Human cells enter mitosis with damaged DNA after treatment with pharmacological concentrations of genotoxic agents

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In the present paper, we report that mitosis is a key step in the cellular response to genotoxic agents in human cells. Cells with damaged DNA recruit γ-H2AX (phosphorylated histone H2AX), phosphorylate Chk1 (checkpoint kinase 1) and arrest in the G2-phase of the cell cycle. Strikingly, nearly all cells escape the DNA damage checkpoint and become rounded, by a mechanism that correlates with Chk1 dephosphorylation. The rounded cells are alive and in mitosis as measured by low phospho-Tyr15 Cdk1 (cyclin-dependent kinase 1), high Cdk activity, active Plk1 (Polo-like kinase 1) and high phospho-histone H3 signals. This phenomenon is independent of the type of DNA damage, but is dependent on pharmacologically relevant doses of genotoxicity.

Entry into mitosis is likely to be caused by checkpoint adaptation, and the HT-29 cell-based model provides a powerful experimental system in which to explore its molecular basis. We propose that mitosis with damaged DNA is a biologically significant event because it may cause genomic rearrangement in cells that survive genotoxic damage.

Key words: camptothecin, checkpoint adaptation, checkpoint kinase 1 (Chk1), cyclin-dependent kinase 1 (Cdk1), mitosis, mitotic catastrophe.

INTRODUCTION

To ensure the fidelity of mitosis, cells use biochemical pathways known as checkpoints, which enable them to detect damaged DNA and prevent entry into mitosis [1]. It is widely believed that failure to repair damaged DNA leads to apoptosis and that this protects cells from transmitting damaged DNA to daughter cells [2]. It has also been observed, however, that cells can enter mitosis after being subjected to radiation [3,4] or diverse genotoxic chemicals [5–7] with the possibility of transmitting damaged DNA to daughter cells. The conditions in which cells can enter mitosis with damaged DNA and the biological significance of doing so are poorly understood [8].

Chk1 (checkpoint kinase 1) and Cdk1 (cyclin-dependent kinase 1) (Cdk1–cyclin B complex) are key enzymes in the DNA damage checkpoint and cell cycle pathways [9]. When cells are challenged with damaged DNA, Chk1 is activated by phosphorylation on Ser177 and Ser345 by the upstream kinase ATR (ataxia telangiectasia mutated- and Rad3-related) [10,11]. In addition to phosphorylating Chk1, the Plk1 (phosphokinase 3-related kinase) family members phosphorylate histone H2AX to convert it into γ-H2AX (phosphorylated histone H2AX) [12]. This histone can be used as a marker to identify damaged DNA in cells [13]. Chk1 prevents the activation of Cdk1 by promoting the degradation and sequestration of the CDC25 phosphatases [14–16], which are required to dephosphorylate Tyr15 of the catalytic subunit of Cdk1. If Cdk1 remains phosphorylated on Tyr15, the enzyme stays inactive despite high cyclin B1 levels and the cell is blocked in the G2-phase of the cell cycle. Active Cdk1 drives cells into mitosis, which is characterized by major structural and biochemical changes, including cell rounding [17], chromosome condensation and phosphorylation of core histones. By measuring cyclin B1 levels, Cdk1 activity and phosphorylation of histone H3 on Ser10, cells can be followed as they exit G2-phase and enter mitosis [18].

We examined how cells treated with genotoxic compounds exit from a cell cycle arrest and enter mitosis despite containing damaged DNA. We used CPT (camptothecin), a topoisomerase I inhibitor, as a genotoxic agent. The pharmacokinetics of CPT and its clinical derivatives (topotecan and irinotecan) have been investigated extensively in human subjects, providing valuable information to model the genotoxic response in cancer cells [19]. In the present paper, we report that when cells were treated with a cytotoxic and pharmacologically relevant amount of CPT, they became rounded and entered mitosis even though their DNA was still damaged. This morphological change was a convenient measure of mitosis, which we exploited to examine how Chk1 participates in restarting the cell cycle. Our findings suggest that mitosis is a component of the cellular response to cytotoxic anticancer agents. Furthermore, HT-29 cells have features that make them a valuable model to investigate the molecular basis of checkpoint adaptation in human cells.

MATERIALS AND METHODS

Cell culture

The human cell line HT-29 was obtained from the A.T.C.C. (Manassas, VA, U.S.A.). HT-29 cells were maintained in RPMI 1640 medium (Invitrogen) supplemented with 10% (v/v) FBS.
Mechanical shake-off

HT-29 cells were plated at 40 0000 cells/25 cm² flask and cultivated for 48 h before treatment with compounds. At desired times after treatment, culture medium was replaced with a small volume of RPMI 1640 medium (40 μl/cm²), and flasks were tapped on all sides with medium force until rounded cells were released.

