

Mechanistic Distinctions between CHK1 and WEE1 Inhibition Guide the Scheduling of Triple Therapy with Gemcitabine

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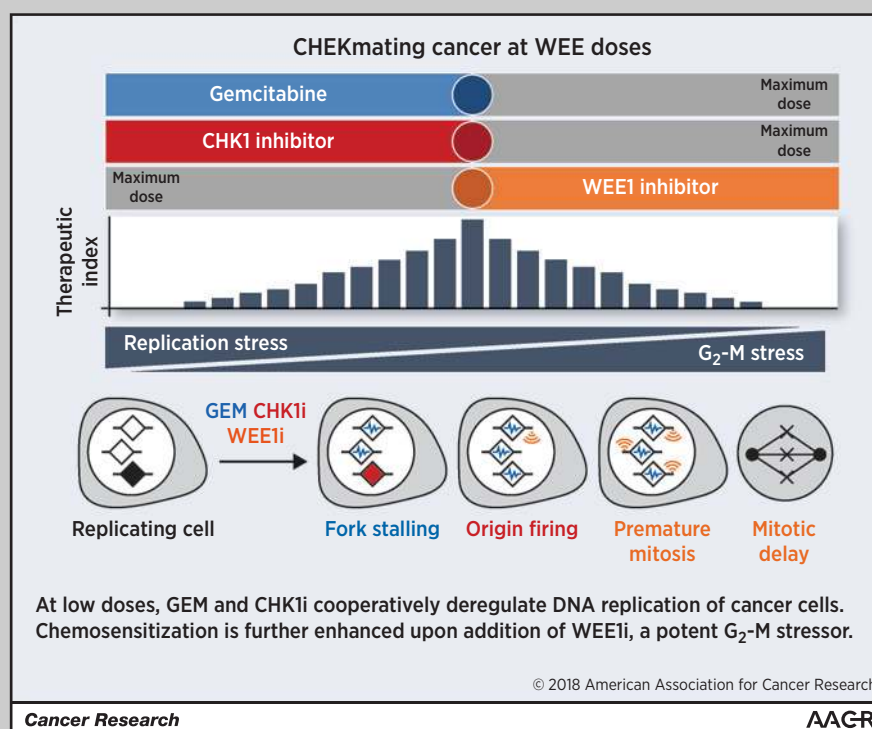


Abstract

Combination of cytotoxic therapy with emerging DNA damage response inhibitors (DDRi) has been limited by tolerability issues. However, the goal of most combination trials has been to administer DDRi with standard-of-care doses of chemotherapy. We hypothesized that mechanism-guided treatment scheduling could reduce the incidence of dose-limiting toxicities and enable tolerable multitherapeutic regimens. Integrative analyses of mathematical modeling and single-cell assays distinguished the synergy kinetics of WEE1 inhibitor (WEE1i) from CHEK1 inhibitor (CHK1i) by potency, spatio-temporal perturbation, and mitotic effects when combined with gemcitabine. These divergent properties collectively supported a triple-agent strategy, whereby a pulse of gemcitabine and CHK1i followed by WEE1i durably suppressed tumor cell growth. In xenografts, CHK1i exaggerated replication stress without mitotic CDK hyperactivation, enriching a geminin-positive subpopulation and intratumoral gemcitabine metabolite. Without overt toxicity, addition of WEE1i to low-dose gemcitabine and CHK1i was most effective in tumor control compared with single and double agents. Overall, our work provides quantitative insights into the mechanisms of DDRi chemosensitization, leading to the rational development of a tolerable multitherapeutic regimen.

Significance: Multiple lines of mechanistic insight regarding DNA damage response inhibitors rationally guide the preclinical development of a tolerable multitherapeutic regimen.

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Introduction

Cancer cells contend with challenging intracellular stress arising from dysfunctional cell-cycle regulations (1). In many tumor subtypes, defects in the p53 gene and in associated signaling pathways impair core processes of the DNA damage response (DDR) program. This scenario renders cancer cells susceptible to exogenous damage, a concept that underpins the use of current chemotherapy. However, because of their nonselective mechanisms of action, many cytotoxic agents operate within a narrow therapeutic window. A strategy to redress this clinical issue is

through the development of targeted agents that exploit cancer-specific cell-cycle functions (2).

The cell cycle is regulated by a network of diverse, partly overlapping and in some cases redundant checkpoints. These fidelity-monitoring checkpoints are invoked upon DNA damage to promote cell-cycle delay and DNA repair. In human cells, the G₂-M checkpoint governs mitotic entry, a transition highly dependent on CDK1. A major negative regulator of CDK1 is WEE1 kinase. Under physiologic condition, WEE1 activity rises during S and G₂ phases in tandem with its protein expression (3). At mitotic onset, WEE1 is inhibited by CDK1 in a double-negative feedback loop to allow for the activation of the latter. Conversely, during genomic crisis, WEE1 phosphorylates CDK1 at its inhibitory residue tyrosine 15, deactivating it and thereby preventing mitotic entry (4). Apart from its involvement in G₂-M transition, more recent studies have begun to clarify the functions of WEE1 during S phase. Similar to CHK1, another CDK-regulating kinase downstream of the ATR and ATM kinases, WEE1 is found to regulate DNA replication initiation and nucleotide consumption (5).

Given its multiple roles in the DDR program, targeting WEE1 in cancer has attracted much attention. Numerous studies have now shown that WEE1 inhibitor (WEE1i) augments the effect of DNA-damaging agents, notably gemcitabine (6, 7). Owing to the functional overlap between WEE1 and CHK1, the current mechanistic paradigm prescribes that WEE1i interacts with gemcitabine similarly as CHEK1 inhibitor (CHK1i) does. The presumption is that both types of inhibition amplify gemcitabine-induced damage via G₂-M abrogation, causing premature mitosis, mitotic catastrophe, and consequently cell death (8, 9). Emerging evidence has also associated WEE1i to aberrant origin firing and nucleotide depletion, culminating in phenotypes of replication stress reminiscent of those observed when ATR/CHK1 signaling is interrupted (10–13). Despite these conceptual advances, several key questions persist. There is considerable ambiguity as to whether WEE1 and CHK1 have distinct modulatory effects on the cell cycle, given that coinhibition of the two kinases in the absence of a cytotoxic agent yields synergistic effects (14, 15). It is not without precedent that kinases acting along the same pathway could have independent molecular cross-talks that lead to different outcomes when their activity is inhibited. For instance, although ATR and CHK1 both coordinate S-phase regulation, suppression of ATR triggers a DNA-PK-mediated pathway that reactivates CHK1, but such compensation is abolished when CHK1 itself is inhibited (16). Moreover, although it is evident that WEE1i and CHK1i perturb both S and G₂-M checkpoints, the extent of these disruptions and how each of them contributes to chemosensitization remain undefined.

The paucity of mechanistic insights into WEE1i and CHK1i chemosensitization has precluded the optimal administration of these inhibitors, even though multiple WEE1i- and CHK1i-based human trials are already underway. Indeed, earlier drug development effort has seen attrition of CHK1/2i following unacceptable toxicities in patients (17, 18). The underlying cause of failure has been attributed to off-target effects than to class specificity, suggesting that tolerability issues to these inhibitors are likely to be related to dose and agent selectivity (19). In the setting of combinations, the challenge of dose-limiting toxicities becomes more prominent. When combined with chemotherapy, WEE1i multiple-dose regimen was more likely to induce toxicities in patients than WEE1i single-dose regimen (20). Likewise, gemci-

tatine plus CHK1i increased the frequency and severity of adverse effects in patients beyond what would be anticipated with gemcitabine alone (21). Previous work by us and others have shown that the combination of gemcitabine and CHK1i could be synergistic at low doses (11, 22). This proposition has now been reinforced by early data demonstrating durable patient response following treatment of CHK1i and gemcitabine at a dose that was several times lower than the standard (23). Nevertheless, it remains debatable how these kinase inhibitors should be deployed as combinations. Based on earlier studies that presuppose G₂-M abrogation as the mechanism of chemosensitization, CHK1i and WEE1i are to be given sequentially after and not concurrently with DNA-damaging agents (24, 25). Many trials have adopted similar approaches of staggered administration, but the outcome for CHK1i thus far has been modest at most (26, 27). Taken together, there is a need to establish optimal dosing schedules for these agents before they could be confidently inducted into the clinic.

We hypothesized that a mechanism-based approach to optimizing the doses and schedules of cell-cycle checkpoint inhibitors with chemotherapy could facilitate the development of more tolerable and effective multitherapeutic strategies. To this end, we sought to build a comparative mechanistic framework of WEE1i and CHK1i in the context of gemcitabine sensitization. Because gemcitabine is a major component of the standard treatment for pancreatic ductal adenocarcinoma, against which few targeted agents have been successfully used, we focused our investigation on preclinical models of pancreatic cancer. Through the concerted application of mathematical modeling and single-cell analyses, our effort uncovered important distinctions in cell-cycle kinetics between WEE1i and CHK1i, leading to the rational implementation of a triple-agent schedule capable of inducing cancer-specific lethality at minimally bioactive single-agent doses.

