Single-Chromosomal Gains Can Function as Metastasis Suppressors and Promoters in Colon Cancer

Highlights
- A trisomy increases metastatic behavior in colon cancer cells by inducing a partial EMT
- Chromosomal instability activates cGAS/STING signaling but suppresses invasiveness
- Whole-chromosome aneuploidies show strong correlations with prognosis across cancers

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In Brief
Tumors often display chromosome copy-number changes that have poorly understood effects on cancer physiology. Vasudevan et al. demonstrate that different aneuploidies have distinct effects on invasive behavior and that specific aneuploidies are tightly correlated with clinical outcomes. They also reveal that chromosomal instability can profoundly suppress metastatic dissemination.
Single-Chromosomal Gains Can Function as Metastasis Suppressors and Promoters in Colon Cancer

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SUMMARY

High levels of cancer aneuploidy are frequently associated with poor prognosis. To examine the relationship between aneuploidy and cancer progression, we analyzed a series of congenic cell lines that harbor single extra chromosomes. We found that across 13 different trisomic cell lines, 12 trisomies suppressed invasiveness or were largely neutral, while a single trisomy increased metastatic behavior by triggering a partial epithelial-mesenchymal transition. In contrast, we discovered that chromosomal instability activates cGAS/STING signaling but strongly suppresses invasiveness. By analyzing patient copy-number data, we demonstrate that specific aneuploidies are associated with distinct outcomes, and the acquisition of certain aneuploidies is in fact linked with a favorable prognosis. Thus, aneuploidy is not a uniform driver of malignancy, and different aneuploidies can uniquely influence tumor progression. At the same time, the gain of a single chromosome is capable of inducing a profound cell state transition, thereby linking genomic plasticity, phenotypic plasticity, and metastasis.

INTRODUCTION

Whole-chromosome aneuploidy is a nearly ubiquitous feature of human tumors, though its role in malignancy remains poorly understood (Sheltzer and Amon, 2011; Knouse et al., 2017; Chunduri and Storchová, 2019). Aneuploidy alters the dosage of hundreds or thousands of genes at once, causing proportional changes in the expression of most transcripts on an affected chromosome (Williams et al., 2008; Sheltzer et al., 2012; Stingele et al., 2012; Dürrbaum et al., 2014). These dosage imbalances have pleiotropic effects on cell physiology and can impair metabolism, protein homeostasis, and the maintenance of genomic stability (Williams et al., 2008; Sheltzer et al., 2011; Donnelly et al., 2014; Passerini et al., 2016). Despite its prevalence in cancer, under most conditions aneuploidy inhibits, rather than promotes, cell proliferation, and many single-chromosome gains can suppress tumorigenesis (Williams et al., 2008; Sheltzer et al., 2017). At the same time, the phenotypic alterations caused by aneuploidy may lead to improved fitness in certain environments (Pavelka et al., 2010; Rutledge et al., 2016), and aneuploidy can serve as a mechanism by which cells increase the dosage of growth-promoting oncogenes (Davoli et al., 2013). Thus, aneuploidy can have multifaceted and sometimes opposing roles during tumor development.

Aneuploidy can arise as a result of a cellular condition called chromosomal instability, or CIN (Holland and Cleveland, 2009). Under normal conditions, the cell’s spindle assembly checkpoint (SAC) maintains a metaphase arrest until all chromosomes are correctly aligned and under tension from the mitotic spindle (Lara-Gonzalez et al., 2012). The kinase Mps1 is the master regulator of this checkpoint, and it phosphorylates multiple proteins capable of monitoring the tension status of each chromosome (Pachis and Kops, 2018). If the SAC is compromised, cells will undergo anaphase prematurely, leading to the random missegregation of chromosomes and the development of aneuploidy.

Clinically, cancer aneuploidy has been recognized as a common indicator of poor patient prognosis (Merkel and McGuire, 1990). The overall degree of aneuploidy tends to increase during tumor progression, and aneuploid tumors from several tissues are correlated with decreased overall survival compared to tumors with near-diploid karyotypes (Sheltzer, 2013; Frankfurter et al., 1985; Friedlander et al., 1984; Choma et al., 2001; Kheir et al., 1988; Kallioniemi et al., 1987). However, most of these analyses have relied on technologies that are only capable of detecting gross deviations from the euploid state (e.g., DNA flow cytometry). Modern array-based and sequencing-based methods can be used to determine the complete set of karyotypic alterations found in a patient’s tumor, but the prognostic importance of most specific aneuploidies is not known. Additionally, the reasons underlying the close link between aneuploidy and outcome remain contested.
Figure 1. Gaining Chromosome 5 Increase Invasion, Migration, and Motility in Colon Cancer Cells

(A) Schematic diagram of the invasion assay. Cells are seeded in the upper chamber in serum-free media while the bottom chamber contains media supplemented with 10% fetal bovine serum (FBS).