Flow cytometry

At desired times after treatment, total cultures or cells in interphase were collected by trypsinization. Ranged cells were collected by mechanical shake-off. Cells were washed in PBS and fixed in 90% ethanol (−20 °C) for at least 24 h. Fixed cell suspensions were blocked for 1 h with labelling buffer (PBS, 5% serum, 1% BSA and 0.1% sodium azide) before 1 h of incubation with anti-(Ser 10-phosphorylated histone H3) antibody (06-570, Millipore; 1:100 dilution), and 30 min of incubation with FITC-conjugated secondary antibody (sc-2012, Santa Cruz Biotechnology; 1:100 dilution) in labelling buffer, separated by wash/centrifuge steps in wash buffer (PBS, 1% BSA and 0.1% sodium azide). For analysis, samples were incubated for 20 min in wash buffer with 0.02 mg/ml propidium iodide (Invitrogen) and 0.2 mg/ml RNase A (Sigma), and analysed by a FACSCanto™ II flow cytometer (BD Biosciences) using BD FACSDiva™ software. Gating was set using control samples without primary antibody. Experiments were carried out at least twice.

Light microscopy

Images were taken with an Infinity 1.5 camera powered by Infinity Capture (Lumenera Corporation) software. Live cells were detected by dye exclusion with Amresco® Trypan Blue 0.4% solution (VWR). Me-CR8 and CR8 were used at 25 μM concentrations with or without 25 nM CPT. Experiments with Me-CR8 or CR8 or MG132 were carried out three times. MG132 was used at 0.3 μM.

Immunofluorescence microscopy

Cells were plated on glass coverslips for 48 h before treatment. Cells collected by mitotic shake-off were attached to glass coverslips coated with poly-L-lysine (Invitrogen). At desired times, cells were fixed in 3% (w/v) formaldehyde for 20 min at room temperature (20 °C) and permeabilized for 5 min in 0.2% Triton X-100. Cells were incubated with anti-γH2AX (05-636, Millipore), anti-(Ser 10-phosphorylated histone H3) (as above) or anti-cyclin B1 (sc-752, Santa Cruz Biotechnology) for 2 h at room temperature [7]. Texas Red-conjugated anti-rabbit secondary antibodies (Jackson ImmunoResearch/Beckman Coulter) for histone H3 and cyclin B1, or Alexa Fluor® 488-conjugated anti-mouse secondary antibodies (Molecular Probes/Invitrogen) for γH2AX were added for 2 h. Nuclei were stained with 300 nM DAPI (4',6-diamidino-2-phenylindole) in PBS for 15 min before mounting. Cells were observed using a Zeiss microscope operated by Axiovision 3.1 software. Images were collected with a Zeiss MR camera within the linear dynamic range. Images were prepared for presentation using identical parameters with Adobe Photoshop CS3 10.0 software. Experiments were carried out at least twice. Confocal images were prepared with a Nikon C1+ confocal system equipped with an inverted Eclipse TE 2000U microscope.

Cytotoxicity assays

Cytotoxicity was measured using the MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl-2H-tetrazolium bromide] assay [20]. Results were expressed as IC50 values, the compound concentration that reduced by 50% the absorbance at 590 nm, compared with DMSO-treated cells. All measurements were carried out in triplicate.

Extract preparation

Cells were resuspended in extraction buffer [50 mM Hepes, 50 mM NaF, 10 mM EGTA, 50 mM 2-glycerophosphate, 1 mM ATP, 1 mM DTT (dithiothreitol), 1% Triton X-100, 10 μg/ml RNase A, 0.4 unit/ml DNase I, with Roche protease inhibitor cocktail] at a concentration of 20 000 cells/μl, on ice for 30 min. After five passages through a 26-gauge needle at 4 °C, the suspension was centrifuged at 12 000 g for 10 min at 4 °C. Extracts were used either for electrophoresis after being boiled for 3 min in the presence of 2× SDS sample buffer or measurement of Cdk/cyclin activity. Cdk activity was measured by incubating extracts with a GST (glutathione transferase)–Cdk1 substrate or GST alone followed by Western blotting with an anti-(phospho-Thr 320 PPK1α) (protein phosphatase 1 catalytic subunit) antibody [21]. Details of the Cdk assay are available from R.M.G. on request.