Materials and Methods

Cell lines and chemicals

All human pancreatic ductal adenocarcinoma lines were obtained from either the European Collection of Cell Cultures or the ATCC, authenticated using either Promega GenePrint10 system or Promega PowerPlex 16HS Kit. MIA PaCa-2 FastFUCCI cell line was generated as previously described (28). Murine pancreatic ductal adenocarcinoma lines DT8082 and K8484 were established from KRas^{G12D} p53^{R172H} Pdx1-Cre mice. All cell lines were routinely verified to be *Mycoplasma*-free using the Mycoprobe Mycoplasma Detection Kit (R&D Systems); the most recent date of testing was 30 January 2018. All cell lines were used within 20 passages following thawing in all experiments. 5-bromo-2'-deoxyuridine (BrdUrd; Sigma), CHIR124 (Selleck Chemicals), gemcitabine (Tocris), MK1775 (AZD1775; Selleck Chemicals), MK8776 (Selleck Chemicals), and roscovitine (Sigma) were dissolved in DMSO (Sigma) in aliquots of 10 to 30 mmol/L as stocks. Final DMSO concentrations were kept constant ($\leq 0.2\%$) in all experiments.

Cytotoxicity and clonogenic assays

For cytotoxicity assays, cells were seeded in 96-well black-walled plates. After 24 hours, cells were treated with a serial dilution of specified agents in an 8 × 8 concentration format, with an extra plate fixed at the time of dosing to determine the T₀ value. After specified days, cells were fixed with trichloroacetic acid and stained with sulforhodamine B. Fluorescent readout was

evaluated using the Infinite 200 PRO microplate reader (Tecan) at excitation and emission wavelengths of 488 and 585 nm, respectively. The T_0 value was subtracted from all wells, and growth inhibition was calculated by expressing it as a ratio of the mean of vehicle control wells. Synergy score was determined as previously described (11). For clonogenic assays, equal number of viable cells were plated 24 hours prior to treatment. After the specified treatment schedule, cells were fixed with 70% methanol and stained with 0.2% crystal violet (Sigma). Colonies were imaged and quantified using GelCount colony counter (Oxford Optronix).

IncuCyte time-lapse imaging

Images were taken with the IncuCyte Live Cell Imaging microscopy (Essen Bioscience) at every three hours under cell culture conditions with 10 to 20× objective. Cell confluence was averaged from multiple fields of view per well. Relative confluence values were calculated by normalizing each value to the time zero (T_0) value.

Quantitative fluorescence-based microscopy

Images of fluorescently labeled samples were acquired using the iCys laser scanning cytometer (CompuCyte; 40× objective) equipped with a motorized Olympus IX71 inverted fluorescence microscope, three lasers (405 nm violet diode laser, 488 nm argon laser, 633 nm helium-neon laser) and three optical filter sets (blue 450/40, green 530/30, far-red 650LP) coupled to photomultiplier (PMT). The in-built iCys software was used to analyze the acquired images. Overlap-proportional Venn diagrams were drawn with the Venn diagram plotter software from Pacific Northwest National Laboratory (<http://omics.pnl.gov/>).

FastFUCCI imaging

The FastFUCCI live-cell assay was performed as previously described (28). Briefly, cells were seeded in glass bottom chamber (ibidi GmbH) and were kept under cell culture conditions. Images were retrieved using a Nikon Eclipse TE2000-E microscope with a 20× long-working distance dry objective and a sCMOS Andor Neo camera. Red and green fluorescence were acquired using a pE-300^{white} CoolLED source of light filtered by Nikon FITC B-2E/C and TRITC G-2E/C filter cubes, respectively. Live-cell time-lapse sequences were split into single channel sequences, and were applied with background subtraction and shading correction. Cell-tracking analysis was performed using the TrackMate plugin available in the Fiji package.

Immunostaining and immunoblotting

For immunostaining, cells seeded in glass bottom chamber (ibidi GmbH) were fixed with 4% paraformaldehyde, stained with antibodies and counterstained with 4',6-diamidino-2-phenylindole (DAPI), 5-Ethynyl-2'-deoxyuridine (EdU) Click-it assay was performed according to manufacturer's instructions (Life Technologies). For immunoblotting, whole-cell extracts were lysed using radioimmunoprecipitation assay (RIPA) buffer (50 mmol/L Tris pH8, 2 mmol/L EDTA, 150 mmol/L sodium chloride, 1% NP-40, 0.5% sodium deoxycholate, 0.1% sodium dodecyl sulfate). Protein concentrations were quantified by the Bio-Rad Protein Assay (Bio-Rad). Equal amounts of protein were resolved using the SDS-PAGE gel system (Life Technologies) and transferred to nitrocellulose membranes using the iBlot Dry Blotting System (Thermo Fisher Scientific). Blots were blocked with Odyssey blocking buffer (LICOR), stained with primary and

secondary antibodies, and analyzed using the Odyssey Infrared Imaging System (LICOR).

Antibodies

Primary antibodies used were from Cell Signaling unless otherwise mentioned: β -actin (Abcam #ab6276), BrdUrd (BD Pharmingen #555627), cleaved caspase-3 (#9664), CDK1 (Abcam #ab18), CDK1 Y15 (#9111), CHK1 (#2360), CHK1 S296 (#2349), CHK1 S345 (#2348), ENT1 (Abcam #ab135756), H2AX (#7631), H2AX S139 (Millipore #05-636), H3 (#9715), and H3 S10 (#3377). For secondary antibodies, Alexa Fluor 488 (#4408, #4412) and Alexa Fluor 647 (#4410, #4414) from Cell Signaling Technology were used in immunostaining. IRDye800-conjugated (#925-32210, #926-33210) and IR680-conjugated (#926-68070, #926-68021) antibodies from LICOR were used in immunoblotting.

Mouse studies

All mouse experiments were carried out in the CRUK Cambridge Institute Biological Resources Unit, in accordance with the UK Animals (Scientific Procedures) Act 1986, with approval from the CRUK Cambridge Institute Animal Ethical Review and Welfare Body. Subcutaneous xenografts of MIA PaCa-2 cells were conceived by implanting 5×10^6 cells in 50% Matrigel/50% PBS, in the right flank of 6 to 10 weeks' old female BALB/c nude mice (Charles River Laboratories). For efficacy study, mice with established tumor ($\sim 300 \text{ mm}^3$) were randomized into three per group and treated accordingly. Gemcitabine (LKT Laboratories) and MK8776 (Selleck Chemicals) were codissolved in 20% hydroxypropyl- β -cyclodextrin (Vehicle I; Sigma) and were given to mice at 50 mg/kg intraperitoneally. MK1775 (Selleck Chemicals) was dissolved in 0.5% methylcellulose (Vehicle II; Sigma) and was given to mice at 60 mg/kg by oral gavage. For each weekly cycle in the efficacy study, Vehicle I or agents (gemcitabine, MK8776) were given first followed 4 hours later by Vehicle II or MK1775, on day 0 and day 3. Mice were killed once the averaged tumor size quadrupled ($\sim 1,200 \text{ mm}^3$). For analysis, the normalized average of the tumor size per group during treatment phase was modeled by means of a mixed model, with treatment groups and time (polynomial of degree 2) as predictors in the fixed part of the model. The within-mouse and time dependences were taken into account by means of random intercepts and slopes as well as a moving average for the within-mouse residuals. Model checks and sensitivity analyses suggested a good model fit. Restricted maximum likelihood estimates, SEs, and the degrees of freedom, t values and P values of the corresponding significance test for all fixed effect parameters were calculated.

Quantitative LC/MS-MS

Snap-frozen tumor tissue was homogenized in ice-cold acetonitrile [50% (v/v) containing tetrahydrouridine] using a Precellys 24 homogenizer (Bertin Technologies). An aliquot (50 μL) of the homogenate was transferred to a clean tube containing 150 μL of ice-cold acetonitrile [50% (v/v) containing internal standards CTP- ^{13}C , $^{15}\text{N}_2$ from Sigma and dFdC- ^{13}C , $^{15}\text{N}_2$ from Toronto Research Chemicals] followed by centrifugation at 21000 G for 20 minutes. The supernatant was transferred to a clean tube and evaporated to dryness under air. The residue was resuspended in water and 15 μL was injected into the mass spectrometer. For plasma samples, 25 μL was added to 150 μL of ice-cold acetonitrile [85% (v/v) containing internal standard dFdC- ^{13}C , $^{15}\text{N}_2$] and the mix was processed as described for tissue homogenate.

Chromatography was performed using an Accela pump (Thermo Fisher Scientific) and the analytes were separated on a PGC Hypercarb column (100 × 2.1 mm ID, 5 μm; Thermo Fisher Scientific) fitted with a guard column (Hypercarb 10 × 2.1 mm, 5 μm; Thermo Fisher Scientific) with (i) 10 mmol/L ammonium acetate, pH10 and (ii) acetonitrile as mobile phases. LC/MS-MS was performed on a TSQ Vantage triple stage quadrupole mass spectrometer (Thermo Fisher Scientific) fitted with a heated electrospray ionization (HESI-II) probe operated in positive and negative mode at a spray voltage of 2.5 KV, capillary temperature of 150°C. Data acquisition was performed using LC Quan2.5.6 (Thermo Fisher Scientific).

