# **Small Molecule Modulation of the Human Chromatid Decatenation Checkpoint**

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After chromosome replication, the intertwined sister<br>
chromatids are disentangled by topoisomerases. The<br>
integrity of this process is monitored by the chromatid<br>
decale stargeting the Topoll family are impor-<br>
decaleratio **molecule modulators of the human chromatid deca- microbial agents [7, 9, 10]. One class of small molecules** tenation checkpoint identified using a cell-based,<br>chemical genetic modifier screen. Similar to 1,2,7-tri-<br>methylyzanthine (caffeine), these small molecules sun-<br>by stabilizing a covalent Topoll-DNA complex. Treatment methylyxanthine (caffeine), these small molecules sup-<br>**prace the G(2)-phase arrest caused by ICBE-193** a of mammalian cells with this class of small molecules press the G(2)-phase arrest caused by ICRF-193, a<br>small molecule inhibitor of the enzymatic activity of<br>topoisomerase II. Analysis of specific suppressors,<br>here named suppopuls for suppressor of Topoisomer-<br>ase II inhibiti

**the activation and fidelity of these checkpoints is important for understanding genomic stability and for the development of improved strategies for the treatment of** diseases involving aberrant cell proliferation or alter-

**mic stability is the proper separation of newly replicated 4Howard Hughes Medical Institute Harvard University sister chromatids prior to chromosome segregation dur-12 Oxford Street ing mitosis (M phase) [3–5]. Sister chromatids are com-Cambridge, Massachusetts 02138 posed of DNA complexed with histones and a variety of associated proteins that collectively constitute chro- <sup>5</sup> Pharmacology matin. During DNA replication (S phase), sister chroma-Harvard Medical School and tids become highly concatenated (intertwined) along Department of Cancer Biology example 3 and 2 strands of DNA** their lengths because the replicating strands of DNA **The Dana Farber Cancer Institute exist in a coil (Figure 1A) [2, 6]. In order to form individual-Boston, Massachusetts 02115 ized chromosomes in preparation for chromosome segregation, sister chromatids are decatenated starting in 6Department of Biochemistry and Molecular Biology but activity of evolution-** the G<sub>2</sub> phase of the cell cycle by the activity of evolution-**University of British Columbia arily conserved proteins known as type II topoisomer-Vancouver, British Columbia V6T 1Z3 ases (TopoII) [6–9]. This process involves the formation Canada of a double-strand DNA break and covalent linkage of the 5 phosphoryl group of each DNA strand to form a protein-tyrosyl intermediate (Figure 1A). These transformations are followed by the passing of a second intact Summary DNA strand through the break and the rejoining of the**

ase II inhibition, revealed distinct effects on cell cycle<br>progression, microtubule stability, nucleocytoplasmic<br>progression, microtubule stability, nucleocytoplasmic<br>transport of cyclin B1, and no effect on the chromatin **been referred to as the chromatid decatenation check-Introduction point [16–21].**

The maintenance of genomic stability in eukaryotic cells<br>involves an evolutionarily conserved set of signaling net-<br>works termed cell cycle checkpoints [1, 2]. These net-<br>works function to detect incompletely replicated, d **was found to be sensitive to the small molecule 1,2,7- \*Correspondence: stuart\_schreiber@harvard.edu trimethylyxanthine (caffeine) and to the structural analog**



**Figure 1. Overview of the Chromatid Decatenation Checkpoint and the Topoisomerase II-Dependent Regulation of Entry into Mitosis (A) Schematic of the role of TopoII in catalyzing the decatenation of sister chromatids and structures of small molecules known to suppress** the DNA damage-independent (caffeine [16, 22, 24], leptomycin [18]) and DNA damage-dependent (okadaic acid [23]) G<sub>2</sub> checkpoint activated **upon treatment with TopoII inhibitors.**

**(B) Chemical structures of two inhibitors of TopoII that stabilize covalent TopoII-DNA complexes.**

**(C) Chemical structure of ICRF-193 (meso-2,3-bis[2,6-dioxopiperazin-4-yl]butane), an inhibitor of the enzymatic activity of TopoII, which traps it in a closed-clamp conformation.**

**(D) Components of the network regulating the ICRF-193-induced chromatid decatenation checkpoint (modified from [18] and [20]).**

