody. This is in marked contrast to staining with p220 antibodies in which cells lacking antibody reactivity were very rare except for those in metaphase and telophase (Figs. 2a and 3b). In cells that were negative for reactivity with anti-phospho-T1270, coilin reactivity was still observed (Fig. 6a). Quantification of relative DAPI intensity of 200 additional cells demonstrated that cells nonreactive with the phosphopeptide antibodies had a lower DNA content than did reactive cells, which is consistent with these cells being in the G1 phase, whereas phospho-T1270 antibody-positive cells had a higher DNA content consistent with S- or G2-phase cells (Fig. 2b). In this separate experiment, a somewhat higher percentage of cells had no detectable focus (data not shown). In agreement with the results with p220 antibodies, the few cells in metaphase and telophase were negative for staining with the phosphopeptide antibody, whereas coilin signals appeared dispersed (Fig. 6d,e).

If cyclin E is responsible for phosphorylation of p220 within CBs, one would predict that p220 would co-localize with cyclin E and that the timing of p220 phosphorylation would be coincident with co-localization. To examine this issue directly, we initially performed co-localization experiments using anti-cyclin E and anti-p220 antibodies. As shown in Figure 7a, cyclin E was concentrated in foci that are coincident with p220 foci. At

**Figure 6.** p220 in Cajal bodies (CBs) is phosphorylated on Cdk sites in a cell cycle-specific manner. Dual detection with anti-phospho-T1270 antibodies and p80coilin in dermal fibroblasts is shown. Focal co-localization was observed in S phase (a) and prophase (b), but antiphospho-T1270 did not detect any foci in metaphase (c) or telophase (d). a also contains three cells that display anti-coilin reactive foci but not anti-phospho-T1270 reactive foci. The DNA content of these cells is consistent with G1 phase (Fig. 2b). Only diffused coilin signals were observed in c and d. (Green) anti-phospho-T1270; (red) anti-coilin; (blue) 4’,6-diamidino-2-phenylindole (DAPI).
longer exposure, cyclin E was evident as faint dust throughout the nucleus as well. These results are consistent with the recent report that cyclin E is concentrated in CBs in S phase (Lui et al. 2000).

To examine whether phosphorylation of p220 correlates with co-localization with cyclin E, we pulse-labeled asynchronous diploid fibroblasts with BrdU and determined the presence of phospho-T1270, cyclin E, BrdU,
and DAPI-stained nuclei [Fig. 7b–e]. Among cells containing two phospho-T1270 foci, both BrdU-positive and BrdU-negative cells were observed, in both cases, however, these foci contained cyclin E [Fig. 7b–d]. In contrast, among cells containing four anti-phospho-T1270 foci, those that were BrdU positive most frequently displayed co-localized cyclin E, whereas those that were BrdU negative typically lacked cyclin E staining [Fig. 7d,e]. Given the data presented previously, we believe the latter class of cells to be G2 cells that have lost cyclin E expression but maintain p220 in a phosphorylated form.

The in situ results with antibodies to p220, phospho-T1270, phospho-T1350, coilin, and cyclin E [Figs. 1–3, 6; Liu et al. 2000] indicate the following: (1) p220 is an in vivo Cdk2 substrate and can be phosphorylated on cyclin E/Cdk2 sites while present in CBs. There is a tight correlation between the appearance of cyclin E in foci and the occurrence of p220 phosphorylation such that (2) cells that are nonreactive with antibody to phospho-T1270 are in the early G1 phase before cyclin E/Cdk2 is present to phosphorylate p220 in CBs. Consistent with this, cells that lacked anti-phospho-T1270 foci also lacked detectable cyclin E. (3) Cells containing two anti-phospho-T1270 reactive foci that co-localize with coilin and cyclin E are in late G1 or early S phases when cyclin E/Cdk2 peaks. (4) Cells that have four co-localized foci are well into S phase, and p220 remains phosphorylated [Figs. 1 and 2]. Although the pattern of p220 phosphorylation persists into prophase, cyclin E staining is lost at some point in late S phase or G2 phase, as exemplified by the presence of BrdU-negative cells containing four phospho-p220 foci but lacking cyclin E co-localization. (5) Around the time of metaphase and later, p220 foci are not detected with either p220 or phospho-T1270 antibodies.

