BIOCHEMISTRY


The authors note that, due to a printer’s error, on page 16967, the keyword “mas spectromrtry” should instead have appeared as “mass spectrometry.” The online version has been corrected.

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CELL BIOLOGY


The authors note that the author name Andrew Kung should instead have appeared as Andrew L. Kung. The online version has been corrected. The corrected author line and author contributions footnote appear below.

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ECOLOGY


The authors note that on page 16298, right column, first full paragraph, the first sentence appeared incorrectly in part. “In the rocky intertidal habitats dominated by mussel beds (dark colored) and barnacle stands (light colored), we found that more than 95% of early postmetamorphic stages of C. concholepas (~2 to 20 mm periostomal length, PL) showed a striking color-matching with the most abundant prey (Fig. 2A and B)” should instead have appeared as “In the rocky intertidal habitats dominated by mussel beds (dark colored) and barnacle stands (light colored), we found that more than 95% of early postmetamorphic stages of C. concholepas (~2 to 20 mm periostomal length, PL) showed a striking color-matching with the most abundant prey (Fig. 2A and B).” Additionally, the authors note that on page 16301, right column, first full paragraph, the second sentence did not appear in full due to a printer’s error. The sentence should instead have appeared as “This modulation of the shell coloration is restricted to snails of small size (ca. less than 3 cm), coincidentally the size at which C. concholepas appear to escape predation by crabs (author’s personal observations).” These errors do not affect the conclusions of the article.

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CBP and p300 are cytoplasmic E4 polyubiquitin ligases for p53

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The ubiquitination and proteasome targeting of p53 by the RING E3 enzyme MDM2 has been considered to generally inhibit p53 function (6). However, the distinction between multiple monoubiquitination (MUM) and polyubiquitination of p53 adds additional complexity to its regulation. MUM has been reported to enhance p53 nuclear export, and in the absence of stress, p53 is very likely polyubiquitinated and then degraded by the proteasome in the cytoplasm (7, 8). In addition, recent evidence points to a potential positive role for p53 ubiquitination in its capacity as a nuclear transcription factor (4, 9, 10). Adding to the difficulty of understanding p53 ubiquitination, a multitude of ubiquitin ligases (E3s) have been identified for p53 in untransformed cells besides MDM2, most notably E4F1 (10), COP1 (11), and ARF-BP1 (12), making it unclear how each E3 contributes to overall p53 regulation in any given cell-type or environmental condition (13).

Whether p53 MUM vs. polyubiquitination is actively and specifically controlled in cells also remains unclear. In stressed cells, the induction of MDM2 expression by activated p53, can specifically increase the abundance of polyubiquitinated p53 adducts (8). In unstressed cells, specific polyubiquitin ligases (E4s), or ubiquitin (Ub) chain-extending factors, exist for p53—namely the coactivator p300 (14) and transcription factor YY1 (15). p300-dependent polyubiquitination of p53 in vitro required priming of p53 by MDM2-driven p53 MUM (16), although p300, and its paralog CREB-binding protein (CBP), both exhibited robust E3 autoubiquitination activity (14). YY1, by contrast, stabilized MDM2/p53 complexes, allowing MDM2 to act more processively toward chain elongation (15). A recognizable E3 or E4 domain could not be identified in p300/CBP, but the N-terminal 595 aa of p300 appeared to harbor both its E3 (autoubiquitination) and E4 activities (14). Recently, Zn-finger like domains have also been characterized as active E3 enzymes (10, 17), and p300 does harbor three conserved Zn2+−binding Cys-His-rich regions, one of which (C/H1-TAZ1) lies within its first 595 aa (18–20).

To better define the role of p300 or CBP in p53 stability regulation, p53 turnover was studied in p300- or CBP-deficient cells. p300 or CBP depletion in untransformed cells led to p53 stabilization, and p300 and CBP E3 activities were exclusively cytoplasmic, with detectable amounts of both proteins localized to the cytoplasm. CBP, like p300, encoded an E3 activity within its N terminus and exhibited E4 activity toward p53 in vitro. p300 and CBP therefore engage in compartmentalized regulation of p53, by cytoplasmic ubiquitination and presumed nuclear acetylation, that is required for the proper homeostasis of p53 in basal and stressed conditions.