**2-aminopurine [22]. Addition of caffeine to ICRF-193- of cyclin B1 [20]. Consistent with this finding, leptomycin treated cells produces aberrant mitotic figures with ab- B (an inhibitor of nuclear export [27]) or overexpression normally elongated and intertwined chromatids con- of cyclin B1 [18] bypassed the ICRF-193-induced arrest, taining a core scaffold of chromosomal proteins [18, 22]. presumably through affecting the equilibrium of active Evidence for the importance of phosphorylation in the nuclear and cytoplasmic complexes of cyclin B1/Cdk1 chromatid decatenation checkpoint was demonstrated involved in regulating mitotic entry [28]. by the ability of the protein phosphatase 1 and 2A inhibi- Besides detecting insufficient decatenation of sister** tors (e.g., okadaic acid, Figure 1A) to overcome the G<sub>2</sub> chromatids, a DNA damage-independent G<sub>2</sub> checkpoint **arrest [23]. More recently, the signal generated by ICRF- delays the entry of cells into mitosis in response to 193 treatment was shown to be independent of the aberrant acetylation of chromatin [30, 31]. This chromaphosphatidylinositol 3 kinase family member ATM tin deacetylation checkpoint is induced upon treatment (ataxia telangiectasia mutated) and p53 [18, 21] but de- of cells with histone deacetylase (HDAC) inhibitors, such pendent upon ATR (ATM and radiation arrest deficient as trichostatin A [31–33]. This checkpoint appears to be 3-related) signaling, which is known to be inhibited by absent in a number of tumor cell lines, which may be caffeine (Figure 1D) [24–26]. In turn, ATR activation, relevant to the differential sensitivity of different tumor through a mechanism involving the tumor suppressor cell lines [31] to treatment with HDAC inhibitors. In the** BRCA1, was proposed to inhibit polo-like kinase 1 (Plk1) case of trichostatin A, the cell cycle arrest can be sup**activity, resulting in altered nucleocytoplasmic transport pressed by a set of 23 recently described small mole-**

**cules [33]. The targets of these small molecules may the other library members screened in at least one of include components of the signaling network required the three assays. After removing descriptors with no for cell cycle arrest. Complexes containing TopoII pos- variance, a total of 155 graph-theoretic, informationsess HDAC and chromatin remodeling activities [34, 35]. theoretic, or physiochemical descriptors were consid-However, the relationship between the decatenation ered for each small molecule. Since many of these mocheckpoint and the chromatin deacetylation checkpoint lecular descriptors are redundant, principal component remains poorly understood. analysis (PCA) was used to reduce the dimensionality**

**chromatid decatenation checkpoint and to clarify its composite descriptors that best represent the variance relationship to the chromatin deacetylation checkpoint, of the data [43, 44]. The resulting representation prowe took an approach similar to that used to identify vides a global, linear model of the information describing small molecule modulators of the DNA-damage-induced the structures of the small molecules. The components G2 checkpoint [36–39] and trichostatin A-induced cell of the first three eigenvectors were used to position cycle arrest [33]. This approach relies on the logic of each of the small molecules in a reduced 3-D chemical classical genetics in which the identification of suppres- space. Accordingly, this model accounted for a total of sor and enhancers of a particular phenotype has been 37% of the variance in the molecular descriptors, with a fruitful means to identify systematically components the full dimensionality spanning 147 dimensions instead of complex biochemical networks. of the original 155 (see Supplemental Figure S1 online**

A chemical genetic modifier screen was performed us-<br>
ing a cytoblot assay (described in full in [40]) to identify<br>
small molecules, yielding a mapping of biologicall<br>
small molecules that modify the cellular effects of IC cules ( $\sim$ 20  $\mu$ M) and two rounds of retesting, a total of

**regions of chemical space selected by the ICRF-193- model of the chemical space were compiled for the local modifier, TSA-modifier, and anti-mitotic screens, the region of chemical space containing only the suppres-132 biologically active small molecules were coded ac- sors of ICRF-193, and PCA was performed again. After cording to the clustering of the screening data. A set of removing descriptors with zero variance, a total of 159 molecular descriptors was then computed for these and descriptors were considered. The first three dimensions**

**To discover small molecule modulators of the human of the descriptor space to a set of linearly independent, at http://www.chembiol.com/cgi/content/full/10/12/ 1267/DC1/). Using the coding determined by the cluster- Results ing of the biological assay data, one of four colors was**

20 small molecules reproducibly increased the number<br>
of AS49 cells capable of retering mitosis in the presence<br>
of CRF-193 Grigure 2).<br>
To investigate the relationship between different cell<br>
of the ICRF-193 suppressors w **ated chemical space and the likelihood that the small Positions of ICRF-193 Suppressors on a Global molecules target the same component of the underlying Map of Chemical Space signaling network. To determine quantitatively these re-To determine global similarities/differences between the lationships, the molecular descriptors used in the global**



**Figure 2. Summary of a Chemical Genetic Modifer Screen of 9600 Small Molecules for Suppressors of ICRF-193**

**(A) Film exposure of cytoblot assay using the TG-3 mAb. A549 cells have an intact chromatid decatenation checkpoint that can be activated** by addition of ICRF-193 (7 μM) and suppressed by two of the modifiers of ICRF-193 (1 pin  $\sim$ 25-50 μM, two pins  $\sim$ 50 μM). Nocodazole (332 **nM) was added at the time of addition of suppressors to trap cells that have bypassed the chromatid decatenation checkpoint and entered mitosis.**