**Mutation of cyclin E/Cdk2 sites in p220 reduces p220-mediated histone H2B promoter activation**

Our data suggest that cyclin E/Cdk2 may regulate p220 during the G1/S-phase transition. To examine this question, we took advantage of the recent finding that p220 expression in tissue culture cells leads to increased expression of histone 2B [H2B] promoter– and histone 4 [H4] promoter–luciferase reporter constructs, independent of its effects on S-phase acceleration [Zhao et al. 2000]. Histone gene expression is complex, involving both message stabilization [approximately sevenfold] and transcriptional activation [approximately fivefold], which together account for a ~35-fold increase in histone mRNA levels during S phase [Harris et al. 1991; Heintz 1991]. But the process by which the histone transcriptional apparatus senses cell cycle position is unknown.

We compared the ability of a vector expressing p220 [pCMV-p220] or p220ACdk [pCMV-p220ACdk] to activate luciferase expression from an H2B (~200/0) promoter–luciferase reporter plasmid in transiently transfected 293T cells [Fig. 8a]. The level of induction by p220 relative to control transfections ranged from two- to 10-fold, depending on the quantity of p220 plasmid used and the level of p220 expression achieved, as determined by immunoblotting [Fig. 8b]. In contrast, p220ACdk displayed a substantially reduced ability to activate the H2B reporter construct when expressed at comparable levels [Fig. 8a–e]. Similar results were obtained with a minimal H2B promoter (~127/~27) [data not shown]. At low levels of expression, p220ACdk displayed levels of H2B reporter activation comparable to control transfected cells [Fig. 8a,b], however, at higher levels of expression, a twofold increase in reporter activity over control transfected cells was typically observed [Fig. 8c,d]. Transfected cells displayed p220 foci as well as diffuse signals throughout the nucleoplasm because of elevated levels of expression from the transfected plasmids [Fig. 8e]. Taken together, these results suggest that phosphorylation at one or more cyclin E/Cdk2 sites contributes substantially to this aspect of p220 function. However, it is possible that elevated levels of p220 can partially bypass a requirement for phosphorylation at these sites. In these experiments, p220 appeared to be primarily in a more slowly migrating phosphorylated form, while p220ACdk remained in a more rapidly migrating form [Fig. 8b,d]. Thus, it appeared that sufficient cyclin E/Cdk2 existed in these cells to phosphorylate fully the transiently expressed p220. This may explain why co-expression of cyclin E/Cdk2 had little effect on the levels of H2B reporter activity in 293T cells [data not shown].

Consistent with a role for cyclin E/Cdk2–mediated phosphorylation in p220 function, we found that co-expression of p57KIP2, which can inhibit Cdk2 activity and block cells at the G1/S-phase transition [Matsuoka et al. 1995], led to a reduction in the ability of p220 to activate H2B promoter activity in transiently transfected cells [Fig. 8f]. In this experiment, the levels of p220 expression used were such that a twofold increase in H2B promoter activity was observed, but p57KIP2 reduced luciferase levels below that obtained with control transfected cells. As expected, expression of p57KIP2 alone also reduced the levels of promoter activity in the absence of p220 expression [Fig. 8f]. This repression likely reflects the fact that cells are blocked in G1 with low cyclin E/Cdk2 activity. Immunoblotting demonstrated comparable levels of expression of p220 and p220ACdk and verified the expression of p57KIP2 [data not shown].

**S-phase entry in quiescent fibroblasts by cyclin E/Cdk2 expression is associated with the accumulation of four p220 foci**

Expression of cyclin E/Cdk in quiescent fibroblasts leads to S-phase entry [Connell-Crowley et al. 1998; Leone et al. 1998]. Because the appearance of four p220 foci is associated with S phase in asynchronous and serum-stimulated cells [Figs. 1 and 2], we wondered whether S-phase entry by an alternative mechanism would also lead to the appearance of four p220 foci. To this end, quiescent WI38 cells were stimulated to enter the cell cycle by infection with adenoviruses (Ad) expressing cy-
clin E and Cdk2. At 24 h after infection, cells were pulse-labeled with BrdU for 1 h before analysis of p220 by immunofluorescence. BrdU-negative cells in control cultures maintained in low serum contained predominantly two p220 foci (Fig. 9a,b). In contrast, a large fraction of BrdU-positive cells in the cyclin E/Cdk2–treated culture contained four foci, whereas BrdU-negative cells in the culture contained predominantly two foci. These results suggest that cyclin E/Cdk2 can function upstream of the pathway responsible for the establishment of four p220 foci during S phase.