Results

Regulation of p53 Abundance and Stability by p300 and CBP. To assess the physiologic contributions of p300 and CBP to p53 regulation in unstressed cells, p300 and CBP were transiently silenced in U2OS cells with specific siRNA duplexes, followed by analysis of cell lysates for p300, CBP, and p53 levels, as well as determination of p53 half-life (t1/2) by cycloheximide decay (Fig. 1A and B). p300 siRNA has been reported to increase p53 abundance but the mechanism was not determined further (21). p300 and CBP siRNAs both caused increases in steady-state p53 abundance (Fig. 1A) and p53 half-life (from 75 min to >2 h and 2 h, respectively; Fig. 1B) compared to that seen in control siRNA-treated cells. Similar gains in p53 abundance were also seen after stable CBP knockdown by two distinct shRNA sequences (CBP-shA, CBP-shB), and CBP-shA also induced p53 stabilization (Fig. S1A). Demonstrating that these results were not cell type-specific or due to the use of a cancer cell line, siRNA depletion of CBP or p300 in nontransformed MCF10A human breast epithelial cells similarly stabilized p53 from t1/2 = 75 min with control siRNA to >2 h for both p300 and CBP siRNAs (Fig. S1B).

Indirect explanations for the increased p53 abundance and stability induced by p300/CBP depletion, such as increased p53 mRNA expression, decreased MDM2 expression, or induction of genotoxic stress (the latter two of which might stabilize p53) were investigated. RT-PCR analysis showed that p53 mRNA levels were increased by p300 depletion (Fig. S2).


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CBP was silenced, with polyubiquitination decreasing and mono-abundance of polyubiquitinated vs. mono-Ub/MUM p53 was quantitated by use of non-chain-forming Ub moieties (14). Input lysates were blotted with the indicated antibodies.

were actually decreased, not increased, in CBP-shA as compared with control-sh cells (Fig. S1C), and MDM2 protein levels were increased or unchanged, not decreased, in CBP-shA or CBP-shB cells as compared with control-sh cells (Fig. S1A). The possibility that CBP depletion might stimulate a p53-stabilizing stress response (22) was investigated by mock or doxorubicin (Dox)-treatment (as positive control for stress stimulus) of control-sh or CBP-shA cells, followed by immunoblotting of lysates for p53 abundance of p53-Ub conjugates, a marker of p53 activation in response to stress (Fig. S1D) (23, 24). The untreated control lines had an equivalent and very low level of serine 15 phosphorylation, and both showed robust induction of phosphorylation after treatment with Dox for 2 h (Fig. S1D). Thus, the lack of basally induced p53 S15 phosphorylation in unstressed CBP-depleted cells does not support a nonspecific stress mechanism as the reason for the observed p53 stabilization.

**CBP Is Required for p53 Polyubiquitination In Vivo.** Given that p300 can act as an E4 for p53 in vitro (14), the abundance of p53-Ub conjugates was analyzed in U2OS cells expressing HA-tagged Ub after control, CBP, or p300 siRNA treatment, and in the presence or absence of MG132 proteasome inhibitor. Lysates of siRNA-treated cells were analyzed by p53 IP under stringent conditions, followed by HA immunoblot (Fig. 1 C and D). Polyubiquitinated p53-Ub conjugates were detected in a p53 IP of lysate from control siRNA-treated cells, as evidenced by the broadly distributed HA immunoreactive species (Fig. 1C, first lane). Based on the pattern of p53-Ub conjugates seen in vitro when non-chain-forming Ub moieties are used, those p53-Ub conjugates <100 kDa represent largely mono-Ub and MUM species, while those >100 kDa represent polyubiquitinated species (14), p53 IPs from lysates of p300 or CBP siRNA-treated cells, by contrast, had significantly reduced levels of polyubiquitinated p53 (Fig. 1C, second and third lanes). Similar results were seen after proteasome inhibition, except for a stronger polyubiquitinated signal in the p53 IP from control siRNA-treated cells (Fig. 1D). Similar results were also seen when the p53 ubiquitination pattern was compared between control-sh and CBP-shA cell lines expressing HA-Ub (Fig. S2A). In this case, the relative abundance of polyubiquitinated vs. mono-Ub/MUM p53 was quantitated showing an inverse correlation between the two forms when CBP was silenced, with polyubiquitination decreasing and mono-Ub/MUM forms increasing (Fig. S2A). p300 and CBP are therefore both required to maintain physiologic levels of polyubiquitinated p53 conjugates in unstressed U2OS cells.