Electrophoresis and Western blotting

Samples (10 μg of total protein per lane) were run on polyacrylamide gels. Protein loading was confirmed by Coomassie Blue staining of gels run in parallel or by anti-actin Western blotting. Molecular-mass markers (Precision Plus) were from Bio-Rad Laboratories. Proteins were transferred on to nitrocellulose membranes with a semi-dry electrobetter system (Bio-Rad Laboratories) for 45 min at 25 V. Subsequently, the membrane was blocked with either 5% (w/v) low-fat milk powder in TBST (Tris-buffered saline with Tween 20: 50 mM Tris/HCl, 150 mM NaCl and 0.1% Tween 20, pH 7.6) or 5% (w/v) BSA in TBST, and incubated overnight with the indicated primary antibody as follows: anti-Chk1 (sc-8408, Santa Cruz Biotechnology; 1:200 dilution); anti-(phospho-Ser 45 Chk1) (2341S, Cell Signaling Technology; 1:1000 dilution); anti-(cyclin B1) (GNS1, Santa Cruz Biotechnology; 1:200 dilution), anti-Cdk1/Cdc2 (21236-2, Signalway Antibodies; 1:500 dilution), anti-(phospho-Tyr 15 Cdk/Cdc2) (11244-2, Signalway Antibodies; 1:500 dilution), anti-(phospho-Thr 232 PPK1α) (protein phosphatase 1α) (2581S, Cell Signaling Technology; 1:1000 dilution); anti-Plk1 (Polo-like kinase 1) (33-1700, Zymed Laboratories; 1:300 dilution); anti-(phospho-Thr 210 Plk1) (558400, BD Biosciences; 1:300 dilution).

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Figure 1  HT-29 cells treated with 25 nM CPT are alive at 48 h and arrested in the G2/M-phase of the cell cycle

(a) Human colon carcinoma cells (HT-29) were treated with graded concentrations of CPT for either 48 h (●) or 96 h (■) and viability was measured by the MTT assay. Results are mean ± S.E.M. percentages of live cells for triplicate experiments. (b) Cells were cultured with 25 nM CPT and prepared for analysis by flow cytometry at 24 and 48 h after treatments. Control cells (not treated) were also analysed. DNA content was determined by propidium iodide staining.

RESULTS

We incubated HT-29 cells with increasing concentrations of CPT and measured their viability at 48 or 96 h by the MTT assay. The IC50 of CPT was 175 nM at 48 h; however, at 96 h the IC50 was 5 nM (Figure 1a). On the basis of these data, we chose 25 nM CPT for subsequent experiments because this concentration is a pharmacologically relevant cytotoxic concentration in human patients [22]. Importantly, at this concentration, more than 80% of the cells were alive at 48 h even though they were destined to die by 96 h. We then treated HT-29 cells with 25 nM CPT and analysed them for cell cycle phase by flow cytometry. Cells accumulated in S-phase and G2/M-phase by 24 h after treatment, and were in the G2/M-phase by 48 h, consistent with the activation of the DNA damage checkpoint (Figure 1b). These data highlighted that HT-29 cells were alive and arrested in the cell cycle at 48 h after treatment with a pharmacologically relevant and cytotoxic concentration of CPT.

We then investigated the fate of the cells after cell cycle arrest. At 24 h after CPT treatment (Figure 2a), nearly all cells were flat and strongly adherent. At 48 h, when most cells were arrested in G2-phase, relatively large rounded cells appeared, which were weakly adherent. These rounded cells could be collected by mechanical shake-off, leaving behind the flattened cells (Figure 2a). The rounded cells were alive as confirmed by vital dye exclusion (results not shown). We re-cultivated the flattened cells that remained attached after mechanical shake-off and found that new rounded cells appeared by 2 h, as a result of a continuous process. For subsequent experiments, we worked with three types of treated cell populations, which we defined as follows: TDCs (total and DNA-damaged cells), a total cell population with damaged DNA composed of interphase and rounded cells; IDCs (interphasic and DNA-damaged cells), flat interphase cells with damaged DNA that were strongly adherent and could not be collected by mechanical shake-off; and MDCs (mechanicallyshake-off cells), cells that were collected by mechanical shake-off and were rounded cells.
examined them for cyclin B1 and Cdk1 by Western blotting. Cells without CPT treatment had lower levels of cyclin B1 than those treated with CPT (Figure 3b). Furthermore, the MDCs expressed relatively high levels of cyclin B1 that were comparable with cells arrested in mitosis by nocodazole treatment. We also examined cells for Cdk1 protein levels and Cdk1 phosphorylation on Tyr15. The levels of Cdk1 phosphorylated on Tyr15 were much lower in extracts prepared from MDCs than in non-treated extracts or IDCs. The Cdk1 signal of MDCs was similar to that of nocodazole-treated cells, suggesting that these cells contained the active form of Cdk1. We then compared the amounts of phospho-Thr210 PP1 in each extract. PP1 is phosphorylated on Thr210 by Cdk1 [21] and is used as an indicator of Cdk1 activity [26]. MDCs contained phospho-Thr210 PP1, whereas IDCs did not, confirming our observation that rounded cells have active Cdk1. We examined the levels of Plk1 [27] and activated Plk1 (phospho-Thr210) [28] in these samples. Plk1 levels were relatively high in IDC and MDC samples and low in non-treated samples, consistent with its expression in the G2/M-phase [29]. Phospho-Thr210 Plk1 levels were highest in MDC samples, yet could still be detected in IDC samples.