Immunohistochemistry

IHC was performed on formalin-fixed, paraffin-embedded sections of tumor after heat-induced epitope retrieval by sodium citrate at 100°C for 10 to 20 minutes, using Bond Polymer Refine Detection Kit on the automated Bond system according to manufacturer's instructions (Leica). Prestaining dewaxing and rehydration, as well as poststaining dehydration and clearing, were performed on Leica ST5020 Multistainer. 3,3'-Diaminobenzidine (DAB) enhancer (Leica) was applied to increase contrast between chromogen-specific staining and the slide background. Slides were mounted using Leica CV5030 Coverslipper Workstation and scanned using a ScanScopeXT (Aperio Technologies). Quantification was performed using the ImageScope (Aperio Technologies). Antibodies used for IHC were geminin (Novocastra #NCL-L-GEMININ), H2AX S139 (Cell Signaling Technology #9718) and H3 S10 (Upstate #06-570).

Statistical analysis

Data were analyzed using GraphPad Prism built-in statistical tests indicated in relevant figure legends. The following asterisk system for *P* value was used: *, *P* ≤ 0.05; **, *P* ≤ 0.01; ***, *P* ≤ 0.001; and ****, *P* ≤ 0.0001.

Results

WEE1i and CHK1i elicit chemosensitivity with different potency

In human (AsPC-1, Capan-1, HPAF-II, MIA PaCa-2, Panc-1) and murine (DT8082, K8484) pancreatic adenocarcinoma cell lines showing different cell doubling times, WEE1i (MK1775) demonstrated a ~6-fold range of single-agent sensitivity, with GI_{50} from 80 ± 19 nmol/L to 455 ± 28 nmol/L (Supplementary Fig. S1A). Irrespective of the degree of sensitivity, all tested cell lines were sensitized to gemcitabine upon WEE1 inhibition (Fig. 1A; Supplementary Fig. S1B–S1E). To analyze the data in pharmacologically meaningful terms, we used two independent mathematical models of synergy (Bliss and Loewe) to assess the combination of gemcitabine and MK1775 across a broad range of concentrations (Fig. 1A). Consistently, both models identified a synergistic interaction between the two agents. However, when we compared gemcitabine + WEE1i synergy metrics to those of gemcitabine + CHK1i (MK8776), we found that noninhibitory concentrations (< GI_{10}) of CHK1i with 10 to 30 nmol/L gemcitabine achieved greater synergy scores than equipotent single-agent concentrations of WEE1i (Fig. 1A). The same conclusion was reached with the use of two other structurally distinct CHK1-specific inhibitors, CHR124 and S1181, suggesting that the differential synergism was not because of off-target effects (11). To further substantiate these findings, we tested the combinations

on a more gemcitabine-resistant cell line Panc-1, again demonstrating that CHK1i elicited greater synergistic inhibition with gemcitabine than WEE1i at noninhibitory concentrations (Supplementary Fig. S1E). Crucially, at equivalent inhibitory concentration ratios (Supplementary Fig. S1E), long-term clonogenic assays show similar extents of growth inhibition in both gemcitabine + WEE1i and gemcitabine + CHK1i, establishing in each instance a bona fide synergy that was durable (Fig. 1B). Together, these data demonstrate that, although WEE1i and CHK1i induce gemcitabine hypersensitivity, there exists a disparity in synergy potential where CHK1i chemosensitizes cells more effectively than WEE1i when given concurrently with gemcitabine.

WEE1i invokes two forms of DNA damage with gemcitabine

WEE1i is commonly presumed to deregulate the DNA replication apparatus in a similar manner as CHK1i. To test this notion, we exposed MIA PaCa-2 cells to the minimum synergistic concentration of WEE1i (300 nmol/L MK1775) and CHK1i (1 μmol/L MK8776) with 10 nmol/L gemcitabine that elicited equivalent growth inhibition, as derived from their respective synergy metrics (Fig. 1A). Notably, these concentrations were clinically attainable and tolerable in humans (20, 29, 30). Although gemcitabine alone induced a marginal increase in S-phase fraction, both WEE1i and CHK1i with gemcitabine caused robust S–G₂ accumulation (Supplementary Fig. S1F). Activation of the CDK members promotes DNA replication origin firing. Accordingly, suppression of origin firing with a broad CDK inhibitor roscovitine partially reversed growth inhibition by gemcitabine + WEE1i and gemcitabine + CHK1i (Supplementary Fig. S1G; ref. 11). However, as opposed to other deoxynucleotides, supplementation of deoxycytidine alone, which antagonizes gemcitabine by competing for incorporation into DNA, was sufficient to rescue proliferation in both combinations (Supplementary Fig. S1H–S1I). These findings indicate that, although nucleotide exhaustion induced by WEE1i and CHK1i has frequently been reported as the cause of gemcitabine sensitization, at synergistic concentrations, increased gemcitabine incorporation following aberrant origin firing represents the predominant source of replication stress.

The differences in replication perturbation by WEE1i and CHK1i, if any, could be masked by bulk responses of cell population. To address this limitation, we turned to measuring molecular markers of genomic stress and damage in each individual cell. Exposure to gemcitabine + WEE1i and gemcitabine + CHK1i for 24 hours induced comparable levels of RPA32 S4/8 and γH2AX with few (<5%) cleaved caspase-3-positive cells, establishing a correlation between extensive replication stress (RPA32 S4/8) and apoptosis-independent DNA damage (γH2AX without substantial cleaved caspase-3; Fig. 1C and Supplementary Fig. S1J). However, although the expression of these markers was comparable in the combinations, quantitative single-cell analysis reveals that cells exposed to WEE1i alone harbored significantly higher levels of γH2AX compared with CHK1i (Fig. 1C; Supplementary Fig. S1K). No difference in replication stress levels as measured by hyperphosphorylated RPA32 was found between these two conditions. Furthermore, a significantly lower fraction (54 ± 5%) of the damaged population exposed to gemcitabine + WEE1i featured replication stress, in contrast to the 72 ± 3% in gemcitabine + CHK1i (Fig. 1D). Together, these data suggest that the genomic damage created by WEE1i in the presence of gemcitabine could evolve from replication stress-dependent and

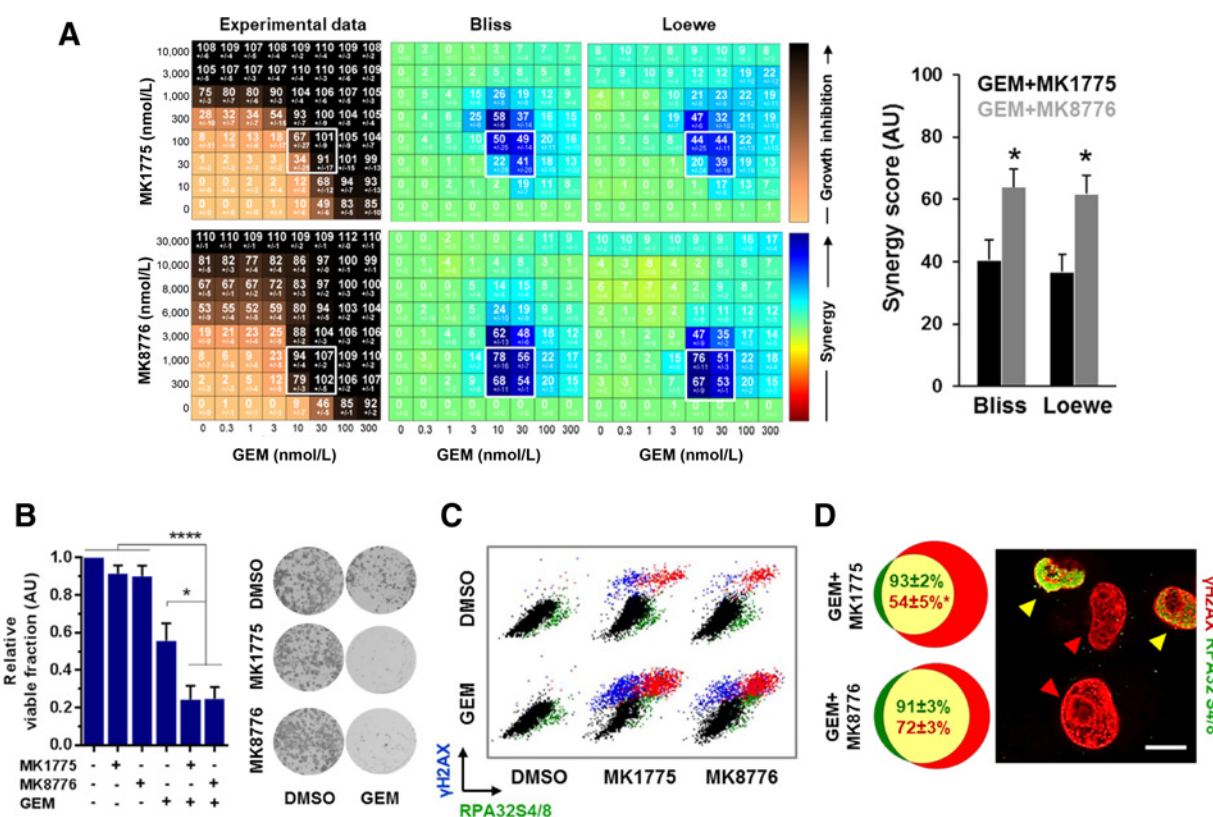


Figure 1.