**CBP N-Terminal Sequences Destabilize p53.** To map the portion of CBP responsible for p53 destabilization, vector control, full-length or truncated CBP cDNAs were expressed as “rescue” alleles in CBP-deficient CBP-shA cells, and p53 stability was measured by cycloheximide-decay analysis. Full-length CBP* (mutated in the shRNA target sequence) expression in CBP-shA cells restored p53 half-life to the normal 60 min from the 120 min observed in CBP-shA cells (Fig. S2B). Given that the p300 N terminus has been proposed as necessary and sufficient for E4 activity (resides 1–595) (14), the homologous portion of the CBP N terminus was analyzed in detail for its ability to rescue p53 instability in CBP-shA cells (Fig. 2A). CBP (1–616) and (1–200), which do and do not, respectively, include the C/H1-TAZ1 domain (resides 348–432), were expressed in CBP-shA cells (Fig. S2C), p53 was destabilized from a t1/2 of 110 min seen in CBP-shA cells to 55 min in cells expressing CBP (1–616) (Fig. 2A), while the p53 t1/2 remained similar to the value in CBP-shA cells when CBP (1–200) was expressed (100 min) (Fig. 2A). Thus, the CBP N terminus is sufficient to destabilize p53, and the sequence between residues 200 and 616, including the C/H1-TAZ1 domain, is required for this activity.

**The CBP N Terminus Exhibits E4 Activity In Vitro and In Vivo.** Given the suggestion from the in vivo data that CBP and p300 both exhibit an E4 function in vivo, we verified that CBP, like p300 (14), could act directly as an E4 in a reconstituted in vitro p53 ubiquitination reaction using purified components (Fig. 2B). p53-MUM conjugates were prepared using purified E1, ubiquitin (E2), Ub, and limiting quantities of purified recombiant MDM2 (14). When purified FLAG-CBP was added to this reaction, there was a dose-dependent increase in the abundance and the maximum molecular weight of high molecular weight species of p53, consistent with polyubiquitination (polyubiquitin conjugates are assumed at a molecular weight of p53 species >100 kDa) (14) (Fig. 2B). Control reactions lacking MDM2 revealed no ubiquitination activity of p300 or CBP toward native p53 (Fig. S3, right), and eliminating p53 from the reaction eliminated all signal on the p53
The CBP N Terminus Regulates Ubiquitination and Stability. (A) CBP shA cells were transfected with vector or the indicated CBP rescue alleles, treated with cycloheximide for 30 min after transfection, and lysed analyzed by p53 immunoblot. (Bottom) p53 levels were quantitated by densitometry, and half-life calculated based on decay of normalized (to GAPDH loading control) p53 levels to 50% of their original level. Values are an average of three independent experiments. Error bars, ±1 S.D. (B) Purified recombinant CBP protein catalyzes p53 polyubiquitination. (Top) Purified recombinant p53-MDM2 protein complexes were preincubated with ubiquitin reaction components at 37 °C for 30 min. Purified FLAG-CBP (1, 10, or 100 ng) was then added, followed by further incubation for 1 h at 37 °C. Reactions were analyzed by p53 immunoblot. A shorter exposure (middle) shows the unmodified p53 protein level. (Bottom) The ratio of high molecular weight (>100 kDa) p53 species: Total p53 abundance was plotted at 20-kDa intervals for the indicated conditions. (C) The CBP N terminus is an active E4 in vivo. H1299 cells were transfected with His-Ub vector, p53 or p53-UbF expression vectors, along with the indicated CBP plasmids or empty vector. (Top) Forty-eight hours after transfection, lysates were purified on Ni-NTA beads, and eluted products immunoblotted with anti-p53 antibody. (Bottom) Input lysates were immunoblotted with anti-FLAG and p53 antibodies. The abundance of high molecular weight p53-Ub conjugates was determined by scanning densitometry and normalized to total amount of p53UbF in the input lysate. Experiment shown is representative of three independent experiments. (D) Sequential E4 assay. Purified FLAG-MDM2/HA-p53 complexes were exposed to E1, E2, Ub, and ATP to generate mono-Ub/MUM p53. HA-p53-Ub conjugates were purified by anti-HA IP and stringent washing, and then exposed to E2 and ATP a second time with or without E1, along with FLAG-Ub and affinity purified CBP (1–616) or (1–200). Further washes, HA-p53-Ub conjugates were analyzed by anti-FLAG immunoblot.