(B) Quantification of the average number of cells per field that were able to cross the membrane in the invasion assay. Averages represent three independent trials in which 15–20 fields were counted.

(C) Representative images of HCT116, HCT116 Ts5 c1, and HCT116 Ts21 c1 invasion stained with crystal violet.

(D) Schematic diagram of migration assay. The setup is similar to the invasion assay except that the membrane does not have a Matrigel layer.

(E) Quantification of the average number of cells per field that were able to cross the membrane in the migration assay. Averages represent three independent trials in which 15–20 fields were counted.

(F) Representative images of HCT116, HCT116 Ts5 c1, and HCT116 Ts21 c1 migration stained with crystal violet.

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As the vast majority of cancer deaths result from metastasis, it is possible that aneuploidy plays a role in the metastatic cascade (Mehlen and Puisieux, 2006). During metastasis, cancer cells must migrate out of their local environment, intravasate into the vascular system, travel to a distant organ, and re-establish tumor growth at a secondary site (Pantel and Brakenhoff, 2004). In epithelial cancers, this process has been associated with the co-option of a developmental program called the epithelial-mesenchymal transition (EMT) (Heerboth et al., 2015). During EMT, cancer cells silence the expression of the cell-adhesion proteins typically found in epithelial tissue and activate a set of mesenchymal genes that drive cell movement and invasion. While the expression of an EMT gene signature has been correlated with poor prognosis (Jolly et al., 2017; Pastushenko and Blanpain, 2019).

Alternatively, it is possible that aneuploidy does not directly affect metastasis and instead arises in aggressive cancers as a by-product of tumor evolution. For instance, mutations in the tumor suppressor TP53 are associated both with poor prognosis and with the acquisition of aneuploidy (Thompson and Compton, 2010; López-García et al., 2017; Smith and Sheltzer, 2018), and the increased levels of aneuploidy in late-stage malignancies could simply be a consequence of the loss of p53 function. Finally, a recent study has suggested that CIN, rather than aneuploidy per se, is a key driver of cancer metastasis (Bakhoum et al., 2018). Thus, in this model, the EMT is either necessary or sufficient for tumor metastasis (Zheng et al., 2015; Fischer et al., 2015). Related programs, including a partial-EMT, in which cells fail to completely adopt a mesenchymal phenotype, have also been proposed to play a role in cancer cell dissemination (Jolly et al., 2017; Pastushenko and Blanpain, 2019).

RESULTS

Single-Chromosome Gains Can Either Promote or Suppress Metastatic Behavior

Using microcell-mediated chromosome transfer (MMCT), we generated derivatives of the human colon cancer cell line HCT116 that harbored single extra chromosomes (Stingele et al., 2012; Domingues et al., 2017). The parental cell line exhibits a near-diploid karyotype, with partial gains of chromosomes 8, 10, 16, and 17. We successfully derived clones of this line that harbored additional trisomies of one of six different chromosomes (Chr. 3, 5, 8, 13, 18, or 21). We then confirmed the presence of the extra chromosome and the absence of any secondary aneuploidies in each clone using short multiply aggregated sequence homologies (SMASH) sequencing (Figure S1A) (Wang et al., 2016).

To test how these alterations affected metastatic behavior, we applied an in vitro invasion assay, in which cells were challenged by a chemotactic gradient to invade through a basement membrane (Figure 1A). In the near-diploid parental HCT116 line, about 20 cells per field were able to successfully invade through the Matrigel layer (Figure 1B). Gaining an extra copy of Chr. 3 or Chr. 21 did not affect the rate of invasion, while gaining an extra copy of Chr. 13 or 18 significantly decreased invasion. However, in four independent clones trisomic for chromosome 5, cellular invasion significantly increased to nearly 100 cells per field (Figures 1B and 1C).