WEE1i and CHK1i synergize with gemcitabine with different potency. **A**, Combination assay. MIA PaCa-2 cells were treated for 72 hours. Data were analyzed with two synergy mathematical models. Combinations of 10 to 30 nmol/L gemcitabine (GEM) with a pair of noninhibitory (<math><G_{H_0}</math>) equivalent concentrations of MK1775 and MK8776 are boxed in white. Bar graph shows the mean synergy score within the boxed surface. Data are represented as mean \pm SEM, $n = 3$. A two-tailed t test was performed; *, $P \leq 0.05$. **B**, Clonogenic assay. Panc-1 cells were treated for 72 hours (30 nmol/L GEM, 300 nmol/L MK1775, 1 μ mol/L MK8776) and were left to grow after washout for 10 days. Data are represented as mean \pm SEM, $n = 3$. A one-way ANOVA analysis was performed; *, $P \leq 0.05$, ****, $P \leq 0.0001$. **C**, Quantitative immunofluorescence of MIA PaCa-2 cells treated for 24 hours (10 nmol/L GEM, 300 nmol/L MK1775, 1 μ mol/L MK8776). Each blue, green, or red dot marks a cell positive for γ H2AX, RPA32 S4/8, or both, respectively. **D**, Quantification of overlap between γ H2AX (red) and RPA32 S4/8 (green), in MIA PaCa-2 cells treated for 24 hours (10 nmol/L GEM, 300 nmol/L MK1775, 1 μ mol/L MK8776). Green, percentage of γ H2AX-positive cells in RPA32 S4/8-positive population; red, percentage of RPA32 S4/8-positive cells in γ H2AX-positive population. Red arrowhead, γ H2AX-positive cell; yellow arrowhead, γ H2AX/RPA32 S4/8 double-positive cell. Data are represented as mean \pm SEM, $n = 5$. At least 2,000 cells per condition per replicate were analyzed. A two-tailed t -test was performed; *, $P \leq 0.05$. Scale bar, 25 μ m.

-independent routes, raising a key question on how these effects are manifested in each phase of the cell cycle.

Passage through mitosis underlies WEE1i chemosensitization

The above findings implied a biological consequence of WEE1 inhibition that was distinct from replication deregulation. To pursue this hypothesis, we first quantified the cell-cycle durations using the live single-cell FastFUCCI assay (28). Compared with vehicle and WEE1i-only conditions, synergistic 10 nmol/L gemcitabine + 300 nmol/L WEE1i resulted in an increase in mean S-G₂ residence time (Fig. 2A). However, the increase in S-G₂ duration induced by 10 nmol/L gemcitabine + 300 nmol/L WEE1i did not differ from that by 10 nmol/L gemcitabine alone. In contrast, mean S-G₂ duration was longer with synergistic concentrations of gemcitabine+CHK1i than with 10 nmol/L gemcitabine (Fig. 2B). This disparity was in spite of both combinations prolonging G₁ to an extent that was as much as that in high (100 nmol/L) concentration of gemcitabine (Supplementary Fig. S2A–S2B). Notably, both WEE1i and CHK1i partially abro-

gated the S-G₂ prolongation induced by 100 nmol/L gemcitabine (Fig. 2A and B), in agreement with the concentration-dependent model of G₂-M abrogation (11). Together, the data reveal differential cell-cycle timing effects by WEE1i and CHK1i at synergistic concentrations with gemcitabine, even though the ultimate consequence is a delay in S-G₂ (Supplementary Fig. S1F).

The integrity of S-G₂ and mitosis are dynamically linked across cell generations (31). To systematically deconstruct this biological flux, we measured the duration of each cell-cycle stage of first- and second-generation cells following gemcitabine + WEE1i. Surprisingly, despite the collective delay in S-G₂ phase (Fig. 2A; Supplementary Fig. S1F), the S-G₂ phase of first-generation cells in the treated sample was comparable with that in control, suggesting an initial reprieve of S-G₂ impediment (Fig. 2C; Supplementary Fig. S2C). This observation was in contrast to CHK1i, which could prolong the S-G₂ residence time as much as two folds in the first cycle (11). Upon entering mitosis, mitotic duration in the gemcitabine + WEE1i-treated cohort was more than three times longer than control (Fig. 2C; Supplementary Fig. S2C). All

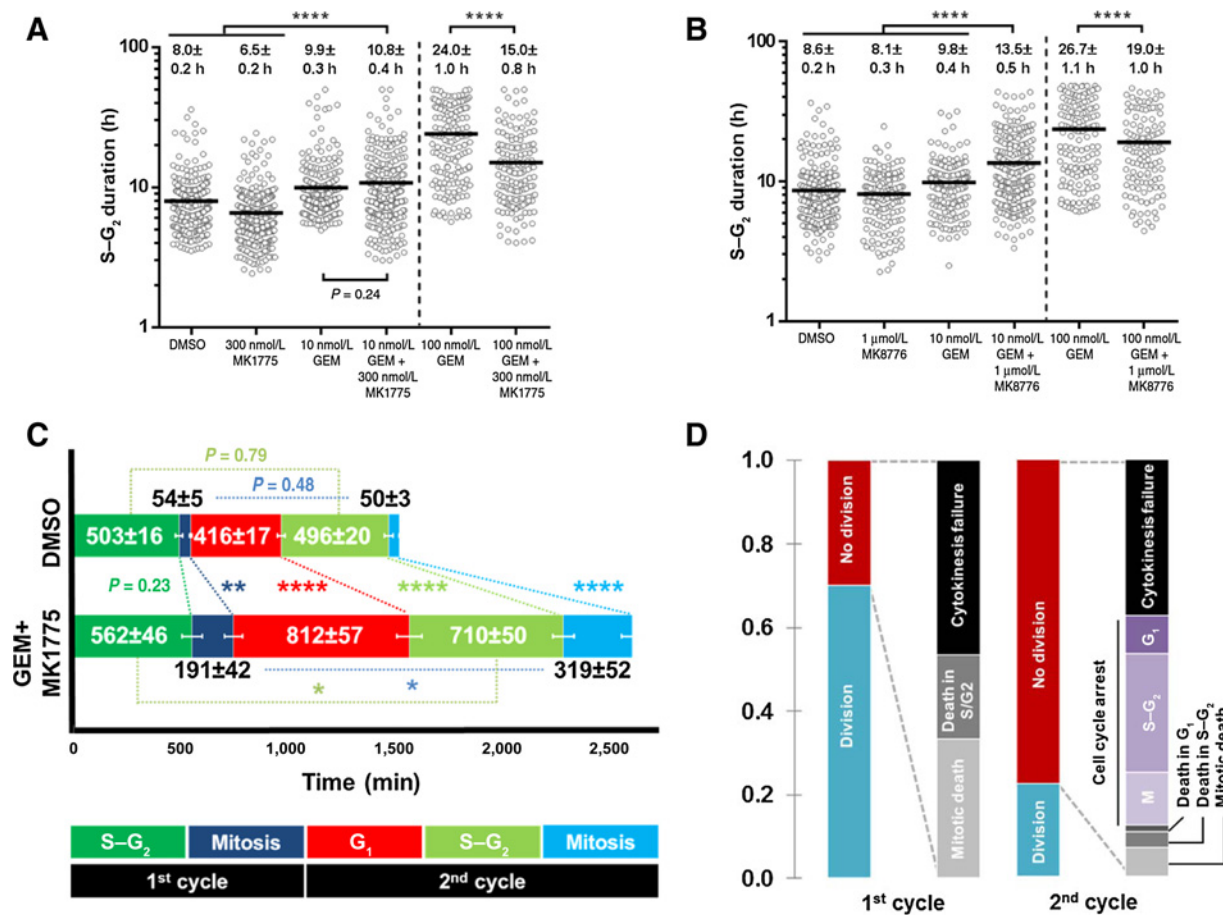


Figure 2.