immunoblots (Fig. S3A, left). To quantitate the E4 effect of CBP, the ratio of p53 signal in the high molecular weight range (>100 kDa) of the gel to total p53 signal was calculated at 20-kDa intervals between 100 kDa and the top of the gel, and plotted vs. mean molecular weight (Fig. 2B, bottom). The clear separation of the control (MDM2-only) curve from the CBP titration curves reveals that CBP had a clearly quantifiable impact on Ub content and, by inference, Ub chain length of p53-Ub conjugates.

To demonstrate that the CBP N terminus could also act as an E4 for p53-mono-Ub or MUM conjugates in vivo, p53 or p53-Ub fusion (Ub cDNA fused in frame to the p53 C terminus; p53-UbF) protein cDNAs were expressed in cells, to determine if p53-UbF could serve as a model for p53-monoUb or MUM conjugates, and act as a direct substrate for the CBP E4 activity. His-Ub and CBP (1–200) and (1–616) polyepitopes were expressed along with p53 or p53-UbF in H1299 cells, followed by Ni-NTA pulldown of lysates and p53 immunoblotting. Analysis of lysates used for Ni-NTA pulldown by FLAG and p53 immunoblot revealed even expression of CBP (1–200) and CBP (1–616) (Fig. 2C, middle), as well as equivalent expression of p53 and p53-UbF (Fig. 2C, bottom). Coexpression of the CBP N terminus with native p53, which is not even multiply monoubiquitinated in H1299 cells (Fig. 2C, top, first lane) had no effect, as ubiquitinated p53 remained undetectable (Fig. 2C, top, second and third lanes). Coexpression of CBP (1–616) with p53-UbF resulted in robust induction of high molecular weight Ub conjugates of the p53-Ub fusion, as compared with the effects of control (empty vector) or CBP (1–200) coexpression, both of which were associated with a similar background level of p53UbF ubiquitination signal (Fig. 2C). The effect of the CBP N terminus on the ubiquitination status of the p53-Ub fusion specifically demonstrates that the first 616 aa of CBP are sufficient to polyubiquitinate p53-monoUb or MUM conjugates in vivo as well as in vitro.

CBP E4 Activity Is Separable from MDM2. To definitively delineate the independent contribution of CBP to p53 polyubiquitination, a two-step in vitro E4 assay was developed. In the first step, p53-MUM conjugates were prepared by incubating with MDM2 in a similar fashion as seen in Fig. 2B, fourth lane, using purified FLAG-MDM2/HA-p53 complex from insect cells. The reaction products were immunopurified on anti-HA-conjugated beads under stringent (RIPA buffer) conditions to remove MDM2. CBP (1–616) or (1–200) polypeptides were then added to the immobilized HA-p53 conjugates with fresh E1, E2, ATP, and FLAG-ubiquitin, followed by washing of the beads and blotting of the reaction products with anti-FLAG antibody. As is seen in Fig. 2D, CBP (1–616), but not (1–200), clearly catalyzed the synthesis of polyubiquitin chains when p53 had previously mono- or multiply monoubiquitinated by MDM2. Moreover, no FLAG-MDM2 signal was seen among the reaction products (absence of signal at 90-kDa section of blot; Fig. 2D), confirming that MDM2 had been effectively removed before the addition of CBP polypeptides. Thus, CBP (1–616) encodes an MDM2-independent p53-directed E4 function.