To further explore the effects of aneuploidy on metastasis-related phenotypes, we performed two additional assays in each trisomic cell line. First, we conducted a migration assay, in which cells were challenged by a chemotactic gradient to move through an 8-μm pore (Figures 1D–1F). Second, we come these limitations, we and others have developed multiple approaches to generate human cell lines with specific whole-chromosome aneuploidies (Williams et al., 2008; Stingele et al., 2012; Sheltzer et al., 2017). By altering chromosome copy number in an otherwise-identical background, we can isolate and study the effects of aneuploidy, absent any other secondary alterations. Similarly, while the causes of CIN in human tumors remain debated, CIN can be generated in vitro using small-molecule inhibitors of the SAC kinase Mps1, allowing us to isolate the specific consequences of CIN (Santaguida et al., 2010, 2015; Sheltzer et al., 2017). Using these systems, we set out to investigate how aneuploidy and CIN affect metastatic behavior in cancer.
performed a scratch-repair assay, in which we used live-cell imaging to follow the rate at which each cell line was capable of closing a cell-free gap (Figures 1G–1I). From these assays, certain patterns emerged: most trisomic lines were either neutral or displayed decreased invasiveness in each experiment. HCT116 Ts13, in particular, exhibited significantly lower rates of cellular invasion and migration and moved nearly twice as slowly as the parental line in the scratch closure assay. In contrast, four of four tested clones that harbored an extra copy of chromosome 5 displayed increased metastatic behavior in each assay. We noted some variability between independent Ts5 clones (e.g., compare Ts5 c1 and Ts5 c12, Figure 1E), but the overall patterns were consistent across this trisomy. In total, these assays demonstrate a complicated relationship between single-chromosome aneuploidies and metastatic phenotypes: specific aneuploidies may either promote, suppress, or fail to affect different metastasis-related processes, and gaining chromosome 5 in particular has a consistently strong effect on invasiveness in this colon cancer cell line.

Next, to test whether the patterns that we observed in vitro translated to changes in metastatic behavior in vivo, we injected luciferase-expressing HCT116, HCT116 Ts3, or HCT116 Ts5 cells into the spleens of nude mice and then quantified the formation of metastatic nodules on the liver and other organs (Figure 1J) (Moriyama et al., 1998). Under these conditions, the wild-type cell line was very weakly metastatic, as evidenced by the lack of luciferase signal and the scarcity of metastatic nodules that had formed by six weeks post-injection (Figures 1K–1N). HCT116 Ts3 cells were also poorly metastatic, forming a similar number of nodules as the parental line (Figures 1M and 1N). However, cells trisomic for chromosome 5 exhibited substantial metastatic activity and formed on average 15 nodules per mouse, significantly more than the parental line (p < 0.0001, Student’s t test). We conclude that the amplification of chromosome 5 in HCT116 cells increases invasive behavior both in vitro and in vivo.

We then set out to investigate the impact of aneuploidy in other human cell lines. Using MMCT, we generated TERT-transformed retinal pigment epithelial (RPE1) cell lines trisomic for Chr. 3, 5, 7, or 21 (Figure S1B; Stingele et al., 2012; Domingues et al., 2017; Durrbaum et al., 2018). Though these cells are non-cancerous, they display several hallmarks of malignant growth (activation of RAS-MAPK signaling, rapid cell division, immortality, and invasive behavior), and they provide an independent genetic background to interrogate the effects of aneuploidy (Di Nicolantonio et al., 2008). Additionally, human developmental trisomies like Down syndrome (Ts21) are commonly associated with neural tube defects and other CNS abnormalities that could potentially result from impaired migration, and we sought to uncover whether single-chromosome gains could also affect invasion and motility in non-tumor-derived cells (Hume et al., 1996; Becker et al., 1997). We therefore tested the behavior of the trisomic RPE1 cell lines in the assays described above. RPE1 Ts3, Ts5, Ts7, and Ts21 were found to suppress metastatic behavior, and Ts3 had a particularly strong inhibitory effect (Figures 2A–2E). While Ts5 increased invasiveness in HCT116 cells, in RPE1 this extra chromosome consistently blocked invasive behavior. This suggests that unique interactions between an aneuploidy and the cell’s genetic background determine whether a specific chromosome gain will have pro-metastatic or anti-metastatic consequences. Interestingly, gaining chromosome 21 strongly decreased monolayer closure in the scratch-repair assay, suggesting that some of the CNS defects observed in Down syndrome could result from Ts21-induced dysregulation of cell motility (Figures 2D and 2E).