Discussion

Cell cycle transitions are driven in part by transcriptional programs that generate proteins needed for subsequent processes. Although it is clear that these transcriptional programs are ultimately linked to the basic cell cycle machinery, how this linkage is accomplished is largely unknown. The best understood connection between transcription and the basic cell cycle machinery in mammalian cells is the activation of E2F by Cdk-mediated phosphorylation of Rb family members (Dyson 1998; Nevins 1998). In this article, we provide evidence, at the cellular and molecular level, that cyclin E/Cdk2 directly regulates the activity of p220NPAT, which in turn controls S-phase-specific activation of histone gene transcription. Thus, cyclin E/Cdk2 not only regulates the production of DNA synthesis machinery through E2F but also supports the production of nucleosome components required for completion of DNA replication.

In normal fibroblasts, p220 is localized to discrete foci in the nucleus, and the number of these foci change during the cell cycle (Zhao et al. 2000; this work). Cells in G1 contain primarily two p220 foci, whereas cells in S phase contain four foci. Anti-p220-reactive foci are only absent during the short span of metaphase and telophase (Figs. 2, 3, and 6). These p220 foci coincide with small nuclear organelles, the CBs (Fig. 3). CBs contain a bewildering number of transcriptional and splicing/polyadenylation proteins, and recent studies have led to the hypothesis that CBs function as sites of assembly of pol II transcriptosomes, complexes of pol II transcription fac-
found little evidence for association with the domains of chromosomes 5, 17, or Y (Fig. 3). Typically, the p220 foci linked with chromosome 6 appeared to be larger than those associated with chromosome 1 (Fig. 3), potentially reflecting the larger numbers of histone genes located in the chromosome 6 cluster. The cell cycle–dependent association of p220 with CBs on chromosomes 6 and 1 then accounts for the oscillation in p220 foci numbers. Zhao et al. (2000) have also demonstrated that p220 is localized with chromosomes 1 and 6 and have shown that p220 is physically linked to histone gene loci.

On a biochemical level, p220 preferentially binds to cyclin E/Cdk2 over other Cdk complexes and is phosphorylated by cyclin E/Cdk2 in vitro and in vivo (Figs. 4 and 5). Indeed p220 foci also contain cyclin E (Fig. 7). Moreover, antibodies specific to p220 (Figs. 1–3) and to specific phosphopeptides of p220 (Figs. 6 and 7) demonstrated that p220 is present in CBs in both Cdk2 phosphorylated form and unphosphorylated forms and that these two forms alternate during the cell cycle. Three major classes of staining patterns were observed with phospho-specific antibodies against p220. Thirty percent of cells in an asynchronous population lacked phospho-T1270 antibody reactivity and displayed predominantly a G1 DNA content while maintaining detectable CBs. Because the vast majority of G1 cells contain two p220 foci co-localized with CBs (Figs. 1 and 2), we conclude that p220 in a distinct population of G1 cells is not phosphorylated on Cdk2 sites. This population of cells also lacked cyclin E staining, which is consistent with these cells being in early G1. Cells in the second class (30%) contain two phospho-T1270 antibody reactive foci that are co-localized with cyclin E. Our analysis indicates that these cells are in either late G1 or early S phase and suggests that chromosome 6–associated p220 can be phosphorylated on Cdk2 sites in advance of the formation of four obvious foci. In asynchronous cells, a small fraction of BrdU-positive cells have two clear p220 foci (Fig. 1c), suggesting that S phase can be initiated before the accumulation of p220 in chromosome 1–associated CBs. This idea is substantiated by the finding that some cells containing two anti-phospho-T1270 and anti-cyclin E reactive foci display partial replication, as determined by quantitation of DNA content or BrdU incorporation (Figs. 2b and 7). However, we cannot rule out the possibility that p220 is already present in chromosome 1–associated CBs but is present at levels below detection. Cells in a third class (40%) each contained four phospho-antibody reactive foci, and a substantial fraction of these cells contained co-localized cyclin E. These cells have a higher DNA content, consistent with p220 being phosphorylated in S and G2. p220 phosphorylation is maintained in prophase, with some cells displaying two foci and some displaying four foci, but p220 foci are absent in metaphase and telophase. Thus, p220 foci appear to be lost sequentially during the prophase-to-metaphase transition. The fate of p220 during mitosis is unclear at present. It could be dispersed and therefore beyond our means to detect at this stage in the cycle. Alternatively, p220 could be degraded. Consistent with