The CBP N Terminus Encodes an Intrinsinc E3 Activity. The E4 effect of CBP would be most consistent with an intrinsic MDM2-independent ubiquitin ligase activity that can only recognize a monoubiquitinated or multiply monoubiquitinated substrate, as has
been observed for p300 (14). We investigated whether sequences arising from the CBP N terminus, which demonstrate E4 activity (Fig. 2 C and D) and promote p53 instability in vivo (Fig. 2 A), encode an intrinsic E3 activity that could be clearly separated from MDM2. Prokaryotically synthesized, purified, GST-CBP (1–616), (1–451), and (452–721) were incubated with Ub, E1, E2 (ubch5a), and ATP in an autoubiquitination reaction, and the reaction products detected by anti-Ub immunoblot. Titration of GST-CBP (1–616) or (1–451) with standard ubiquitin reaction components, revealed robust E3 activity (Fig. S3C), GST-CBP (452–721), however, was inactive as an E3 (Fig. S3C). Thus, the CBP polypeptide that promotes p53 degradation in vivo and p53 polyubiquitination in vivo and in vitro, also encodes an intrinsic, MDM2-independent, E3 ubiquitin ligase domain. The core sequences responsible for this activity appear to lie within at least the N-terminal 451 aa of CBP, and include the C/H1-TAZ1 domain.

CBP Promotes p53 Degradation in the Cytoplasm. Since p300/CBP are considered to be nuclear coactivators (25), but p53 and MDM2 both shuttle between nucleus and cytoplasm (26), the cellular localization of CBP’s regulation of p53 ubiquitination and degradation was explored further. Levels of unmodified p53 were proportionately increased in both nuclear and cytoplasmic fractions in CBP-shA vs. control cells, and controls for nuclear (Rb) and cytoplasmic (actin) proteins revealed that there was little cross-contamination of the fractions (Fig. 3A). Mono-Ub/MUM p53 was found predominantly in the cytoplasm, and levels of this species in the cytoplasm were significantly increased in CBP-shA vs. control cells, either due to the overall increase in total p53, or a loss of an (E4) activity required to convert p53-mono-Ub or p53-MUM conjugates to polyubiquitinated forms (Fig. 3A).

To determine if the differential ubiquitination of nuclear and cytoplasmic p53 correlated with different rates of turnover, nuclear and cytoplasmic p53 half-life was measured by cycloheximide decay in control and CBP-shA cells (Fig. 3B and Fig. S3D). PARP (nuclear marker) and α-tubulin (cytoplasmic marker) immunoblots of the fractions from each time point revealed negligible cross-contamination (Fig. 3B). Surprisingly, CBP silencing led to stabilization of the normally rapidly degraded cytoplasmic p53 (t1/2 increase from 50 to 85 min, P = 0.01), but had no significant effect on the slower turnover of nuclear p53 (t1/2 increase from 95 to 100 min, P = 0.63; Fig. 3B and Fig. S3D). Given that p300/CBP are considered to be nuclear proteins (27, 28), but the major effect of CBP depletion was on cytoplasmic p53 half-life, the possibility that cytoplasmic pools of CBP or p300 might influence p53 metabolism was explored.

CBP/p300 Are Partly Cytoplasmic. CBP and p300 are localized in PML oncogenic domains (PODs) and senescence-associated nuclear bodies, respectively (27, 29). To gain a more precise understanding of p300 and CBP subcellular localization, their abundance in cytoplasmic and nuclear fractions of U2OS cells was determined (Fig. 3C). The nuclear-cytoplasmic abundance ratios of Rb and actin, as determined by densitometry, revealed that the upper limit of cross-contamination of one fraction into the other was no greater than 10% in either direction (Fig. 3C). CBP and p300 were predominantly nuclear, as expected, but a substantial fraction of CBP was cytoplasmic, whereas a small but reproducible fraction of p300 was also observed in the cytoplasmic fraction (Fig. 3C). These fractionation results were further confirmed by immunofluorescent staining of U2OS cells for CBP and p300, where a clear cytoplasmic signal was detected for CBP, but p300 appeared predominantly nuclear (Fig. S3E). The specificity of the CBP immunofluorescence signal was confirmed by the near complete loss of CBP staining seen in CBP-shA cells (Fig. S3E).