To determine whether cells enter mitosis after CPT treatment, we examined them for phospho-Ser10 histone H3 and DNA content by flow cytometry. A non-treated cell culture displayed a typical DNA distribution with approximately 2% of the cells positive for phospho-Ser10 histone H3, compared with 64% of the cells treated with nocodazole (Figure 4a). At 48 h after CPT treatment, 24% of the TDC population was positive for phospho-Ser10 histone H3. MDCs, which were collected at 48 h, were 73% positive for phospho-Ser10 histone H3, whereas the IDCs had only 1.9% positive cells remaining. We then measured Cdk enzyme activity in extracts prepared from CPT-treated cells. MDCs had much higher Cdk activity than IDCs or non-treated cells as shown by phosphorylation of an artificial Cdk substrate, but similar to that of nocodazole extracts (Figure 4b). The negative control substrate, which did not have a Cdk1 phosphorylation site engineered into it, was not phosphorylated. These results indicate that CPT-treated cells are capable of restarting the cell cycle and entering mitosis.

We then tested whether entry into mitosis after CPT treatment would occur in a cell line other than HT-29. We chose M059K cells, which, in contrast with HT-29, are of brain (glioma) origin, fibroblastic and have a long (48 h) cell cycle. Like HT-29, M059K cells are sensitive to CPT, which is a therapeutic option for glioma. M059K cells were alive after treatment with clinically relevant cytotoxic concentrations of CPT at 48 h but were destined to die by 96 h (Supplementary Figure S1a at http://www.BiochemJ.org/bj/446/bj4460373add.htm). We observed cell rounding in CPT-treated populations and the appearance of new rounded cells 2 h after mechanical shake-off of a 48 h IDC population (Supplementary Figure S1b). Because M059Ks have a longer cell cycle than HT-29 cells, fewer cells were in mitosis after CPT treatment at any time tested, including those arrested with nocodazole (7%). Importantly, starting at 48 h after CPT treatment, phospho-Ser10 histone H3-positive cells appeared at frequencies that were higher than for non-treated cells (1.5%) or cells under a DNA damage checkpoint at 24 h (0.8%). These data suggest that the phenomenon of entry into mitosis with damaged DNA is common to many cell types.

If cells enter mitosis after CPT treatment, we reasoned that a Cdk inhibitor might prevent them from doing so. Cells were treated with DAPI (blue) and with anti-(cyclin B1) antibodies (red). Cells were analysed by immunofluorescence microscopy. Scale bar, 50 μm. (b) Extracts were prepared from cells either treated (right) or not treated (left) with CPT. CPT-treated cells were analysed further as total cultures (TDC) or cultures fractionated by mechanical shake-off into adherent interphase cells (IDC) or rounded cells (MDC). Extracts were also prepared from cells treated with nocodazole (NocO). Samples were processed by Western blotting with antibodies against cyclin B1, Cdk1, phospho-Tyr15 Cdk1 (P-Y15 Cdk1), phospho-Thr320 PP1a (P-T320 PP1), Plk1, phospho-Thr210 Plk1 (P-T210 Plk1) or actin. Molecular masses are indicated in kDa.

Figure 3 MDCs express cyclin B1 and the active form of Cdk1

(a) Cells were grown on coverslips and either not treated (upper row) or treated with CPT for 48 h (lower row), stained with DAPI (blue) and with anti-(cyclin B1) antibodies (red). Cells were analysed by immunofluorescence microscopy. Scale bar, 50 μm. (b) Extracts were prepared from cells either not treated (NT) or treated with CPT. CPT-treated cells were analysed further as total cultures (TDC) or cultures fractionated by mechanical shake-off into adherent interphase cells (IDC) or rounded cells (MDC). Extracts were also prepared from cells treated with nocodazole (NocO). Samples were processed by Western blotting with antibodies against cyclin B1, Cdk1, phospho-Tyr15 Cdk1 (P-Y15 Cdk1), phospho-Thr320 PP1a (P-T320 PP1), Plk1, phospho-Thr210 Plk1 (P-T210 Plk1) or actin. Molecular masses are indicated in kDa.