**(B) Summary of screening results after two rounds of retesting using the TG-3 mAb cytoblot assay.**

(C) Example of retest data for the 20 suppressors (two pin transfers;  $\sim$ 25–50  $\mu$ M) showing the average (n = 2) value of TG-3 signal relative **to the DMSO (n 10) control alone (labels correspond to structures shown in (D). Error bars correspond to one standard deviation of the mean. Increased TG-3 signal corresponds to increased bypass of the decatenation checkpoint and accumulation in mitosis. Small molecules that had a TG-3 mAb signal equal to (or greater than) the mean plus 1.5 standard deviations of DMSO treated control cells (in at least two independent retests) were considered to be biologically active.**

**(D) Chemical structures of 20 small molecule suppressors of the ICRF-193-induced chromatid decatenation checkpoint. Small molecules characterized further in this work are: 22 P11 (suptopin-1), 28 E19 (suptopin-2), and 33 N15 (suptopin-3).**

**the variance in the molecular descriptors, with the full the small molecules named suptopin-1, suptopin-2, and dimensionality spanning 21 dimensions instead of the suptopin-3 (for** *sup***pressor of** *top***oisomerase** *in***hibition) original 159 (Figure 4A). The set of ICRF-193 suppres- were selected for further cell biological analysis (Figsors is therefore less diverse than the whole library. ure 4D). The coordinates of the compounds in the 21-D reduced space were then used to construct a Euclidean distance Mitotic Spreads and Immunofluorescence matrix, which was then subject to hierarchical cluster- Although the screen we used had the capacity to identify ing. From this analysis, three distinct clusters emerged small molecules directly targeting components of the**

**in the reduced space accounted for a total of 55% of within the graph (green arrows) (Figure 4C) from which**



**Figure 3. Multidimensional Analysis of Three Cell-Based Screens of 9600 Small Molecules: Suppressors of ICRF-193, Suppressors of Trichostatin A, and Anti-mitotics.**

**(A) Summary of combined screening results.**

**(B) Hierarchical clustering of biological assay data depicting four clusters of small molecules corresponding to 20 suppressors of ICRF-193, 21 suppressors of ITSA1, 89 anti-mitotics, and 2 small molecules that scored in both the anti-mitotic and trichostatin suppressor screen. (C) Position of the four clusters of small molecules in a 3-D molecular descriptor space formed from the first three principal components (PCA) axes (1–3), which account for 37% of the total variation in the data. Axes shown run parallel to indicated PCA axes.**

**(D) Composite representations of chemical space depicting biologically active (coding) and inactive (noncoding) small molecules colored based upon the clustering of assay data from cell-based assays for suppressors of ICRF-193 (red), suppressors of trichostatin (green), and anti-mitotics (blue). Axes shown run parallel to indicated PCA axes (see Supplemental Figure S1 for an enlarged representation of the entire chemical space of the 15,120 small molecules).**

**chromatid decatenation checkpoint (e.g., as caffeine might affect directly the phosphorylation status of prodoes with ATR), the assay itself is an indirect measure teins that are recognized by the TG-3 mAb. Furthermore, based upon the level of the phosphorylated epitope although no compounds scored in both ICRF-193-modrecognized by the TG-3 mAb [36]. Consequently, small ifier and anti-mitotic screens (Figure 3B), additional phemolecules that scored positive in the primary assay notypic effects at higher concentrations/durations of**



**Figure 4. Principal Component Analysis of a 159-D Chemical Space Derived from the Molecular Descriptors of the ICRF-193 Suppressors (A) Variance accounted for by each eigenvalue/eigenvector and plot of cumulated variance of the reduced 21-D space.**

**(B) Positions of the 20 ICRF-193 suppressors according to their coordinates from the first three PCA axes (1–3). Shading (blue to red)** indicates increased weighting on  $\Psi$ 1.

**(C) Hierarchical clustering of the 20 ICRF-193 suppressors showing three main clusters (green arrows) of small molecules from which suptopin-1 to -3 were chosen for further analysis.**

**(D) Chemical structures of suptopin-1 to -3.**

**nocodazole might occur. To explore these possibilities, DMSO control, as expected due to the dominance of mitotic spreads were performed on cells treated with ICRF-193 pretreatment over the anti-mitotic effect of suptopin-1, -2, or -3 (Figures 5A and 5B). In the absence nocodazole. Upon the addition of caffeine, suptopin-2, of ICRF-193, neither caffeine nor suptopin-3 had an ef- or suptopin-3, there was a strong increase in the number fect on the percentage of mitotic cells compared to of mitotic cells compared to DMSO, whereas suptopin-1 the control solvent dimethylsulphoxide (DMSO) alone. had a weaker effect and caused some visible cell death. In contrast, both suptopin-1 and suptopin-2 caused an In contrast to the hypercondensed chromosomes seen increase in the number of mitotic cells, with the resulting upon treatment of cells with only suptopin-2, cells chromosomes hypercondensed compared to normal treated with suptopin-2 in the presence of ICRF-193 mitotic chromosomes. The fact that suptopin-1 and sup- appeared to have elongated chromosomes with paired topin-2 caused a mitotic defect in the absence of ICRF- sister chromatids. This chromosomal morphology was 193 but did not score as anti-mitotics in the original similar to the effect of caffeine and suptopin-3. assay is likely due to differences in concentration and To test for additional phenotypic effects, A549 cells duration of treatment. In the presence of ICRF-193, the treated with suptopin-1 to -3 were fixed and incubated**