Finally, to ensure that our results were not a consequence of the MMCT protocol used to generate these aneuploidies, we applied an alternate methodology to derive trisomic cells. We treated near-diploid DLD1 colon cancer cells with AZ3146, a small-molecule inhibitor of the SAC kinase Mps1, and isolated single-cell-derived clones. The DLD1 parental cell line harbors partial or focal gains on chromosomes 2, 11, and 16, and we successfully derived clones with additional whole-chromosome trisomies of either Chr. 4, 6, or 11 (Figure S1C). We subjected these clones to each metastasis assay, and we found that these trisomies were either neutral or suppressed invasive behavior (Figures 2F–2J). In total, our results indicate that aneuploidy can have complex effects on metastasis-related phenotypes. Single-chromosome aneuploidies can be sufficient to either promote or suppress invasiveness, depending on the cell’s genetic background and the identity of the aneuploid chromosome.

**Trisomy 5 Promotes Invasive Behavior by Causing a Partial EMT and Upregulating Matrix Metalloproteinases**

We next sought to investigate how gaining chromosome 5 in HCT116 causes an increase in invasiveness. As metastasis is commonly associated with an EMT, we assessed the expression of canonical markers of epithelial cell identity in each clone of HCT116 trisomy. We found that wild-type HCT116 cells and HCT116 derivatives trisomic for Chr. 3, 8, 13, 18, or 21 expressed high levels of the epithelial adhesion genes EpCAM, E-cadherin, and claudin 7 (Figure 3A). In contrast, EpCAM, E-cadherin, and claudin 7 were silenced in nine out of ten clones trisomic for chromosome 5, suggesting that this aneuploidy caused a loss of epithelial cell identity. To test whether Ts5 converted cells to a mesenchymal state, we next assessed the expression of the mesenchymal markers fibronectin, N-cadherin, and vimentin. We found that these genes were not expressed in any trisomic derivative of HCT116, including Ts5, indicating that this aneuploidy induces a “partial” EMT phenotype (Figure 3B).

Each trisomic cell line was derived via single-cell cloning, and HCT116 exhibits a very high mutation rate (Glaab and Tindall, 1997). While our data indicated that gaining chromosome 5 induces a profound alteration in cell identity, it remained possible that these results were an artifact of a mutation or some other alteration acquired during the MMCT process. Notably, our observation that a single Ts5 clone (“clone 11”) retained epithelial gene expression raised the possibility that our results were due to a secondary alteration that occurred during cell line derivation. We hypothesized that, if gaining chromosome 5 caused epithelial gene silencing, then eliminating that trisomy should restore epithelial gene expression. Alternately, if this phenotype was a byproduct of a mutation acquired during cloning, then the loss of epithelial cell identity should be independent of the presence of the extra chromosome. To differentiate between these possibilities, we used two approaches to generate derivatives of HCT116 Ts5 that had lost the extra chromosome (Figure 3C). First, we grew these cells as a subcutaneous xenograft in a nude
mouse, a process that we previously demonstrated selected for cells that had lost the extra chromosome (Sheltzer et al., 2017). Second, we cultured HCT116 Ts5 cells for several weeks to allow them to missegregate the trisomy, and then we performed a second round of single-cell cloning. Using SMASH sequencing, we confirmed that the post-xenograft cell line had lost the extra chromosome, and we identified three “evolved” clones that had lost most or all of the extra chromosome (Figure S1D). We found that all four Ts5-loss cell lines had restored EpCAM, E-cadherin, and claudin 7 expression (Figure 3D), and this correlated with a decrease in Matrigel invasion (Figure 3E). This indicates that the prior silencing of these epithelial genes was likely a consequence of the initial aneuploidy, while the aberrant expression profile in Ts5 clone 11 is most likely due to a secondary alteration or other clonal artifact.

In addition to the EMT, cancer cell invasion is also associated with the upregulation of matrix metalloproteinases (MMPs), a set of enzymes capable of degrading extracellular matrix components (Gialeli et al., 2011). Using qPCR, we found that HCT116 Ts5 clones upregulated several classes of MMPs, including...
Figure 3. Gaining Chromosome 5 Induces a Cell State Transition in Colon Cancer Cells

(A) Western blot analysis of epithelial marker expression in HCT116 trisomies. Alpha-tubulin was used as a loading control.

(B) Western blot analysis for fibronectin, vimentin, and N-cadherin indicates that HCT116 trisomies do not express mesenchymal genes. The sarcoma cell line U2OS was analyzed as a positive control.