To determine whether the MDC morphology was related to genotoxic responses other than CPT, we treated cells with 3 μM etoposide. Like the conditions used for CPT, 3 μM etoposide is a pharmacologically relevant cytotoxic concentration [23]. Etoposide induced the appearance of rounded cells (Figure 2b), indicating that the MDC morphology seen with CPT is probably common to genotoxic agents. The frequency of rounded cells was approximately 2–3% of the total culture at 48 h and 10–12% by 72 h, whereas rounded cells made up approximately 1% of an untreated culture (Figure 2c). This number of MDCs, and the likelihood that cells were continuously entering this population, suggested that this change in cell shape might reflect an important event after cell cycle arrest.

In non-treated cultures, rounded cells that can be collected by mechanical shake-off are typically in mitosis [24]. Therefore we tested whether MDCs induced by CPT treatment were in mitosis. In non-treated cells, immunofluorescence microscopy revealed that only a subpopulation of cells were cyclin B1-positive, which was consistent with cell-cycle-dependent expression of cyclin B1 [25] (Figure 3a). However, in cultures treated for 48 h with 25 nM CPT, nearly all cells were positive for cyclin B1, which was consistent with cells arrested in the G2-phase of the cell cycle. We separated cells into IDC and MDC populations and...
Figure 4 Rounded cells (MDC) are positive for phospho-Ser^10 histone H3 and for Cdk activity

(a) Cells were either not treated (NT) or treated with nocodazole (Noco) or 25 mM CPT for 48 h (TDC). A second TDC population was fractionated into IDC and MDC populations by mechanical shake-off. All samples were analysed by flow cytometry for DNA content and for phospho-Ser^10 histone H3 signals. The percentage of cells positive for phospho-Ser^10 histone H3 (P-H3) is listed in the upper right-hand quadrant. (b) Extracts were prepared from IDCs, MDCs, cells not treated (NT) or cells treated with nocodazole (Noco). Extracts were incubated with a GST–Thr^320 Cdk1 substrate (lanes 1, 2 and 5–7) or a GST-control substrate (lanes 3, 4 and 8–10). Samples were analysed by Western blotting with an anti-Cdk1 phospho-substrate antibody (upper panel, Anti P-T320). Equal amounts of substrate were used in each sample (lower panel, Anti GST). IDC, MDC, NT and Noco extracts were tested at concentrations representing 500 cells. Blank indicates buffer was used in place of extract. Molecular masses are indicated in kDa.

Figure 5 Cell cycle inhibitors block the entry into mitosis of cells treated with CPT

(a) Cells were either not treated (upper row) or treated with CPT (lower row). At 24 h after treatment, cells were either treated with methyl-CR8 (Me-CR8) or with CR8 or no further treatment (NT) and cultivated for a further 24 h. Scale bar, 200 μm. (b) The number of MDCs collected by mechanical shake-off were counted at 48 h under each treatment condition and presented as percentages relative to non-treated cells (NT). (c) Cells were either not treated or treated with 100 mM BI2536 for 24 h, 25 mM CPT for 48 h, or co-treated with 25 mM CPT for 48 h with 100 mM BI2536 for the last 24 h. Samples were analysed by flow cytometry for DNA content and for phospho-Ser^10 histone H3 signals. The percentage of cells positive for phospho-Ser^10 histone H3 (P-H3) is listed in the upper right-hand quadrant.

(Tables 5a and 5b). In contrast, methyl-CR8 had little effect upon the number of MDCs. These results confirmed that CPT-treated cells are able to enter mitosis and that this event can be blocked by addition of Cdk1 inhibitors. We then incubated CPT-treated cells with the Plk1 inhibitor BI2536 to determine whether Plk1 activity was required for entry into mitosis with damaged DNA (Figure 5C). First, we demonstrated that 100 mM BI2536 was sufficient to induce a mitotic arrest in HT-29 cells. We then treated cells for 24 h with CPT and then for a further 24 h with CPT and BI2536. Unlike the Cdk1 inhibitor, the Plk1 inhibitor consistently reduced the number of phospho-Ser^10 histone H3 cells by approximately 40%.