Mitotic stress underlies WEE1i cytotoxicity. **A** and **B**, S-G₂ duration of MIA PaCa-2 FastFUCCI cells treated as indicated. At least 100 cells per condition were analyzed. Data are represented as mean ± SEM. A one-way ANOVA analysis was performed; ****, $P \leq 0.0001$. **C**, Cell-cycle duration of MIA PaCa-2 FastFUCCI cells treated with DMSO or 10 nmol/L gemcitabine (GEM) + 300 nmol/L MK1775. A total of 243 cells were analyzed. Data are represented as mean ± SEM. A two-tailed t test was performed; *, $P \leq 0.05$, **, $P \leq 0.01$, ****, $P \leq 0.0001$. **D**, Percentage of MIA PaCa-2 FastFUCCI cells in the first and second cycles that underwent division or not following 10 nmol/L gemcitabine + 300 nmol/L MK1775. Fraction of nondividing cells was further categorized according to cell fate. A total of 122 cells were analyzed.

subsequent cell-cycle stages in the treated sample were also significantly protracted. To interpret these spatiotemporal details in the context of cellular responses, we tracked the fate evolution of gemcitabine + WEE1i-treated cells using three approaches. First, quantification of individual cell fates shows that (i) the majority of nondivision events in the first cycle arose during or after mitosis (47% cytokinesis failure and 33% mitotic death, compared with 20% death in S-G₂), and that (ii) the incidence of nondivision escalated from 30% in the first cycle to 78% in the second cycle (Fig. 2D). Second, cross-generation analysis indicates that (i) 75% of second-generation cells arising from productive first-generation cells subsequently experienced a deleterious event (cytokinesis failure, cell cycle arrest, cell death), but (ii) all unproductive first-generation cells (i.e., those that underwent cytokinesis failure) were again unproductive in the second cycle (Supplementary Fig. S2D). Third, pairwise sister cell examination reveals that (i) a productive sister did not necessarily predict a productive counterpart, but (ii) if one of the sisters was unproductive, the other sister tended to share the same fate (Supplementary Fig. S2E). Together, these data demonstrate that WEE1i chemosensitization is most profound only after a cell launches

into its first mitosis, with deleterious events occurring during or after mitosis and persisting over generations.

WEE1i effects on cell-cycle kinetics are temporally coordinated

Abrogation of G₂-M checkpoint is frequently regarded as the mechanism of WEE1i and CHK1i, but precisely when and in which cell-cycle stage the event could arise are unknown. We previously showed that CHK1i at synergistic concentrations with gemcitabine in fact did not effectively override G₂-M checkpoint (11). In contrast, gemcitabine + WEE1i induced a demonstrable reduction in inactive CDK1 Y15 that correlated with an increase in mitotic marker H3 S10, suggesting CDK hyperactivation and accumulation of mitotic cells (Fig. 3A). Comparative single-cell analysis shows that the mitotic DNA content in gemcitabine+WEE1i sample was significantly lower than its single-agent controls and CHK1i, further indicating a mitotic subset with incomplete (<4N) genome (Fig. 3B; Supplementary Fig. S3A). As measured by the level of γ H2AX, both WEE1i and CHK1i combinations with gemcitabine invoked comparable degrees of genomic damage in S-G₂-M cells (Supplementary Fig. S3B). However, quantitative immunofluorescence shows that WEE1i

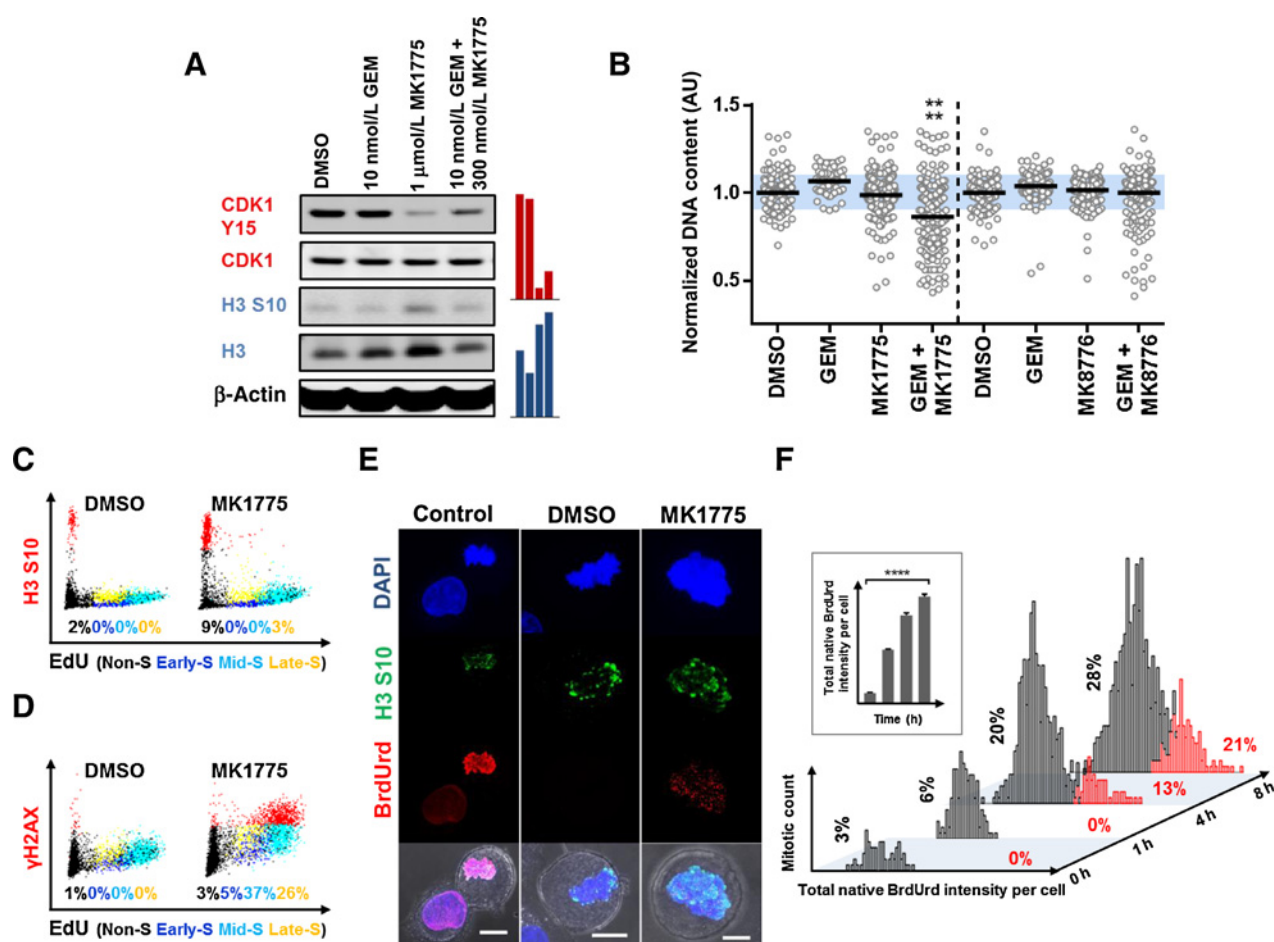


Figure 3.

Inimical effects of WEE1i are spatiotemporally defined. **A**, Immunoblotting for MIA PaCa-2 cells treated for 24 hours. The graph shows densitometric analysis of CDK1 Y15/CDK1 or H3 S10/H3, relative to DMSO. **B**, Quantification of DNA content of mitotic MIA PaCa-2 cells treated for 24 hours [10 nmol/L gemcitabine (GEM), 300 nmol/L MK1775, 1 μmol/L MK8776]. Data are represented as mean, normalized to DMSO. A one-way ANOVA analysis was performed; ****, $P \leq 0.0001$. **C**, Quantification of mitotic Panc-1 cells in S phase. Cells were treated with 10 μmol/L EdU for 45 minutes, followed by 3 μmol/L MK1775 for 1 hour. S and non-S phase cells were identified based on EdU and DNA contents. Percentage of H3 S10-positive cells is shown. **D**, Quantification of damaged Panc-1 cells in S phase, treated as in **C**. Percentage of γ H2AX-positive cells is shown. **E** and **F**, Quantification of mitotic Panc-1 cells harboring ssDNA. Cells were grown with 10 μmol/L BrdUrd for 48 hours, treated with 3 μmol/L MK1775, and immunostained for native BrdUrd. In **E**, the first column shows positive control, where sample was acid-denatured to confirm BrdUrd incorporation. Scale bar, 10 μm. In **F**, percentage of mitotic cells is in black and percentage of native BrdUrd-positive mitotic cells (out of the respective mitotic fractions) in red. At least 2,000 cells were analyzed per time-point. Inset shows the total native BrdUrd intensity per mitotic cell. A one-way ANOVA analysis was performed; ****, $P \leq 0.0001$.

alone induced greater genomic damage in the mitotic fraction than CHK1i alone. This difference was particularly pronounced when WEE1i was combined with gemcitabine, where 95% of mitotic cells exhibited dramatic DNA fragmentation decorated with intense γ H2AX, compared with 60% in gemcitabine+CHK1i (Supplementary Fig. S3B). Collectively, these data illustrate the superiority of WEE1i over CHK1i in driving cancer cells into stressful mitotic states.