CBP/p300 E3 Activities Are Exclusively Cytoplasmic. To determine if nuclear and cytoplasmic CBP or p300 molecules have any differential capacities to influence p53 ubiquitination, nuclear and cytoplasmic CBP/p300 were assayed for E3 autoubiquitination activity. Nuclear and cytoplasmic fractions of U2OS cells were IP’d with anti-CBP or p300 antibody, and the IPs were assayed for E3 activity by the addition of ubiquitin, E1, E2, (ubch5a), and ATP, followed by Ub, CBP, or p300 immunoblotting (Fig. 4A and S4A). Dropout control reactions lacking E1/E2, Ub, or ATP were also performed to demonstrate that any Ub chains observed were the result of bona fide in vitro E3 activity and not contamination by cellular Ub chains binding nonspecifically to the IPs (Fig. 4A and S4A). CBP and p300 were observed in the cytoplasm and nuclear fractions as expected, and controls indicated little cross-contamination (Fig. 4A, bottom, and Fig. S4A, bottom). Surprisingly, CBP and p300 derived from cytoplasm demonstrated robust E3 autoubiquitination activity dependent on the simultaneous addition of E1/E2, Ub, and ATP, whereas nuclear-derived CBP/p300 exhibited much lower or undetectable activity (Fig. 4A and S4A). CBP/p300 immunoblotting of the reactions revealed that the lack of E3 activity associated with nuclear CBP/p300 could not be explained by lesser abundance (Fig. 4A and S4A). The lack of nuclear CBP autoubiquitination activity could also not be explained by nonspecific inhibition by extraction conditions or inhibitory factors within the nuclear fraction, as immunopurified nuclear MDM2 exhibited easily detected autoubiquitination activity, which if anything, had a higher specific activity than
cytoplasmic MDM2, based on the relative abundance of MDM2 in the two fractions (Fig. S4B).

To further assess the specific dependence of the observed E3 activity on the actual presence of CBP or p300 in the reaction, cytoplasmic fractions of control, CBP-shA, and p300-sh cells were IP'd with anti-CBP or anti-p300 antibodies, as appropriate, and the IPs assayed for E3 activity with or without added E1 and E2 (Fig. S4 C and D). There was no significant cross-contamination of nuclear and cytoplasmic markers (Fig. S4 C and D, bottom), and the IPs were also probed for p300 and CBP to assure that the levels of p300/CBP in the IPs reflected the respective shRNA knockdowns for each (Fig. S4 C and D, middle). CBP and p300 IPs from control cell cytoplasm were both highly enriched for E3 autoubiquitination activity dependent on added E1/E2 (Fig. S4 C and D, upper). However, CBP and p300 IPs from both CBP-shA and p300-sh cytoplasm were substantially depleted of E3 activity (Fig. S4 C and D, upper), confirming that the E3 activity seen in the p300 and CBP IPs from control cell cytoplasm did depend on the specific presence of p300 or CBP proteins in the respective reactions.

**Fig. 4.** p300/CBP E3 activities are predominantly cytoplasmic. (A) (Top and Middle) Nuclear and cytoplasmic fractions of U2OS cells were immunoprecipitated with anti-CBP antibody, and the washed IPs incubated with E1/E2, Ub, and ATP as indicated, followed by anti-Ub and CBP immunoblotting of the reactions. (Bottom) Immunoblots of CBP and Rb (nuclear marker) in nuclear and cytoplasmic fractions. (B) (Top and Middle) Nuclear and cytoplasmic fractions of insect-cell-derived p53/MDM2 complexes (insect-cell-derived) were incubated with E1/E2, Ub, or methyl-Ub, ATP and CBP IPs from control-sh or CBP-shA cytoplasmic fractions as indicated, followed by anti-p53 (Top) or anti-CBP (Middle) immunoblotting of the reactions. Migration positions for native, mono-Ub/MUM and polyubiquitinated p53 species are indicated. (Bottom) Relative abundances of native, mono-Ub/MUM, and polyubiquitinated p53 species were quantitated by densitometry.