**treatment or unexpected effects due to synergy with number of mitotic cells was decreased compared to the**



**cells (n total number of cells counted in two independent treat- mean side scatter (SSC), a measure of cellular granularments). All cells received an equivalent concentration of DMSO. ity or complexity, but not forward side scatter (FSC) Nocodazole (332 nM) was added at the time of addition of the (data not shown), a measure of cell size. To test whether suppressors only to ICRF-193 (7** -**M)** suppressors only to ICRF-193 (*I* LM) pretreated (5 hr) cells, and<br>cells were incubated an additional 18 hr. Suptopin-2 (60 <sub>L</sub>M, 20 g/mL) and to a lesser extent suptopin-1 (32  $\mu$ M, 20  $\mu$ g/mL) in**creased the mitotic index both in the absence and presence of treatment of cells with ICRF-193 and a checkpoint sup-ICRF-193. Caffeine (2 mM) and suptopin-3 (51 μ.Μ, 20 μ** 

**(B) Mitotic spreads of cells treated with caffeine and suptopins at and none of the chromatid decatenation checkpoint** the same concentrations as for (A). All cells received an equivalent<br>concentration of DMSO. Nocodazole (332 nM) was added at the suppressors altered the SSC levels themselves (Figure<br>time of addition of the suppressors on Mode of the pre-<br> **M** and cells incubated (6 hr) cells pre-<br> **Mode of the suptopin-2 and suptopin-3, decreased SSC in cells pre-**<br> **A** absence of ICRF-193. Suptopin-1 and suptopin-2 caused hypercon-<br> **Treated with ICRF-193** absence of ICRF-193, suptopin-1 and suptopin-2 caused hypercon**densed mitotic chromosomes unlike those of DMSO treated cells, creased granularity of cells upon treatment with ICRFwhereas mitotic cells treated with caffeine and suptopin-3 appeared 193 is due to altered chromatin conformation that occurs normal. In the presence of ICRF-193, caffeine, suptopin-2, and sup- upon activation of the chromatid decatenation checktopin-3 showed abnormally elongated chromosomes similar to point. those described in [22]. Suptopin-1-treated cells showed very few to no abnormally elongated chromosomes (none visible in experiment Effect of Suptopins shown).**

**with fluorescently labeled phalloidin to observe actin microfilaments, a fluorescently labeled anti--tubulin antibody to observe the microtubule cytoskeleton, and Hoechst 33342 to observe chromatin (data not shown). None of the compounds appeared to affect the actin cytoskeleton or cause significant alterations in chromatin. However, in interphase cells, suptopin-1 destabilized microtubules, while suptopin-2 and suptopin-3 had no observable effect. In mitotic cells, suptopin-1 caused hypercondensed chromosomes with clearly aberrant spindles, reminiscent of the effects of nocodazole, suptopin-2 caused a disorganized mitotic spindle with an apparent absence of cells in telophase, and suptopin-3 had no observable effect.**

## **Analysis of Cell Cycle and Nuclear Size Using Flow Cytometry**

**Since suptopin-2 and suptopin-3 had no effect on interphase microtubules, they were chosen for further characterization by dual-parameter flow cytometry, using the TG-3 mAb to measure the number of A549 cells in mitosis, and propidium iodide staining to quantify DNA content (Figures 6A and 6B) [45]. Similar to the mitotic spread analysis, in the absence of ICRF-193, suptopin-2 caused approximately a 6-fold increase in the number of mitotic cells (area III), whereas caffeine and suptopin-3 had no effect. In the presence of ICRF-193 and nocodazole, the number of DMSO-treated mitotic cells (area III) decreased by over half. The number** of DMSO-treated cells in G<sub>2</sub> phase, which were negative **for TG-3 mAb staining but had a 4 N amount of DNA, increased by over 3-fold. In the presence of ICRF-193 all three compounds—caffeine (15-fold), suptopin-2 (10 fold) and suptopin-3 (6-fold)—caused an increase in the** number of TG-3 mAb positive cells with a 4 N amount **of DNA, indicating a bypass of the checkpoint arrest.**

**Besides arresting A549 cells prior to mitosis, an increase in the average diameter of interphase nuclei occurs upon treatment with ICRF-193, a phenotype visible Figure 5. Effect of the Suptopins on the Mitotic Index and Chromo- by fluorescent microscopy (data not shown). Correlated somal Morphology of A549 Cells with this increase in nuclear size was an increase that (A) Mitotic index obtained from counting the frequency of mitotic was observed during flow cytometry (Figure 6C) in the M, 20 status, SSC measurements were analyzed upon pre-** pressor plus nocodazole. While FSC did not change **increased the mitotic index in the presence of ICRF-193. significantly in any of the treatments (data not shown)**