Figure 9. S-phase entry in quiescent fibroblasts by cyclin E/Cdk2 expression is associated with the appearance of four p220 foci. WI38 fibroblasts were subjected to serum deprivation for 72 h before infection with Ad-cyclin E/Cdk2 and maintained in 0.1% serum. Twenty-four h after infection, cells were pulse-labeled with BrdU for 1 h before analysis of p220 staining by immunofluorescence. (a) An example of an S-phase cell from a cyclin E/Cdk2 infection containing four p220 foci adjacent to a non-S-phase cell containing two p220 foci. Thirty to 100 nuclei of each class were counted.
the latter possibility is the finding that 293T cell extracts from mitotic cells (obtained by mitotic shake-off) have no detectable p220 by immunoprecipitation/immunoblotting analysis (data not shown). Thus, if p220 is degraded in mitosis, new p220 must be synthesized early in G1 phase and be incorporated into CBs in the unphosphorylated form before the activation of cyclin E/Cdk2 at late G1 and early S phase. In HeLa cells released from mitosis, p220 foci and colin foci reappear 2–3 h after release, consistent with the formation of foci in early G1 phase (data not shown).

The link between p220, a cyclin E/Cdk2 interacting protein (Zhao et al. 1998; Ma et al. 1999), and the transcription of histone 2B and histone 4 genes (Zhao et al. 2000) led us to address whether cyclin E/Cdk2 directly regulates this aspect of p220 function. In principle, cyclin E/Cdk2 could function to relay cell cycle positional information to p220, thereby playing a role in controlling the timing of S-phase-specific histone gene transcription. We found that p220 lacking five Cdk2 phosphorylation sites, four of which were phosphorylated in vivo, displayed a reduced ability to activate transcription from an H2B reporter construct (Fig. 8), consistent with a role for cyclin E/Cdk2 in p220 activation and H2B transcription. p220-dependent activation of the H2B promoter requires the Oct-1 element (data not shown) known to be involved in S-phase-specific induction of H2B transcription (Fletcher et al. 1987; Segil et al. 1991). Using a U2OS-based tissue culture system, Zhao et al. (2000) found that the ability of p220 to activate H4 transcription was stimulated by co-expression of cyclin E, again pointing to a role for cyclin E in this process. Although we did not observe a stimulatory effect of cyclin E/Cdk2 co-expression in our system, we found that the vast majority of ectopic p220 in 293T cells is in the slower mobility phosphorylated form (Fig. 8), suggesting that cyclin E is not a limiting component in these cells at the levels produced with our p220 expression plasmid. Apparently, at the levels of expression achieved in U2OS cells, cyclin E/Cdk2 is limiting. Regardless of these differences, both studies indicate that the cyclin E/Cdk2–mediated, cell cycle–dependent activation of p220 is an important component of the S-phase-specific histone transcriptional program.