**Cytoplasmic CBP Is an Active p53 E4.** The specific localization of E3 autoubiquitination activity of CBP/p300 to the cytoplasm suggests that their E4 activity, likewise resides in the cytoplasm. To confirm that cytoplasmic CBP was an active E4 for p53, immunopurified cytoplasmic CBP derived from control-sh or CBP-shA cells was exposed to MDM2/p53 complexes in the presence or absence of E1/E2, along with Ub and ATP (Fig. 4B). To identify the migration positions of p53-mono-Ub/MUM species, p53/MDM2 complexes were incubated with E1/E2, methyl-Ub (nonconjugatable), and ATP (Fig. 4B, first lane). CBP IP'd from control-sh cells clearly catalyzed the formation of polyubiquitinated p53 (p53-polyubiquitin conjugates ~50% of total p53 species; Fig. 4B), and this activity was significantly reduced in proportion to the reduction in CBP abundance when a CBP IP from CBP-shA cells was used as the source for cytoplasmic CBP (p53-polyubiquitin conjugates ~20% of total p53 species; Fig. 4B). Thus, cytoplasmic CBP is an active E4 for previously mono/multiply monoubiquitinated p53.

**Discussion**
p300 and its paralog CBP have been identified as physiological regulators of p53 ubiquitination and stability. Depletion of CBP or p300 resulted in p53 stabilization, and CBP and p300 were both required for physiologic p53 polyubiquitination. The CBP N terminus (1-616) was found to encode a p53 destabilization function in vivo, and residues 1-616 and 1-451, respectively, constitute minimal functional E4 and E3 domains within CBP. CBP and p300 E3 activities were only observed from cytoplasmic fractions, and p53-mono-Ub and MUM conjugates accumulated in the cytoplasm of CBP silenced cells. The degradation of cytoplasmic p53 was specifically altered by CBP silencing, consistent with the cytoplasmic localization of CBP E4 function in vivo.

The N terminus of p300 harbors its E3/E4 activity (14), and the same region of CBP encodes its E3/E4 activity and shares significant areas of sequence conservation (18, 30). Within this region of p300/CBP is a cys-his-rich region, but no signatures for RING, HECT, U-box, or PHD domains can be found. Based on findings with the A20, Rabex-5, and E4F1 proteins (10, 17, 31), a generic Zn$^{2+}$-binding region (cys- or cys-his-rich) can encode an active E3, with no clear sequence or structural homology to the Zn$^{2+}$-binding RING E3 domain. Perhaps more directly relevant, the evolutionarily related HAT P/CAF was recently shown to encode a noncanonical E3 that targets MDM2 (21). In this case, the protein domain responsible for the activity contains no structural motifs found in known E3 proteins (21).

Subcellular fractionation revealed that p300/CBP E3 activities are limited to the cytoplasm and presumably account for the activity that converts mono-Ub/MUM p53 conjugates to polyubiquitinated unstable p53 in the cytoplasm. The mechanism of localization of p300/CBP E3 activity to the cytoplasm cannot yet be explained. Either a cytoplasm-specific modification of p300/CBP or p300/CBP-interacting factor activates the activity or an interacting factor or modification in the nucleus represses it. The spatial separation of p300/CBP's E3/E4 activity from their nuclear HAT functions is intriguing and solves the seeming paradox of how these two presumed opposing regulatory functions exist within the same molecules (32).

This work raises a number of questions critical to p53, ubiquitin/proteasome, and cancer biology. How the plethora of p53 E3s, and now E4s, are coordinated to achieve p53 homeostasis remains unclear, although it is likely that the subcellular localization of these enzymes will be of critical importance to understanding their roles. The biochemical mechanism by which CBP/p300 recognizes ubiquitinated, and not native, p53 for further ubiquitination is also presently not understood. The simplest explanation for the preferred recognition of ubiquitinated substrate would be the existence of a Ub recognition motif within the p300/CBP N-terminus—as is seen in the Rabex-5 atypical E3 (31), although no such domain is readily recognized in p300/CBP by homology search. Finally, the implications of these findings for cancer biology and therapy are potentially significant. As MDM2 is now a bona fide clinically relevant drug target for cancer therapy (33), other enzymes in the p53 stability regulation pathway, such as the p300/CBP E4 domain, might also be effectively targeted in human cancers.
Materials and Methods