The persistent presence of a mitotic subset with normal 4N DNA under WEE1i conditions suggested that not all affected cells were compelled into mitosis directly from S phase (Supplementary Fig. S3C). To resolve how WEE1i impacted the S and mitotic phases, we segregated asynchronous cell population into early, mid and late S phases as well as mitosis using quantitative multiparametric image-based cytometry. Acute WEE1 inhibition for 1 hour delayed mitotic progression but drove a fraction of only

late S-phase cells into mitosis (Fig. 3C; Supplementary Fig. S3D). Simultaneously, about one in every three cells in mid and late S-phase compartments experienced DNA damage (Fig. 3D; Supplementary Fig. S3E). Costaining of H3 S10 and native BrdUrd confirms that mitotic accumulation started within 1 hour; however, it was after this timepoint that single-stranded DNA (ssDNA) became evident in the mitotic population (Fig. 3E and F; Supplementary Fig. S3F). Notably, ~40% to 50% of <4N premature mitotic cells harbored ssDNA at 4 and 8 hours following treatment, suggesting that these cells were under-replicated and may still be replicating (Supplementary Fig. S3F). Pulse labeling with EdU indicates that there was indeed active DNA synthesis predominantly in late-S and prophase cells, in line with stress-triggered mitotic DNA replication (Supplementary Fig. S3G; ref. 32). Within the same timeframe, there was neither mitotic block nor accumulation of ssDNA-harboring mitotic cells with

CHK1 inhibition, reinforcing the notion that the mitotic consequences of CHK1i are not acute (Supplementary Fig. S3H). Together, these data show three temporally coordinated but independent responses following WEE1 inhibition: (i) Upon initial insult, replication stress-independent mitotic block is enacted in mitotically competent (4N) cells. (ii) Concurrently, a fraction of mid to late S-phase cells encounter genomic damage. (iii) At the same time, a subset of late S-phase (<4N) cells prematurely slip into mitosis. These premature mitotic cells persist to synthesize DNA ineffectively, with WEE1i-induced CDK hyperactivation further aggravating the mitotic state by delaying its resolution.

Scheduled gemcitabine/CHK1i/WEE1i sustains tumor cell inhibition

Our data thus far show that the respective interactions of WEE1i and CHK1i with gemcitabine are synergistic but differ in several

other pharmacologic terms (Fig. 4A). First, although both inhibitors induce gemcitabine hypersensitivity to equal synergistic levels, CHK1i generates greater synergy at equivalent single-agent inhibitory concentrations than WEE1i (Fig. 1; Supplementary Fig. S1). Second, although both inhibitors trigger replication stress, WEE1i prolongs S-G₂ only after the first cell cycle, with catastrophic events occurring chiefly during or following mitosis, coincident with cross-generation damage accrual (Fig. 2; Supplementary Fig. S2). Third, unlike CHK1 inhibition where mitotic distress originates from a deregulated replication checkpoint, WEE1i directly affects both mitotic entry and progression by potent disinhibition of mitotic CDK (Fig. 3; Supplementary Fig. S3). From a therapeutic perspective, these distinctions are consistent with the reported synergy between WEE1i and CHK1i (14, 15). Furthermore, analysis of two cancer cell line databases shows that WEE1 and CHK1 expression were significantly correlated in diverse cancer subtypes (Fig. 4B; Supplementary Fig. S4A).

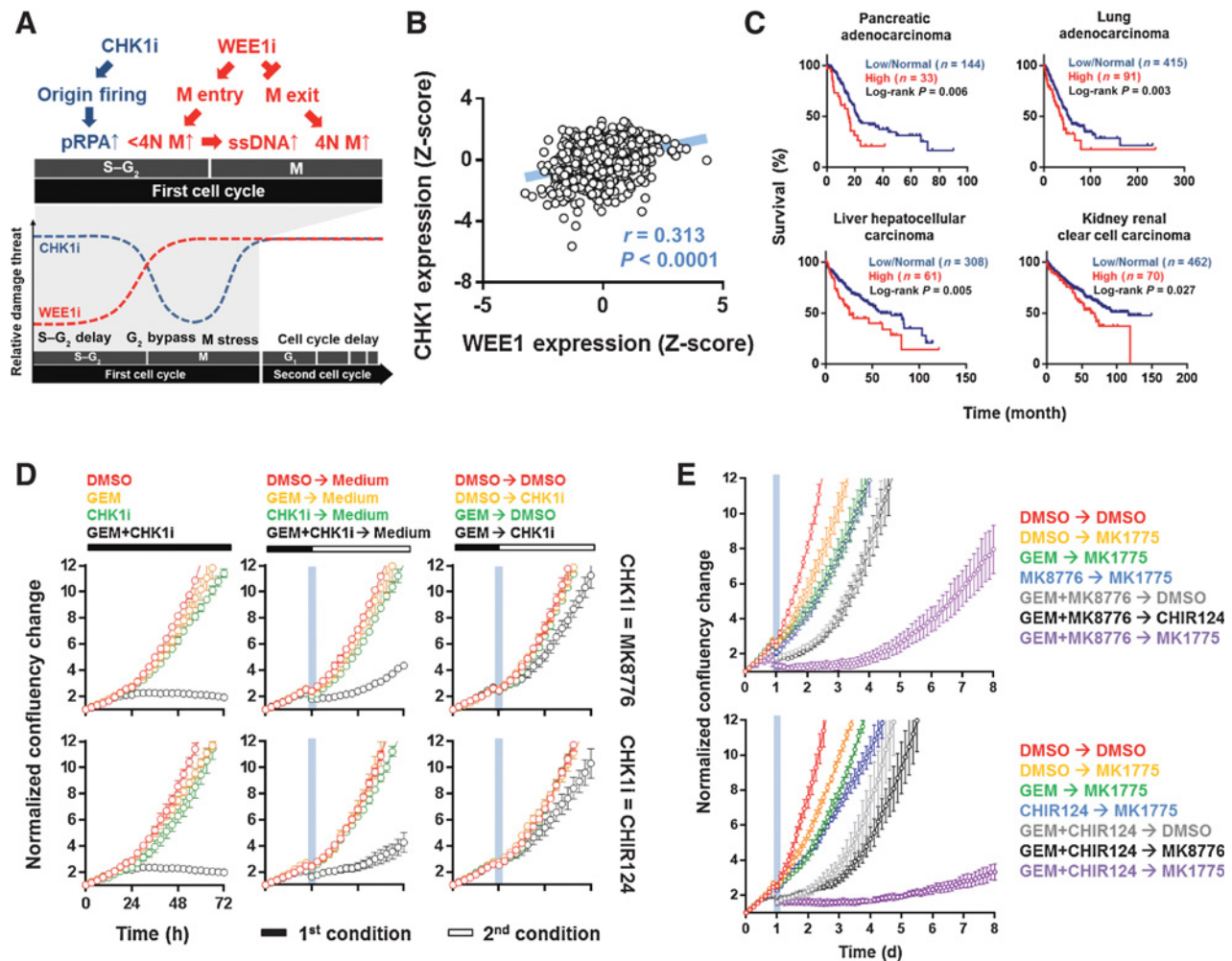


Figure 4. A gemcitabine/CHK1i/WEE1i regimen enhances tumor cell suppression. **A**, Schematics of the spatiotemporal effects of WEE1i and CHK1i. **B**, Correlative analysis between WEE1 and CHK1 mRNA expression in 967 tumor cell lines from the Cancer Cell Line Encyclopedia project. Pearson correlation coefficient *r* and *P* values are indicated. **C**, Kaplan-Meier analysis of RNASeq V2 data on WEE1 or CHK1 expression and patient survival in indicated primary tumor samples. Tumors with mRNA expression Z-score +1.5 were considered as tumors with high expression. Data were sourced from the TCGA Research Network. **D-E**, Real-time growth kinetics of MIA PaCa-2 cells treated as indicated [10 nmol/L gemcitabine (GEM), 1 μmol/L MK8776, 20 nmol/L CHIR124, 300 nmol/L MK1775]. Data are represented as mean ± SEM, *n* = 3.

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In agreement, cell lines with high WEE1 or CHK1 expression were more sensitive to a WEE1/CHK1 dual inhibitor 681640, but such association was less robust in a CHK1/2-specific inhibitor AZD7762, arguing from a genetic perspective the advantage of cotargeting the two kinases (Supplementary Fig. S4B; ref. 33). Importantly, the trend of a positive correlation between WEE1 and CHK1 expression was also observed in multiple patient samples of different tumor origins (Supplementary Fig. S4C). In primary pancreatic ductal adenocarcinoma, high expression of these kinases was a strong determinant of poor overall survival and disease-free survival (Fig. 4C; Supplementary Fig. S4D). Equally, overexpression of these kinases was associated with shorter overall survival in three other aggressive tumors, including lung adenocarcinoma where gemcitabine is also used routinely. Together, these findings offer a compelling rationale to combine WEE1i with CHK1i to maximize gemcitabine sensitivity.