### **on Nucleocytoplasmic Transport**

**Since overexpression of cyclin B1 or treatment of cells with leptomycin B was reported to attenuate the deca-**



**Figure 6. Dual-Parameter Flow Cytometry of the Effect of Suptopin-2 and Suptopin-3 on the Cell Cycle Distribution and Side Scatter (SSC) of A549 Cells**

**(A) DNA content (y axis) and TG-3-FITC signal (x axis). Suppressors were added for a total of 18 hr. All cells received an equivalent concentration** of DMSO. Nocodazole (332 nM) was added at the time of addition of the suppressors only to ICRF-193 (7 µM) pretreated (5 hr) cells. Left panel (DMSO pretreated), suptopin-2 (60 μM, 20 μg/mL), but not caffeine (2 mM) or suptopin-3 (51 μM, 20 μg/mL), increased the mitotic index (area III). Right panel (ICRF-193 pre-treated cells), caffeine (2 mM), suptopin-2 (60 μM, 20 μg/mL), and suptopin-3 (51 μM, 20 μg/mL) all **increased the mitotic index (area III).**

(B) Quantification of cell cycle distributions in the three areas in (A). Area I, G<sub>1</sub> phase (2 N amount of DNA, low TG-3-FITC signal); Area II, G<sub>2</sub> **phase, 4 N amount of DNA, low TG-3-FITC signal); Area III, M phase, 4 N amount of DNA, high TG-3-FITC signal.**

**(C) Side scatter (SSC), a measure of cellular granularity/complexity, of cells treated with small molecules as in (A). Caffeine and to a lesser extent suptopin-3 and suptopin-2 decreased SSC in the presence of ICRF-193 and nocodazole.**

**tenation checkpoint [18], we tested whether caffeine or 193. In cells pretreated with DMSO (4 hr), none of these suptopin-1 to -3 affected the cellular localization of cyclin small molecules had a visible effect on the level of ex-B1 in A549 cells in the presence and absence of ICRF- pression of cyclin B1 (data not shown). Interestingly,**



**Figure 7. Effect of Suptopin-2 and Leptomycin B on the Intracellular Localization of Cyclin B1 and Fluorescent Properties of Suptopin-2 in A549 Cells**

**(A) Nuclear (blue, Hoechst 33342) accumulation of cyclin B1 (red; anti-cyclin B1 antibody)** in suptopin-2 (120  $\mu$ M) and leptomycin B **(37 nM) treated cells (4 hr) detected using fluorescence microscopy. Nuclei are blue when cyclin B is excluded from the nucleus and pink when it accumulates in the nucleus. (B) Percentage of cells (average n 2 treatments, 100 cells counted per treatment) with cyclin B1 predominantly nuclear detected as in (A). Error bars indicate one standard deviation from the mean.**

**(C) Intranuclear localization of intrinsic fluorescence (488 nm excitation) from suptopin-2 (151** -**M, 20 hr).**

**suptopin-2 and leptomycin B, but not caffeine, sup- molecules affect the nucleocytoplasmic transport of topin-1, or suptopin-3, caused the nuclear accumulation other proteins involved in regulating the chromatid deof cyclin B1 after a 4 hr treatment (Figure 7A) and in- catenation checkpoint. creasingly so after a 20 hr treatment (Figure 7B). However, in cells pretreated with ICRF-193 (18 hr), the in- Relationship of the Chromatid Decatenation, creased nuclear localization induced by both leptomycin Chromatin Deacetylation, and Spindle B and suptopin-2 was not observed, perhaps due to Assembly Checkpoints altered accumulation of cyclin B1 in the nucleus upon One interpretation of the absence of "hit" overlap beprolonged cell cycle arrest (data not shown). In addition, tween the ICRF-193-modifier, TSA-modifier, and antian accumulation of green fluorescent bi- or single-lobed mitotic screens is that the chromatid decatenation, globular structures within the nucleus of interphase cells chromatin deacetylation, and spindle assembly checkwas noted upon suptopin-2 treatment (Figure 7C). We points use different signaling networks in mammalian** speculate that the formation of such structures may cells. As a further test for similarities between the signal**sequester nuclear proteins that affect the chromatid de- ing networks we determined whether caffeine, an inhibicatenation checkpoint. The intrinsic fluorescence of tor of ATR that suppresses the decatenation checkpoint, suptopin-2 may prove to be useful in the identification could suppress the cell cycle effects of trichostatin A.**

**topin-2 prompted us to test whether any of the suptopins of ICRF-193. These experiments showed that caffeine affect the nucleocytoplasmic transport of other proteins has no effect on the cell cycle arrest or on the level of regulated by the Crm1-dependent transporter. To do histone H3 acetylation and -tubulin acetylation (data so, we used as a reporter U2OS cells stably transfected not shown) induced by trichostatin A and that ITSA1** with a human immunodeficiency virus (HIV) Rev-green had no effect on the cell cycle arrest induced by ICRF**fluorescent fusion protein (RevGFP) containing a nuclear 193 (Supplemental Figure S3). Collectively, these results export sequence recognized by Crm1 [46, 47]. Unlike support the notion that the chromatid decatenation leptomycin B, which caused nuclear accumulation of checkpoint functions independently from the chromatin the RevGFP reporter [47], suptopin-1 to -3 had little deacetylation checkpoint. to no effect (Supplemental Figure S2). Although these assays were performed in a different genetic back- Discussion ground than the A549 cells and with a reporter protein, this result suggests that suptopin-2 affects the localiza- The small molecule library used in this study is a com-**