CPT induces a DNA damage checkpoint [30], which arrests cells in the cell cycle, yet we observed that treated cells could enter mitosis. Therefore we wanted to know whether the mitotic cells no longer had damaged DNA, or whether the DNA damage checkpoint was extinguished. We examined cells by immunofluorescence microscopy for γH2AX staining, which is recruited to sites of damaged DNA (Figure 6). Cells were treated with CPT and at various times fixed and stained to detect damaged DNA and mitotic chromosomes. At 24 h after treatment, 97–100% of treated cells displayed γH2AX foci, indicating that this population had damaged DNA (Figure 6a), whereas no signals were observed in cells without CPT treatment. By 48 h, IDCs were still strongly positive; however, cells with condensed chromosomes were also strongly positive, suggesting that cells were entering mitosis even though their DNA was still damaged. It had been reported previously that cells in mitosis might be positive for γH2AX [31], therefore we compared γH2AX signal intensity in mitotic cells after arrest by nocodazole with those treated with CPT. We found that γH2AX staining was much more intense in MDCs, which have damaged DNA but not mitotic DNA. Therefore, we conclude that CPT-treated cells can enter mitosis even though their DNA was still damaged. However, CPT-treated cells that entered mitosis were strongly positive for γH2AX, indicating that their DNA was still damaged.
DNA, so that they could be readily distinguished from mitotic cells without induced DNA damage (Supplementary Figure S2 at http://www.BiochemJ.org/bj/446/bj4460373add.htm). We then collected MDCs by mechanical shake-off, mounted them on coverslips and observed them by confocal microscopy for DNA organization, phospho-Ser10 histone H3 and γ-H2AX antibodies (right-hand panels). Cells were observed after 24 and 48 h of treatment by immunofluorescence microscopy. Scale bar, 20 µm. (b) MDCs were collected, attached to coverslips and stained with DAPI (left) and antibodies against γ-H2AX (centre) or phospho-Ser10 histone H3 (right, P-H3) and observed by confocal microscopy. Scale bar, 20 µm.

Figure 6 MDCs are in mitosis with damaged DNA
(a) Cells were either not treated or treated with CPT and stained with DAPI (left-hand panels) and γ-H2AX antibodies (right-hand panels). Cells were observed after 24 and 48 h of treatment by immunofluorescence microscopy. Scale bar, 10 µm. (b) MDCs were collected, attached to coverslips and stained with DAPI (left) and antibodies against γ-H2AX (centre) or phospho-Ser10 histone H3 (right, P-H3) and observed by confocal microscopy. Scale bar, 20 µm.

DNA, so that they could be readily distinguished from mitotic cells without induced DNA damage (Supplementary Figure S2 at http://www.BiochemJ.org/bj/446/bj4460373add.htm). We then collected MDCs by mechanical shake-off, mounted them on coverslips and observed them by confocal microscopy for DNA organization, phospho-Ser10 histone H3 and γ-H2AX staining (Figure 6b). The majority of MDCs had poorly formed chromosomes, typical of cells in mitotic catastrophe [32]. They were also phospho-Ser10 histone H3-positive, but, strikingly, they were positive for γ-H2AX signals. This suggested that these mitotic cells had damaged DNA [33], in addition to atypical chromosome arrangements. We concluded that MDCs had entered mitosis with damaged DNA.

To determine whether mitosis was a frequent event in a population of CPT-treated cells, we counted the number of MDCs at regular intervals after CPT treatment until 96 h, when most, but not all, cells were dead. The time of the greatest number of mitotic cells was 62 h after treatment, although mitotic cells could be detected as early as 40 h. Using video microscopy and cell counting, we found that more than 90% of a cell population entered mitosis before they die (results not shown).

Cells are believed to be able to escape the DNA damage checkpoint by inactivation of Chk1 by either proteolysis [34] or by dephosphorylation [3]. We tested both of these hypotheses using our model. We treated cells with 25 nM CPT and asked whether cells would enter mitosis in the presence of MG132, a proteasome inhibitor (Figure 7a). Cells were cultivated with CPT for 48 h and MDCs were removed by mechanical shake-off. The remaining IDCs were then incubated with MG132 for another 6 h and the number of new MDCs were counted and compared with cells incubated for 6 h without MG132. We detected the same number of MDCs in the presence or absence of the proteasome inhibitor (Figure 7b). We verified that we used an adequate amount of MG132 because we could arrest cells in mitosis by treatment with 300 nM MG132 (see also [35]).

We then compared Chk1 protein levels in extracts prepared from cells treated with increasing half-log concentrations of CPT ranging from 0.03 to 3 µM (Figure 7c). We found that Chk1 protein levels were relatively stable up to 0.3 µM CPT, but were much lower when cells were treated with CPT concentrations greater than 0.3 µM. The activated form of Chk1, phospho-Ser145, was present in treated cells, except when Chk1 was low. These data suggested that cells might enter mitosis with damaged DNA by mechanisms other than, but in addition to, Chk1 degradation, depending on the concentration of the genotoxic agent.