(C) Schematic of two strategies to select for HCT116 Ts5 chromosome-loss revertants.

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gelatinases (MMP2), stromelysin-like proteinases (MMP11), transmembrane MMPs (MMP14 and MMP24), and GPI-type proteinases (MMP17) (Figure S2A). However, MMPs were not uniformly upregulated by Ts5, as we observed no consistent changes in MMP15, MMP21, or MMP25 expression. Non-invasive HCT116 Ts3 cells did not upregulate any of the MMPs that we tested (Figure S2A).

Finally, we sought to discover whether the loss of epithelial gene expression and the upregulation of MMPs caused the increased invasive behavior in the Ts5 cells. First, we assessed invasiveness in Ts5 clone 11 (which maintained EpCAM and E-cadherin expression), and we verified that significantly fewer cells from this clone were able to invade through a basement membrane compared to other Ts5 clones that silenced epithelial genes (Figure 3F). Next, we expressed either EpCAM or E-cadherin cDNA under a constitutive promoter in two different Ts5 clones. Western blotting revealed that these constructs led to EpCAM and E-cadherin expression levels comparable to those found in the near-diploid parental HCT116 line (Figures 3G and 3H). We then tested the effects of restoring EpCAM or E-cadherin expression on invasive behavior, and we found that the expression of either protein was sufficient to decrease Matrigel invasion to a level indistinguishable from the wild-type control (Figures 3I and 3J). Treating Ts5 cells with an MMP inhibitor also decreased Matrigel invasion, though not to wild-type levels (Figure S2B). In total, these results demonstrate that gaining an extra copy of chromosome 5 increases invasiveness in colon cancer cells by multiple mechanisms, and reestablishing epithelial gene expression is sufficient to block invasive behavior.

**Triggering Chromosomal Instability with Mps1 Inhibitors Suppresses Invasive Behavior**

Our findings indicated that specific aneuploidies can directly affect metastasis, suggesting that the frequent appearance of aneuploidy in metastatic tumors is not simply a by-product of CIN. We therefore set out to investigate the relationship between CIN, whole-chromosome aneuploidy, and invasive behavior. In order to generate CIN, we used two inhibitors of the Mps1 kinase, AZ3146 and BAY1217389, at two different concentrations. We performed live-cell imaging in HCT116 cells that expressed H2B-GFP and we verified that significantly fewer cells from this clone were able to invade through a basement membrane compared to other Ts5 clones that silenced epithelial genes (Figure 4A). Furthermore, after exposure to 1 μM AZ3146, single-cell sequencing of HCT116 revealed that 92% of cells exhibited whole-chromosome aneuploidy, compared to only 8% of cells in an untreated population (Figures 4B and 4C). Thus, Mps1 inhibitors allow us to trigger transient periods of

CIN, which generated populations of cells with different aneuploid karyotypes.

Next, we tested the effects of these Mps1 inhibitors on invasion and motility in HCT116 and DLD1 cells. To generate CIN, we cultured cells in low concentrations of AZ3146 or BAY1217389 for 24 h, then washed-out the drug and performed the assays described above (Figure 4D). Additionally, we performed a set of experiments in which we allowed AZ3146-treated cells to recover for 24 h prior to commencing the assays. Surprisingly, we found that exposure to these drugs significantly suppressed invasive behavior in nearly every condition tested (Figures 4E–4Q). For instance, treatment with either 2 μM AZ3146 or 6 nM BAY1217389 decreased Matrigel invasion in HCT116 cells by more than 80% (Figures 4E and 4F) and to a lesser degree in DLD1 cells (Figure 4N). These drugs also decreased pore migration and slowed the closure of a monolayer scratch (Figures 4G–4I and 4O–4Q). Finally, HCT116 cells treated with 2 μM AZ3146 formed significantly fewer metastatic nodules compared to the parental line when injected into the spleens of nude mice (Figures 4J–4M). These results suggest that Mps1 inhibitors are capable of suppressing invasive behavior.

To confirm that these results were not specific for colon cancer cells, we performed additional sets of Matrigel invasion assays following Mps1 inhibitor treatment in the near-diploid Cal51 breast cancer cell line, the highly-aneuploid A375 melanoma cell line, and in RPE1-hTert cells. As we observed with HCT116 and DLD1 cells, blocking Mps1 significantly reduced Matrigel invasion, particularly at the higher concentration of either AZ3146 or BAY1217389 (Figures 5A and 5B). Thus, in a variety of different cancer types and genetic backgrounds, exposure to a drug that increases CIN suppresses rather than enhances invasive behavior.