Cell Culture and Plasmids. U2OS cells were grown in DMEM supplemented with 10% FBS and antibiotics. Cells were treated, where noted, with 100 μg/ml cycloheximide (CHX), or 10 μg/ml MG-132 (Sigma). Plasmid transfection was done with Fugene 6 (Roche), and siRNA transfections used Oligofectamine (Invitrogen). pRSV-CBP Myc (Sigma) and pCDNA3-p53-UbF, which has the Ub cDNA cloned (14), and (1–200) were generated by cloning the indicated PCR-amplified silent mutation of the shRNA target sequence (AACTCCAATAGC mutated to AACTCCAATAGA) within pRSV-CBP-Myc. pGEX-2tk (1–616) and (1–200) were generated by cloning the indicated PCR-amplified fragments into pGEX-2tk. pCDNA3-p53-UbF, which has the Ub cDNA cloned in-frame to the 3’ end of the p53 CDS, was the generous gift of Christine Blattner, and MT107 His-Ub expression vector has been described (35).

Western Blotting, Immunofluorescence, and Immunoprecipitation. For western blot analyses, cells were lysed in cold NETN240 buffer (20 mM HEPES, pH 7.4, 2 mM EDTA, 5 mM NEM and protease inhibitors) using A-22, N-15, or D-7 antibodies followed by protein A Sepharose. The IPs were washed in high salt buffer three times followed by two washes in Ub buffer (25 mM HEPES, pH 7.4, 10 mM NaCl, 3 mM MgCl2, 0.05% Triton X-100, 0.5 mM DTT, 3 mM Mg-ATP). The washed and equilibrated IPs were then incubated with 100 ng E1 (rabbit; Boston Biochem), and 5 μg Ub (human recombinant; Boston Biochem), and 25 ng E2 (UbHsa, human recombinant; Boston Biochem) at 37 °C for 30 min. FLAG-CBP (1, 10, or 100 μg) immunoprecipitated from baculovirus-infected Sf9 insect cells with anti-FLAG resin/FLAG peptide elution (35) or CBP IP from U2OS cytosolic fraction (see In Vitro E3 Assay supporting the Kimmel Scholar Award, and R01CA107532 from the National Cancer Institute (NCI)).

In Vitro E3 Assays. To determine CBP/p300 or MDM2 E3 ligase activity, CBP, p300, or MDM2 were immunoprecipitated from 0.7 mg cytoplasm or 0.2 mg nuclear extracts diluted with high salt buffer (100 mM HEPES (pH 7.5), 150 mM NaCl, 150 mM KCl, 1 mM MgCl2, 0.5% Triton X-100, supplemented with fresh 5 mM NEM and complete EDTA-free tablets (Roche). Immunoprecipitations from whole cell lysates were performed in the lysis buffer overnight, followed by capture on protein A Sepharose. Ubiquitin conjugates in lysis buffers in western blot signals were quantified after visualization of primary antibody by HRP-conjugated secondary antibody and enhanced chemiluminescence or by fluorescently labeled secondary antibody and detection by Odyssey blot scanner (LiCor), using ImageJ National Institutes of Health (NIH) software.

To determine E4 ligase activity of CBP, a two-step ubiquitination reaction was performed. First, insect-cell-derived p53 was ubiquitinated by MDM2 as described above. To remove MDM2 and nontagged ubiquitin, p53 was then immunoprecipitated and washed three times with RIPA buffer, two times with PBS, and two times with Ub buffer. Immobilized p53 was then mixed with second ubiquitination reaction components (100 ng E1, 25 μg UbHsa, 5 μg FLAG-tagged Ub (Boston Biochem)) along with afffinity-purified Flag-CBP (1–200) or (1–616) polypeptides obtained from transfected U2OS cells and incubated at 37 °C for 60 min. To remove CBP auto-ubiquitination products, beads were washed three times with RIPA buffer. p53 was eluted by boiling with NuPAGE LDS sample buffer, separated by SDS-PAGE, and visualized by immunoblotting with anti-Flag antibodies.

Additional Materials and Methods are available in SI Materials and Methods.

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References

6. An shRNA-resistant CBP allele (pRSV-CBP*myc) was generated by cloning the indicated PCR-amplified silent mutation of the shRNA target sequence (AACTCCAATAGC mutated to AACTCCAATAGA) within pRSV-CBP-Myc.