We then examined the level of Chk1 in MDCs and compared it with IDCs or IDCs. Chk1 levels were similar in extracts prepared from cells under either condition (Figure 7d). We observed a slight decrease in electrophoretic mobility of Chk1 in MDCs and cells treated with nocodazole, as had been described previously [36]. Strikingly, we found that Chk1 had very low levels of phosphorylation on Ser145 in MDCs. We also observed a loss of phosphorylation on Ser317, a second phosphorylation site on active Chk1 (results not shown). We explored Chk1 dephosphorylation further by examining an IDC population that was re-incubated for 2 h so that new MDCs could be collected (Figure 7e). Under these conditions, we observed again that Chk1 was stable after entry into mitosis and that the dephosphorylation step at Ser145 was closely correlated with entry into mitosis.

Discussion
The events that occur after cell cycle arrest in cells with damaged DNA are poorly understood. To study them, we set up a cell-based assay in which we treated cells with genotoxic agents and observed them at various times. We found that cells became rounded, starting at approximately 40 h after treatment with CPT. The rounded cells are in mitosis, express high levels of cyclin B1, have active Plk1, do not express Cdk1 phosphorylated on Tyr15, have Cdk activity and are positive for phospho-Ser10 histone H3. The relationship between cell rounding and Cdk activity was demonstrated by applying the Cdk inhibitor CR8, which prevented cell rounding. The inactive form of this inhibitor class had no effect upon the transition from IDCs to rounded mitotic cells (MDCs). This cell-based model enables us to study pathways that cells use to exit a DNA damage checkpoint, enter mitosis with damaged DNA and, in some cases, survive.

We set up the assay under conditions that may be relevant for studying the cellular response to cancer drugs in humans. We chose a concentration of CPT that was cytotoxic, as nearly all cells will die by 96 h. Furthermore, this concentration was within the range of those obtained in the plasma of patients treated with CPT or its derivatives [22]. We confirmed that HT-29 cells responded to CPT in the manner described previously: nearly all cells had multiple γ-H2AX foci after treatment [7,37], induction of Chk1 by phosphorylation on Ser145 [11] and cell cycle arrest in G2-phase by 48 h.

The rounded cell morphology provided a simple means to examine how cells with damaged DNA enter mitosis. Because the MDCs were rounded and weakly adherent, they could
Figure 7 Chk1 is dephosphorylated and not degraded in cells in mitosis with damaged DNA

Cells were either not treated or treated with MG132 for 6 h or with CPT for 48 h. At 48 h, MDCs were removed by mechanical shake-off from a total culture and the remaining IDCs were either re-incubated with CPT for 6 h (IDC 54 h) or incubated with CPT and MG132 for 6 h (IDC + MG132 54 h). Scale bar, 100 μm. (b) The number of MDCs collected by mechanical shake-off was counted at 48 h under each treatment condition and presented as percentages relative to non-treated cells (NT). (c) Cells were either not treated (NT) or treated with increasing concentrations of CPT (half-log; 0.03–3 μM). Extracts were prepared and analysed by Western blotting with antibodies against either Chk1 or phospho-Ser345 Chk1 or by Coomassie Blue staining. Molecular masses are indicated in kDa. (d) Cells were treated with CPT for 48 h (TDC) and separated into IDC and MDC populations. Mitotic cells without damaged DNA were prepared from nocodazole-treated culture (Noco). Samples were processed by Western blotting with antibodies against either Chk1 or phospho-Ser345 Chk1. Molecular masses are indicated in kDa. (e) Cells were treated with CPT for 48 h (TDC) and separated into IDC and MDC populations (MDC1). The IDC cells were re-cultivated for 2 h and new mitotic cells were collected (MDC2). Samples were processed by Western blotting with antibodies against either Chk1 or phospho-Ser345 Chk1. Molecular masses are indicated in kDa. p, phospho-

be readily distinguished and isolated from IDCs. Furthermore, by performing sequential mechanical shake-offs, we confirmed that the transition from IDCs to MDCs is a continuous event. At each time tested, starting at approximately 40 h after CPT treatment, cells would enter mitosis. The IDC population appeared to be uniform in their shape and DNA content, yet MDCs appeared asynchronously, hinting that they are following a timing mechanism that we have not yet identified. The lack of synchrony made it difficult to count precisely the proportion of cells that enter mitosis over a period of 48 h. To address this, preliminary observations using video microscopy have revealed that more than 90% of the TDCs enter mitosis before dying, indicating that this is a major cellular event.