Although cyclin E/Cdk2 activation seems to be central to p220 function, several issues remain to be addressed. First, what is the significance of p220 localization in CBs? Although p220 is clearly localized in these organelles and these organelles are physically linked to target genes, it is conceivable that the localization of p220 reflects its accumulation before release from CBs in a form that then activates histone gene transcription. Many transcription factors accumulate in inactive pools and are not present in detectable levels at target gene loci. Nevertheless, the co-localization of p220 with cyclin E, CBs, and histone gene clusters, the cell cycle–dependent phosphorylation of p220, the persistent association of phosphorylated p220 with CBs throughout S phase when histone genes are transcribed, as well as the activation of histone gene transcription by Cdk2-dependent p220 phosphorylation are most striking and strongly point to a functional role for p220 and cyclin E localization in CBs, as summarized in Figure 10. Second, what is the basis of the appearance of p220 foci on chromosome 1 during S phase, and how is this regulated? Presumably, the association of p220 foci with chromosome 1 reflects a role in S-phase-dependent histone transcription, but why then do p220 foci exist on chromosome 6 throughout most of the cell cycle? In this regard, it is important to determine whether transcription from endogenous histone genes is linked to accumulation of p220 at histone gene clusters. Third, the histone gene cluster on chromosome 6 contains >50 copies of the four classes of core histones as well as the linker histone H1 (Ahn and Gruen 1999). Does p220 coordinate the transcription of all classes of histone genes in the locus, and if so, how is this achieved? One possibility is that p220 could generate a chromosomal context that allows transcriptional activation of the whole region during S phase. If this is the case, p220 might also regulate the expression of nearby genes during S phase. Analysis of the 6p21 region reveals a large number of nonhistone genes, and it is possible that one or more of these genes are under the control of a p220-dependent S-phase transcriptional program. Alternately, p220 might function within the CB to assemble histone-specific transcription complexes, in keeping with the proposed function of CBs (Gall et al. 1999). Fourth, although p220 overexpression increases the S-phase population in transiently transfected cells (Zhao et al. 1998), it remains to be determined whether this activity is related to histone transcription and whether the phosphorylation events that we have identified are relevant to this activity. Finally, the finding that cyclin E is concentrated in CBs suggests the possibility that these organelles are important control centers linking the cell cycle machinery to S-phase-specific pro-

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cesses. It will therefore be important to determine whether other relevant cyclin E substrates gain access to the kinase through localization in CBs.

Materials and methods

Cell culture

Normal diploid fibroblasts (dermal fibroblasts, WI38, and BJ/TERT), 293T, and HeLa cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS). To generate quiescent fibroblasts, cells were plated at ~50% confluence before culture for 72 h without serum. Cells were released into DMEM containing 10% FBS for various lengths of time. In some experiments, the cells were exposed to BrdU (10–50 µg/mL) for 1 h before harvest to reveal cells in S phase. To examine cell cycle entry via cyclin E/Cdk2 expression, WI38 cells were maintained for 72 h in 0.1% FBS and infected with adenoviruses expressing cyclin E and Cdk2 (generously provided by J. Nevins, Duke University, Durham, NC) [Leone et al. 1998] at a multiplicity of infection of 100.

Antibodies and immunofluorescence assays

Bacterial GST-p220 [residues 1054–1397] was used to generate antibodies in rabbits. Antibodies were depleted of reactivity to the GST protein and affinity-purified using immobilized GST–p220. Anti-colchin monoclonal antibodies [α-isotype] were provided by M. Carmo-Fonseca (University of Lisbon, Portugal; Almeida et al. 1998). Anti-cyclin E [HE12] came from Pharmingen. Antibodies against Thr-1270 [Asp-Leu-Pro-Val-Pro-Arg-Ala-Ala-Val-Thr-Arg-Thr-Ser-Ala-phosphoThr-Pro-Leu-Lys-Asp-Asn-Thr-Cys] and Thr-1350 [Ser-Arg-Thr-Thr-Ser-Ala-phosphoThr-Pro-Leu-Lys] were generated in rabbits after coupling to keyhole limpet hemocyanin. For immunofluorescence, cells were fixed in either ethanol or formalin and permeabilized with 0.1% Triton X-100. Detection of rabbit antibodies was accomplished using secondary antibodies in rabbits. Antibodies were depleted of reactivity to human, mouse, and rat IgG by affinity-purified anti-p220 antibodies bound to protein A-Sepharose. The slides were DAPI stained before imaging.

Interphase chromosome painting

Asynchronous primary human dermal fibroblasts were fixed and stained with antibodies to p220, and signals were developed with tyramide as described above. Slides were fixed again with 4% paraformaldehyde and treated with RNase A (100 µg/mL) in 2× SSC (1× SSC is 0.15 M NaCl and 0.015 M sodium citrate) for 1 h at 37°C. They were then ethanol-dehydrated and denatured in 70% formamide, 2× SSC, pH 7.0, for 2 min at 72°C, dehydrated in ethanol, and hybridized individually with chromosome paints overnight at 37°C. The chromosome 1 and chromosome 17 paints were a direct FITC conjugate or cyanine 3 conjugate, respectively, from Vysis (Downers Grove, IL). Signals were detected according to the manufacturer’s protocol. For chromosome 5, 6, or 17 were digoxigenin probes from Oncor (Gaithersburg, MD). They were detected using an anti-dig Texas Red antibody. The slides were DAPI stained before imaging.