**of its target(s) by biochemical means. Similarly, we tested again whether ITSA1, a suppressor The apparent control of cyclin B1 localization by sup- of trichostatin A, could suppress the cell cycle effects**

**tion of cyclin B1 independent of Crm1-mediated export. mercially available collection of compounds possessing Furthermore, the possibility remains that these small a diverse range of structural properties and chemical** **functionalities. For example, the values (low, high, aver- Of the compounds analyzed here, suptopin-3 was the age, median) for molecular weights (g/mol) are 157.21, most like caffeine in terms of its ability to bypass the 698.9, 337.73, 322.36, respectively; for calculated solu- decatenation checkpoint without having an effect on bility (cLogP) are 4.69, 9.76, 2.97, 2.99, respectively; microtubule stability or causing nuclear accumulation for the number of hydrogen bond donors are 0, 10, 1.33, of cyclin B1. Thus, we hypothesize that the cellular inter-1, respectively; and for the number of hydrogen bond actions of suptopin-3 are more specifically related to acceptors are 0, 24, 5.69, 5, respectively (see http:// the chromatid decatentation checkpoint in the check point of the check iccb.med.harvard.edu/screening/compound\_libraries/ suptopin-1 or suptopin-2.**  $R$ chembridge\_additional.html for more details). However, **without rigorous comparison to a collection of reference duction of the chromatid decatenation checkpoint with molecules or some other standard, it is difficult to draw ICRF-193 has been reported to be independent of ATM precise conclusions about the chemical diversity of the signaling and to occur in the absence of Chk1 and Chk2 library. Regardless, the dispersion of the four classes of phosphorylation [18, 20]. However, it is important to note biological activity across the global map of the chemical that although there is evolutionary conservation of the space derived from principal component analysis of the chromatid decatenation checkpoint across at least two** associated molecular descriptors indicates that no par-<br>ticular region of chamical space was favored by any tence, at least to date, depends entirely on the use of

**port based upon the Rev-GFP assay (Supplemental Fig- point suppressor leads to mutations in otherwise normal ure S2), it may specifically interact with a component cells. unique to cyclin B transport or through a mechanism In summary, the analysis of multidimensional datasets independent of Crm1. We note that in contrast to the using methods of dimensionality reduction and unsuperresults described in [18], we did not observe a lepto- vised clustering, such as hierarchical clustering and mycin B-induced increase in the number of condensed principal component analysis (for example, Figures 3 mitotic chromosomes in the presence of ICRF-193 (data and 4), provide a biologically based means of analyzing**

**ent small molecule-protein interactions, the stability of molecule modulators and other methods of chemical different organic molecules, and the abundance of cellu- perturbation, such as ribonucleic acid interference (relar proteins, some small molecules may have multiple viewed in [53]). By combining the results of multiple celland sometimes competing effects in cells at different based assays, phenotypic effects of small molecules concentrations. Such pleiotropy would explain the abil- can be mapped onto a multidimensional space derived ity of suptopin-1 and suptopin-2 to increase the mitotic from molecular descriptors analysis yielding a "funcindex in the absence of ICRF-193. Alternatively, since tional map" of chemical space (Figure 3D). 13-hydroxy-15-oxozoapatlin, a recently identified suppressor of the DNA-damage induced G<sub>2</sub> checkpoint, also Significance causes mitotic spindle defects [39], such effects might** be a common property of suppressors of the G<sub>2</sub> check-**Although there is growing interest in the use of small points that act different mechanistically from caffeine. molecules as basic research tools, there remains a**

ticular region of chemical space was favored by any tenno. at least to date, depends entirely on the use of the Higher angle and molecules with very different stuctures can have to the closed-clamp form of Topoll have addi

the diversity of small molecules and their effects on **ent genetic backgrounds of the cell lines used, duration biological systems. These global methods of analysis** may become increasingly powerful discovery tools in **Since variation exists in the relative affinities of differ- the context of rapidly growing numbers of diverse small**