We used this model to test the role of Chk1 in exiting the DNA damage-induced cell cycle arrest. Previous studies observed that Chk1 is inactivated by either degradation [34] or dephosphorylation [3]. We found that Chk1 was present in similar amounts in IDCs as in MDCs, although it was no longer phosphorylated on Ser345. A reduced electrophoretic mobility of Chk1 in mitotic cells has been reported to be caused by phosphorylation on Ser345 and Ser301 by Cdk1 [38]. As the MDCs are in mitosis, it is likely that the reduced Chk1 mobility that we observed was due to Cdk1 phosphorylation. Entry into mitosis and Chk1 dephosphorylation on Ser345 were closely linked because we detected Chk1 dephosphorylation in cells that had only entered mitosis after a very short time. We did not detect Chk1 degradation in cells that were treated with pharmacological, yet cytotoxic, concentrations amounts of CPT. Furthermore, treated cells could still enter mitosis even with co-treatment with the proteasome inhibitor, MG132. Our data suggest that when cells are challenged with cytotoxic amounts of damaged DNA, they can use one of two mechanisms to inactivate Chk1. We speculate that the selection of the Chk1 inactivation mechanism (dephosphorylation or degradation) will be linked to the amount of damaged DNA.

The cellular process that we have observed is similar to what has been described in yeast as checkpoint adaptation. Checkpoint adaptation involves three key steps: arrest in the cell cycle due to damaged DNA, overcoming this arrest and resuming the cell cycle even though cells still have damaged DNA [39]. We have detected cell cycle arrest after treatment with CPT and entry into mitosis with poorly formed chromosomes that were γH2AX-positive in human cells. Checkpoint adaptation has been described in human cells that were irradiated [3]. Our data demonstrate that CPT and possibly most genotoxic agents induce checkpoint adaptation.

Our observations suggest that checkpoint adaptation is one of the steps that lie between cell cycle arrest and cell death (Figure 8). Interfering with these steps is likely to modify the outcome of the genotoxic response [40,41]. Cells that escape the DNA damage checkpoint with poorly organized chromosomes might activate a mitotic checkpoint [7] and cause the accumulation of MDCs [5]. Furthermore, we confirmed that this phenomenon occurs in osteosarcoma U2OS cells as reported previously [3], and we have detected it in human M059K glioma cells, which indicates that checkpoint adaptation may be a common mechanism in cell death.

Although cells treated with cytotoxic concentrations of DNA-damaging agents are destined to die, it is important to understand all of the steps that lead to cell death (Figure 8). Entry into mitosis before cell death and mixing of mitotic cells with dying cells may account for previous reports of Cdk activity occurring during apoptosis [42,43]. Knowing that mitosis precedes cell death may help to design strategies to improve the outcome of the majority of current cancer treatments, which use genotoxicity as their mechanism of action. For example, Chk1 inhibitors are being evaluated for use in association with genotoxic agents to improve...
Cells respond to cytotoxic pharmacological amounts of genotoxic agents through a series of key biochemical events that include: increase in γ-H2AX levels and increase in phospho-Ser345 Chk1 levels before adaptation. Cells that have undergone checkpoint adaptation still have γ-H2AX, but a greatly reduced level of phospho-Ser345 Chk1, high Cdk1 activity and phospho-Ser10 histone H3. Although the majority of cells will die, a small number of cells survive, giving rise to cells that have major changes in their genome.

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SUPPLEMENTARY ONLINE DATA

Human cells enter mitosis with damaged DNA after treatment with pharmacological concentrations of genotoxic agents

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Figure S1 M059K cells treated with CPT are alive at 48 h and enter mitosis

(a) Human fibroblastic glioma cells (M059K) were treated with graded concentrations of CPT for either 48 h (●) or 96 h (▲) and viability was measured by the MTT assay. Results are mean ± S.E.M. percentages of live cells from triplicate experiments. M059K cells were either not treated or treated with 25 nM CPT and observed by phase-contrast light microscopy at 24, 48, 72 and 96 h. Rounded cells were visible in non-treated samples, but rarely at 24 h after treatment and commonly by 48 h after treatment. Rounded cells were collected after mechanical shake-off (MDC) leaving behind flattened interphase cells (IDC). New rounded cells appeared within 2 h in the adherent culture. Scale bar, 100 μm. (b) Cells were either not treated (NT) or treated with nocodazole (Noco) or 25 nM CPT for 24, 48, 72 and 96 h. Samples were analysed by flow cytometry for DNA content and for phospho-Ser10 histone H3 signals. The percentage of cells positive for phospho-Ser10 histone H3 (P-H3) is listed in the upper right-hand quadrant. DNA content was determined by propidium iodide staining.

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Figure S2  MDCs can be distinguished from nocodazole-induced mitotic cells by γH2AX staining

Cells were not treated, treated with nocodazole (Noco) or treated with CPT. Mitotic cells were collected after nocodazole or CPT treatment by mechanical shake-off, plated on to coverslips and stained with DAPI (left-hand panels) and γH2AX antibodies (right-hand panels). Cells were observed by immunofluorescence microscopy under identical detection conditions. Scale bar, 10 μm.