Immunoprecipitation and phosphorylation

For nuclear extracts, 293T cells were lysed in 50 mM Tris-HCl, pH 7.5, containing 1 mM EDTA, 100 mM NaCl, 0.3% Nonidet P-40, and protease/phosphatase inhibitors, and nuclei pelleted by centrifugation. Nuclear proteins were solubilized using RIPA buffer (50 mM Tris-HCl at pH 7.5, 150 mM NaCl, 0.5% deoxycholate, 1% Nonidet P-40, 0.1% SDS, and protease/phosphatase inhibitors) before centrifugation. Extracts were precleared with protein A–bound normal rabbit IgG before incubation with affinity-purified anti-p220 antibodies bound to protein A-Sepharose. Beads were washed with RIPA buffer. Proteins were separated by SDS–PAGE before staining with Coomassie Brilliant Blue R 250 or transferred to nitrocellulose filters for immunoblotting with anti-p220 antibodies. Detection was accomplished using enhanced chemiluminescence (Amersham).

For expression of p220 in insect cells, the coding sequence [Imai et al. 1996] was cloned into pUNI-50, and in vitro recombination was then used to generate a pVL1392-based plasmid with p220 fused at its N terminus to the Flag tag as described [Liu et al. 1998]. Viruses were made using Baculogold [Pharmingen]. To examine interaction with Cdks, cyclin/Cdk complexes were purified with glutathione-Sepharose beads [Ma et al. 1999]. Immobilized complexes were then incubated with insect cell extracts with or without Flag-p220. Complexes were washed with lysis buffer and with 20 mM Tris-HCl, pH 7.5, and 10 mM MgCl2. A portion of each mixture was used for kinase assays employing 50 µM γ-[32P]ATP. Samples were separated by
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SDS–PAGE and transferred to nitrocellulose before immunoblotting and autoradiography.

Matrix-assisted laser desorption/ionization mass spectrometry [MALDI/TOF] with delayed extraction [Voyager-DE, Perseptive Biosystems, Framingham, MA] was used for the identification of phosphopeptides, as described by Zhang et al. [1998]. An electrospray ion trap mass spectrometer (LCQ, Finnigan, San Jose, CA) coupled on-line with a capillary high-precision liquid chromatograph [Magic 2002, Auburn, CA] was used for identification of phosphorylation sites. A MAGICS C18 column [5 μm particle diameter, 150 Å pore size, 0.1 X 50 mm dimension] was used for the LC/MS/MS analysis.

Plasmids and reporter assays

Mutations were generated using a Gene Editor kit [Promega]. Mutated and wild-type p220 coding sequences were cloned into pCDNA3.1 for expression. To generate histone gene reporter plasmids, H2B regulatory sequences (~200/0 or ~127/~27) were cloned into the luciferase reporter pGL3 [Promega]. To examine histone transcription, pCMV-p220, pGL3 reporter plasmids, and pCMV-β-galactosidase plasmids were co-transfected into 293T cells at 50% -galactosidase activity before measurement of luciferase activity.

Acknowledgments

We thank J. Gall for discussions, J. Zhao, A.G. Matera, and E. Harlow for communicating results before publication, M. Carmo-Fonseca for anti-coilin antibodies, and Richard Atkinson, Brian Streib, and Heather Benedict-Hamilton for assistance. J.W.H. was supported by U.S. Public Health Service (USPHS) grant GM54137 and by the Welch Foundation. L.T.C. was supported by USPHS grants CA36200 and DE/CA11910. B.A.V.T was partially supported by the University of Alabama Medical Scientist Training Program. The Digital Imaging Microscopy Facility at the University of Alabama was supported by grant DE/CA11901.

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Cell cycle–regulated phosphorylation of p220NPAT by cyclin E/Cdk2 in Cajal bodies promotes histone gene transcription

Tianlin Ma, Brian A. Van Tine, Yue Wei, et al.

*Genes Dev.* 2000, 14:
Access the most recent version at doi:10.1101/gad.829500

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