**relative paucity of such molecules and only limited number of cells in mitosis was assayed using the TG-3 mAb as** approaches to enable their discovery. The chemical described in [40] using a norseradish peroxidase-conjugated anti-<br>genetic modifier screen used here to identify small<br>mouse IgM antibody and enhanced chemiluminescent reag **tenation checkpoint illustrates an approach that should was placed on top of the plate for 1–5 min and then developed in be applicable to the dissection of signaling networks** a Kodak M35A X-OMAT processor. After two rounds of retesting,<br> **involved in the requiation of other aspects of cell prolif-** small molecules that had a TG-3 mAb signa **involved in the regulation of other aspects of cell prolif-** small molecules that had a TG-3 mAb signal equal to (or greater than)<br>**eration** as well as the mechanism of action develon- the mean plus 1.5 standard deviation eration, as well as the mechanism of action, develop-<br>ment of resistance to, and metabolism of chemothera-<br>were considered to be biologically active. peutics. Using this approach in the current study, we<br>provide evidence that the chromatid decatenation<br>functions independently from the chromatin deacety-<br>was performed using the unweighted pair-group average method **lation checkpoint. Through consideration of multiple with the Tanimoto distance metric (XLStat-Pro v5.2). A structure screening datasets, these results provide an example descriptor file containing each small molecule was imported into of mapping chemical space using a combination of** QSARIS (SciVision, Inc.) for enumeration of a set of 221 graph and<br> **proportion** and molocular descriptors. The supproper contraring information theoretic and physiochemic phenotypic and molecular descriptors. The suptopins<br>provide new molecular tools for investigating the mo-<br>lecular mechanism regulating genomic stability in gen-<br>lecular mechanism regulating genomic stability in gen-<br>into M **eral and, more specifically, for illuminating whether (XLStat-Pro v5.2). In the calculation of the global chemical space, inhibiting the chromatid decatenation checkpoint small molecules with missing values (for example due to the presence of a counter ion) were removed from the analysis leaving a**<br>anents in the calculation of the local chemi-<br>anents

From Sigma. Propidium iodide, Hoechst 33342, Hoechst 33258, and<br>
Texas methyl red-conjugated anti-mouse IgG antibody were pur-<br>
chased from Molecular Probes. ITSA1 (5253409), suptopin-1 (5146479),<br>
suptopin-1 (5146479),<br> **located from Santa Cruz. Anti-TG-3 map** and  $\frac{1}{20}$  -20°C, 10 min). Cells were pelleted and all but 50–100 μl of the<br>B1 antibody was purchased from Santa Cruz. Anti-TG-3 mAb was supernatant aspirated. After resuspendi a gift from Dr. Peter Davies (Albert Einstein College of Medicine).<br>384-well pin arrays were purchased from Genetix. X-OMAT AR film<br>were expressive the material spotted from a height of 10 cm onto glass slides and allowed

L-glutamine (1% L-Gln) (DMEM<sup>+</sup>). U2OS (human osteosarcoma)containing 400 µg/mL G418.

A549 cells were seeded in 45  $\mu$ I DMEM<sup>+</sup> (4000 cells/well) in white **384-well plates (Nalge Nunc, tissue culture treated) using a liquid no emission occurring at or above 543 nm. Images of mitotic spreads dispenser (Multidrop 384, Labsystems) and allowed to attach over- were collected on a Leitz Laborlux 2 microscope with a 40 PHACO2**  $\boldsymbol{\mu}$  at 37°C with 5% CO $_{2}$ . ICRF-193 (5  $\boldsymbol{\mu}$ I) ([final] = 7  $\boldsymbol{\mu}$ **added to each well, and incubation continued for 5 hr. Since ICRF- shown) and cyclin B1 localization were collected on a Zeiss LSM510 193 is dominant to the addition of nocodazole, cells are prevented confocal scanning laser microscope using a 63 objective using** from entering mitosis and the basal signal from the mitosis-specific the accompanying software. For the RevGFP export assay, U2OS**phosphorylation-sensitive epitope recognized by the TG-3 mAb was RevGFP cells were seeded onto clear-bottom, black 384-well plates** low [36]. Library compounds (9600 molecules, Diverse E set, Chem-**Bridge) dissolved in DMSO (5 mg/ml) were then pin transferred allowed to attach and grow overnight before compound treatment.** (50–200 nL to each well, [final]  $= \sim$  20  $\mu$ M/well) using disposable **384-well pin arrays and nocodazole (332 nM final concentration), a a separate 384-well plate. Media was aspirated from cells before microtubule destabilizer, added to arrest cells that have bypassed the diluted compounds were transferred onto them. Cells were incuthe checkpoint and entered mitosis. Following 16 hr incubation, the bated with compound for 1 hr before fixation with 3.7% formalde-**

**total of 15,120 small molecules. In the calculation of the local chemi- agents. cal space of only the ICRF-193 suppressors, missing values for two small molecules (51 N16 and 37 A17) were replaced by the descriptor Experimental Procedures means. Visualization of 3-dimensional PCA models was performed by importing the coordinates of compounds after PCA into Spotfire Materials Descision Site software (v7.0). Trichostatin A, nocodazole, and ribonuclease A were purchased**