To determine optimal dosing schedules for WEE1i and CHK1i with gemcitabine, we used real-time *in vitro* imaging and found that delayed administration of CHK1i at 24 hours relative to gemcitabine did not lead to substantial growth inhibition (Fig. 4D). Conversely, concurrent administration yielded dramatic growth suppression, even when gemcitabine and CHK1i were removed after 24 hours. Given that G₂-M abrogation was not the predominant mechanism of synergy for gemcitabine+CHK1i, we hypothesized that the combination could be further enhanced by driving G₂-M bypass using WEE1i. We determined the growth kinetics of MIA PaCa-2 cells exposed to the schedule of gemcitabine + CHK1i (MK8776 or CHIR124) for 24 hours followed by WEE1i (Fig. 4E). Cell growth inhibition was remarkably durable with the triple regimen compared with vehicle and dual agent controls. Notably, substitution of WEE1i in the triple regimen with another CHK1i (i.e., gemcitabine + MK8776 followed by CHIR124, or gemcitabine + CHIR124 followed by MK8776) failed to recapitulate the durable response. Long-term clonogenic assays confirm effective growth suppression of the proposed triple regimen and further show that administering gemcitabine + CHK1i followed by WEE1i was more effective than administering gemcitabine + WEE1i followed by CHK1i (Supplementary Fig. S4E and S4F). Together, the sequence-dependent cooperativity of WEE1i with gemcitabine + CHK1i validates the differential mechanistic properties of WEE1i and CHK1i, affirms the lethality of G₂-M bypass, and demonstrates the antitumor potential of the proposed triple regimen.

***In vivo* studies show antitumor potential of minimal-dose multitherapy**

To evaluate whether the *in vitro* mechanistic findings could be recapitulated *in vivo*, we chose a low dose of gemcitabine (25 mg/kg) compared with the "full" maximum tolerable dose (MTD) dose (100–150 mg/kg) administered to mice bearing MIA PaCa-2 xenografts. An intraperitoneal bolus of 25 mg/kg MK8776 increased gemcitabine-induced CHK1 S345 phosphorylation and abrogated CHK1 S296 autophosphorylation for at least the first 4 hours, indicating target engagement (Supplementary Fig. S5A–S5B). These changes were accompanied by an increase in γ H2AX and RPA32 S4/8 without overt CDK1 hyperactivation or H3 S10 upregulation, consistent with the lack of G₂-M checkpoint bypass we previously established with CHK1i S1181 (Fig. 5A; Supplementary Fig. S5B–S5C; ref. 11). Quantitative IHC further confirms high genomic damage and low mitotic index with aberrant mitoses in the combination cohort (Fig. 5B; Supplementary Fig.

S5D–S5E). Moreover, gemcitabine + MK8776 induced an accumulation of geminin-positive cells by 8 hours, indicative of high S–G₂ fraction as observed in cell lines (Fig. 5C). This accumulation was in concert with intratumoral elevation of active gemcitabine metabolite 2',2'-difluoro-2'-deoxycytidine triphosphate (dFdCTP; Fig. 5D). MK8776 did not significantly alter the pharmacokinetics of native gemcitabine (2',2'-difluoro-2'-deoxycytidine, dFdC) in tumor and in plasma, implying that there was neither increased drug uptake in tumor nor decreased drug clearance in blood (Supplementary Fig. S5F). There was also no obvious difference between both treatment arms in the expression of a major gemcitabine transporter ENT1 (Supplementary Fig. S5B–S5C). Having determined that acute administration of low gemcitabine and MK8776 in mice induced molecular responses observed *in vitro*, we tested the efficacy of the proposed triple schedule. For stringent comparison, we used doses and treatment frequencies that were equal to or lower than those previously established to be suboptimal in MIA PaCa-2 xenograft model, which typically exhibits marginal response even to the highest doses of gemcitabine combinations (34–37). As the *in vitro* findings predicted, the triple minimal-dose regimen was most effective in the growth suppression of established (~300 mm³) tumors compared to single- and double-agent arms (Fig. 5E). During the treatment phase (day 0–26), the triple regimen was the only group that trended towards a difference from the vehicle group in terms of tumor volume ($P = 0.04$ by *t* test, $P = 0.08$ by Dunnett's multiple-comparison test). Following the treatment phase, the regimen delayed tumor growth by about 10 days relative to control, before the averaged tumor size quadrupled. Importantly, there was no overt weight loss in mice treated with the triple combination, suggesting that the regimen did not impact physiological functions at least within the period of investigation (Supplementary Fig. S5G). Together, the *in vivo* data support the *in vitro* mechanistic findings and demonstrate the underappreciated feasibility of minimal-dose multitherapy in achieving tumor control.

Discussion

Targeting cancer with cell-cycle checkpoint inhibitors in combination with chemotherapy is conceptually attractive, but the success has thus far been largely dependent on serendipity (38, 39). Here, we used a mechanism-guided approach to systematically combine classic cytotoxic agent gemcitabine with two DDR clinical candidates CHK1i and WEE1i. We first established synergy between pairs of these agents through mathematical modeling of individual dose–response curves, simultaneously identifying the concentration ratios that yielded optimal growth inhibition. From a series of single-cell studies conducted at these synergistic ratios, we uncovered distinct cell-cycle kinetics between CHK1i and WEE1i, contrary to the common presumption that these inhibitors affect the cell-cycle similarly. Notably, upon initial exposure, CHK1i induced S-phase deregulation in cells more readily than WEE1i, which itself had more direct inimical effects on mitosis. Based on these distinctions, we devised a minimal-dose gemcitabine/CHK1i/WEE1i triple regimen that achieved tumor control without additional toxicity.

Our work represents a preclinical proof of concept that MTD is not necessarily the best approach in developing multitherapeutic regimens. To date, most clinical studies use MTD as a standard for dose selection of chemotherapy. This paradigm stems from

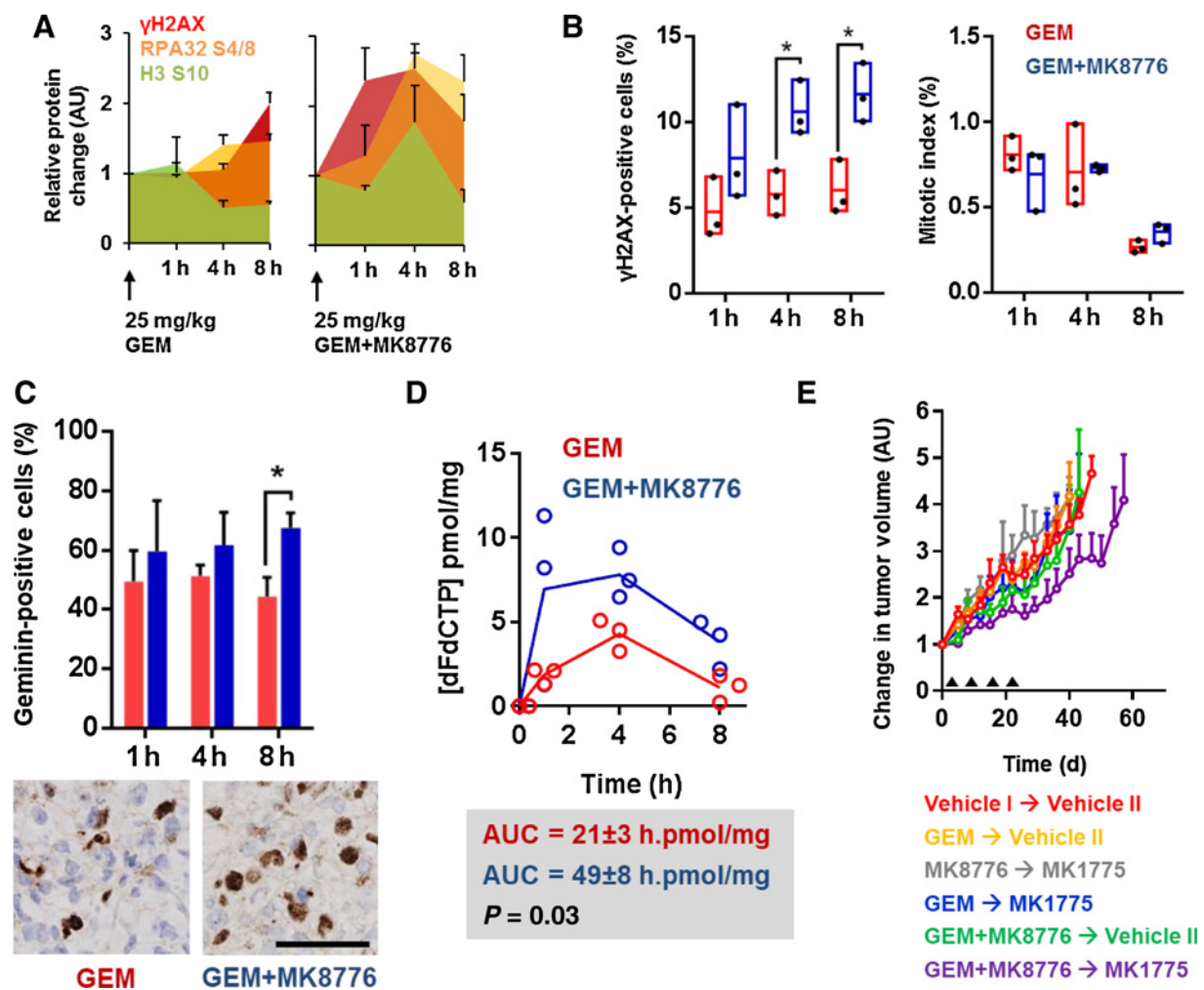


Figure 5.