384-well pin arrays were purchased from Genetix. X-OMAT AR film<br>was purchased from Kodak Corporation. FuGENE 6 transfection<br>regents were purchased from Boehinger Mannheim. All cell culture<br>reagents were purchased from Gibc in cellular medium. Cells were fixed in glutaraldehyde (0.2%) in a<br> **Cell Culture permeabilization buffer (50 mM K-Pipes [pH 6.8], 5 mM EGTA,**<br>
A 549 cells (ATCC, human lung carcinoma) were maintained at 37°C, and MgCL, A549 cells (ATCC, human lung carcinoma) were maintained at 37<sup>°</sup>C, and MgCl<sub>2</sub>, and 0.1% Triton X-100) for 15 min, quenched (10<br>5% CO<sub>2</sub> in Dulbecco's modified Eagle medium (DMEM) supple-<br>min) in sodium borphydride (10 mg/ 5% CO<sub>2</sub> in Dulbecco's modified Eagle medium (DMEM) supple-<br>min) in sodium borohydride (10 mg/mL) in TBS/0.1% Triton X-100<br>G sodium, 100  $\mu$ g/ml streptomycin sulfate (1% P/S), and 2 mM<br>G sodium, 100  $\mu$ g/ml streptomycin **g/ml streptomycin sulfate (1% P/S), and 2 mM (FITC)-conjugated anti--tubulin antibody (DM1; 1:500), TMR-con**jugated phalloidin (2 μg/mL), and Hoechst 33342 (1 μg/mL) diluted RevGFP cells were maintained as for A549 cells and grown in DMEM<br>
containing 10% Fetal Clone (DMEM<sup>o</sup>). These cells were established<br>
by transfecting U2OS cells with pRev(1.4)-GFP+PKI NES (gift from<br>
Dr. Beric Henderson [ Dr. Benc Henderson [47]) with FuGENE 6 according to the manufac-<br>turer's directions. Stable clones were selected in complete media<br>B1 antibody (1:50) and Hoechst 33342 (1 uq/ml) dve diluted ADB B1 antibody (1:50) and Hoechst 33342 (1  $\mu$ g/ml) dye diluted ADB **g/mL G418. followed by a secondary TMR-conjugated anti-mouse IgG antibody (1:500). For detection of suptopin-2 fluorescence, A549 cells were TG-3 Cytoblot Assay illuminated between 364 and 514 nm using a longpass filter, with a** peak emission occurring at an excitation wavelength of 488 nm, and **M) was objective. Images of microtubule/actin cytoskeleton (data not** (Costar) at a density of  $\sim$ 4500 cells/well in 50  $\mu$ I DMEM<sup>c</sup>. Cells were **M/well) using disposable Compounds were diluted 1:125 in DMEM and serially diluted 1:2 in** hyde and nuclei staining with Hoechst 33258 at 2  $\mu$ g/mL, RevGFP **localization was then imaged with an inverted fluorescence micro- topoisomerase II. Biochim. Biophys. Acta** *1400***, 155–171.** scope containing a 20 $\times$  **ELWD** objective (Nikon) and Metamorph **software (Universal Imaging). the ATM and ATR kinases. Genes Dev.** *15***, 2177–2196.**

were added at the indicated concentrations for an additional 18 hr.<br>Samples were rinsed in PBS, trynsinized, fixed (30 min. 3.7% formal. **The above and all tures. Genes Dev. 14, 927–939.** Samples were rinsed in PBS, trypsinized, fixed (30 min, 3.7% formal-<br>
dehyde in TBS, room temperature; 5 min 100% methanol  $[-20^{\circ}C]$ ,<br>
washed in TBS/1% bovine serum albumin (TBS-B), and stored over-<br>
inght in TBS-B (4°C (2 mL) of TBS, resuspended in ribonuclease A (100 µL, 100 µg/ (2 mL) of TBS, resuspended in ribonuclease A (100  $\mu$ L, 100  $\mu$ g/<br>
16. Downes, C.S., Clarke, D.J., Mullinger, A.M., Gimenez-Abian, mL), and incubated 10 min (37°C). To measure the DNA content,<br>
T.F., Creighton, A.M., an propidium iodide ([PI] 400 µL, 50 µg/mL) was added and incubated propidium iodide ([PI] 400  $\mu$ L, 50  $\mu$ g/mL) was added and incubated<br>
1 hrat room temperature. Samples were analyzed using a FACScanll<br>
flow cytometer (Becton-Dickinson) at the Dana Farber Cancer Institute, exciting at

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other members of the Schreiber group for helpful discussions. S.L.S. dependent G2 checkpoint **is an Investigator at the Howard Hughes Medical Institute in the Biol. Chem.** *277***, 36832–36838. Department of Chemistry & Chemical Biology, Harvard University. 21. Kaufmann, W.K., Campbell, C.B., Simpson, D.A., Deming, P.B., We thank the National Institute for General Medical Sciences for Filatov, L., Galloway, D.A., Zhao, X.J., Creighton, A.M., and support of this research (GM38627); the National Cancer Institute Downes, C.S. (2002). Degradation of ATM-independent deca- (NCI), Merck KGaA, Merck & Co., and the Keck Foundation for sup- tenation checkpoint function in human cells is secondary to port of ICCB; and the NCI for support of the Initiative for Chemical inactivation of p53 and correlated with chromosomal destabili-Genetics. K.M.K. was supported by a Damon Runyon Cancer Re- zation. Cell Cycle** *1***, 210–229.**

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