In vivo studies show antitumor potential of the triple regimen. **A**, Quantification of immunoblotting of tumor samples from MIA PaCa-2 xenografts treated and harvested as indicated. Data are represented as mean \pm SEM, $n = 3$. **B**, Quantification of IHC of tumor samples from **A**. γ H2AX and H3 S10 were used as a marker of DNA damage and mitosis, respectively. Middle line, mean. A two-tailed t test was performed; *, $P \leq 0.05$. **C**, Quantification of geminin-positive cells in tumor samples from **A**. Data are represented as mean \pm SEM, $n = 3$. A two-tailed t test was performed; *, $P \leq 0.05$. Scale bar, 50 μ m. **D**, Pharmacokinetic profile of GEM. Tumor samples from MIA PaCa-2 xenografts treated with either 25 mg/kg GEM or 25 mg/kg gemcitabine + MK8776 were analyzed for the active metabolite of gemcitabine (dFdCTP) at specified time-points. Area under the curve (AUC) and P values are indicated. **E**, Change in tumor volume of MIA PaCa-2 xenografts. Mice were treated as indicated for four consecutive weekly cycles. Black triangle on the x-axis denotes start of each dosing cycle. Data are represented as mean \pm SEM, $n = 3$.

historical observations that response to cytotoxic agents often correlates with dose. Recent advances in tumor evolution have begun to question the basis of MTD, given the potential rapid expansion of resistant clones from intensive drug-induced selection (40, 41). The advent of targeted therapies, which unlike cytotoxic drugs exploit cancer-specific features, has also challenged the relevance of MTD (42). In one instance, meta-analysis of 24 clinical trials shows similar outcomes between patients treated with low- and high-dose targeted agents, with the latter cohorts having higher dropout rates due to cumulative toxicities (43). These findings were supported by another study demonstrating through computational modeling that drug concentrations lower than the MTD could be equally efficacious (44). The complexity of identifying optimal doses escalates in the context of drug combinations, where their clinical benefits are tempered by

further risks of off-target effects. Evidently, drug-specific dose-response kinetics and dose-dependent drug effects are formidable challenges in the design of multitherapy, and preclinical drug development should be primed to address these considerations from the outset.

An immediate clinical impact of our study is the re-evaluation of current schedules used in human trials for CHK1i and WEE1i. We demonstrate that concurrent, not sequential, treatment of gemcitabine and CHK1i leads to better tumor cell inhibition. This inhibition is further enhanced by subsequent addition of WEE1i and not continuation of CHK1i. Our proposal is grounded on the mechanisms of synergy we identified at optimal concentration ratios (Fig. 6). Synergy between gemcitabine and CHK1i relies chiefly on the collapse of the S-phase replication checkpoint (11). Higher concentrations of either agents induce G₂-M bypass as a

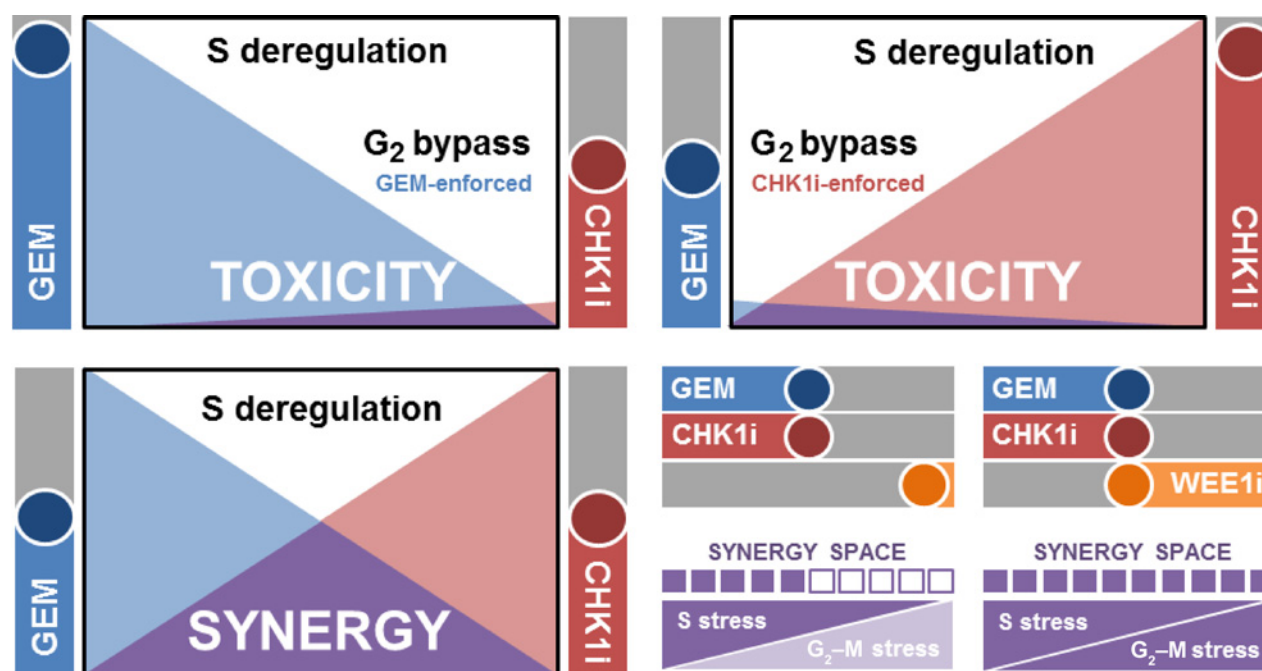


Figure 6.

Combination of gemcitabine, CHK1i, and WEE1i maximizes synergy space. Gemcitabine (GEM) and CHK1i at optimal noncytotoxic concentrations enforces synergy primarily via S-phase deregulation. Complementing this combination with WEE1i expands the synergy space of gemcitabine + CHK1i by more robust induction of G₂ bypass and mitotic stress.

secondary mechanism, but inherent to these scenarios is a greater risk of toxicity, as has been clinically observed (17, 21). Introduction of WEE1i, which we have determined to be a much superior G₂-M abrogator and mitotic stressor, expands the synergy space of the dual therapy through its complementary but independent modes of action. Because this triple regimen rationally staggers the kinase inhibitors with all agents titrated at minimally bioactive doses, it should in principle strike a balance between therapeutic activity and tolerability.

The finding that the low-dose triple regimen can indeed confer tumor suppression without acute toxicity in mice warrants further exploration. A proximate extension of this tripartite DDR model is permutations of similarly acting modalities. For instance, combinations of radiotherapy with gemcitabine and WEE1i are now entering early-phase clinical trials (45). Understanding their mechanisms at optimal dose ratios can inform decisions on doses, treatment sequences, and timing of administration. Equally, the triple regimen (gemcitabine, CHK1i, WEE1i) or its variant is testable in preclinical models tailored to pursuing specific questions on tumor initiation and progression. In particular, overcoming the evolution of treatment resistance is a key aspect of invoking drug combinations. Therefore, evaluating clinic-pathologic parameters such as long-term survival, metastasis, and disease relapse, as well as understanding the resurgence of resistance will be crucial to innovating more effective multitherapeutic strategies. The same tenet of rationalizing multipronged regimens should apply to current treatments in the clinic. Already, gemcitabine is being replaced by gemcitabine plus nab-paclitaxel and FOLFIRINOX (folinic acid, 5-fluorouracil, irinotecan, oxaliplatin) as the standard of care for pancreatic ductal adenocarcinoma, with enthusiasm in further combining them with immunotherapy

(46). There is a need to identify logical ways to induct these cocktails, and we anticipate mechanism-led preclinical studies directed at defining optimal doses and schedules to be instrumental in this endeavor. Admittedly, the scope of these translational works must also be expanded to include the physical and biological properties unique to each tumor type. For instance, the heterogeneity in drug penetration to target sites and the diversity of the tumor microenvironments are factors that could influence treatment outcome in patients. These features can only be comprehensively addressed with further development of preclinical tools and models, as well as the integration of laboratory discoveries and clinical insights.

Computational modeling on patients has shown that, although dual therapy confers clinical benefits, patients with large disease burden require triple therapy (47). There is currently no established bioinformatic tools that can predict the interaction of more than two therapeutic agents. Our approach obviates this barrier by coupling existing synergy metrics with mechanistic reasoning anchored at the single-cell level. This effort, which weight efficacy with tolerability, is a distinct component in the emerging trend of using high-resolution cellular data for the rational design of multitier therapy (48, 49). The implications of these detail-driven translational studies on the next generation of drug development could be profound.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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Development of methodology: S.-B. Koh

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Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): S.-B. Koh, C.R. Dunlop, F.M. Richards

Writing, review, and/or revision of the manuscript: S.-B. Koh, Y. Wallez, C.R. Dunlop, F.M. Richards, D.I. Jodrell

Administrative, technical, or material support (i.e., reporting or organizing data, constructing databases): S.-B. Koh, S.B. de Quirós Fernández, T.E. Bapiro
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