Many small molecules can exhibit promiscuous activity against other kinases, leading us to consider the possibility that an off-target effect of these drugs suppressed invasion (Giuliano et al., 2018; Klaeger et al., 2017; Lin et al., 2019). To test whether this phenotype represented an on-target effect of Mps1 inhibition, we used CRISPR-mediated homology-directed repair (HDR) to introduce two mutations into the MPS1 gene (C604W and S611G; Figure S3A; Table S1A) that have been reported to block the binding of AZ3146 to Mps1 (Gurden et al., 2015). Live-cell imaging verified that HCT116 cells harboring these mutations displayed minimal CIN when cultured in 2 μM AZ3146 (Figures S3B and S3C). Correspondingly, while 2 μM AZ3146 resulted in a significant decrease in Matrigel invasion in wild-type HCT116, treatment of cells harboring either Mps1C604W or Mps1S611G with AZ3146 had no effect on invasion (Figures S3D and S3E). These results demonstrate that AZ3146
Figure 4. Chromosomal Instability Is Insufficient to Drive Metastasis

(A) Quantification of mitotic error frequency after Mps1 inhibitor treatment. A mitosis with “minor errors” exhibited a single-lagging chromosome, anaphase bridge, or micronucleus, while a mitosis with “major errors” displayed more than one of these phenomena.

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suppresses invasion specifically through its inhibition of the SAC kinase Mps1.

We next considered the possibility that our experimental approach (in which cells were examined after Mps1i washout) blocked the pro-invasive effects of CIN, perhaps by restoring normal chromosome segregation fidelity. As aneuploidy is sufficient to increase CIN (Sheltzer et al., 2011; Passerini et al., 2016) and as micronuclei that result from prior missegregation events are liable segregate incorrectly during subsequent mitoses (Soto et al., 2018), we anticipated that cells would still display instability following drug washout. Indeed, live-cell imaging 24 h after Mps1i washout demonstrated that cells continued to exhibit a significant increase in mitotic errors compared to untreated control cells (Figure S3F). To further verify that Mps1 inhibitors decreased cellular invasion, we repeated the Matrigel invasion assays in the presence of low doses of AZ3146 or BAY1217389, without drug washout. Under these conditions, drug treatment caused a significant decrease in Matrigel invasion, verifying that increased invasiveness is not an obligate consequence of ongoing CIN (Figure S3G).

As MPS1 is an essential gene (Maciejowski et al., 2010), we next sought to test whether cellular toxicity caused by transient exposure to Mps1 inhibitors could explain the decrease in invasion that we observed. First, we assessed PARP cleavage and caspase-3 cleavage (markers of apoptosis) and β-galactosidase staining (a marker of senescence) in cells treated with Mps1 inhibitors. 3 nM BAY1217389, 6 nM BAY1217389, and 1 μM AZ3146 treatment had no effect on apoptosis, while only the highest concentration of AZ3146 (2 μM) caused a slight increase in the appearance of cleaved PARP and cleaved caspase-3 (Figure S4A). Similarly, fewer than 1% of Mps1i-treated cells stained positive for senescence-associated β-galactosidase (Figures S4B and S4C). Thus, under multiple conditions, Mps1 inhibitors suppress invasion without significantly increasing apoptosis or senescence. As an alternate approach to investigate this question, we exposed cells to AZ3146 treatment for 24 h and then cultured them in drug-free media for either 7 or 14 days. We reasoned that these conditions would allow cells to recover from any acute toxicity caused by the Mps1 inhibitor. Instead, we hypothesized that aneuploidy and cytoplasmic DNA that

Figure 5. The Effects of Mps1 Inhibitor Treatment on Invasive Behavior

(A, C, and E) Quantification of the average number of cells per field that were able to cross the membrane in the invasion assay in (A) A375, (C) Cal51, and (E) RPE1. Averages represent two independent trials in which 15–20 fields were counted.

(B, D, and F) Representative images of invasion in the indicated Mps1i-treated cell lines. Bars represent mean ± SEM. Unpaired t test *p < 0.05; **p < 0.005; ***p < 0.0005. Scale bar, 50 μm.

Bars represent mean ± SEM. Unpaired t test *p < 0.05; **p < 0.005; ***p < 0.0005. Scale bar, 50 μm.
arise from Mps1-induced CIN may persist (though the least-fit aneuploid cells could be out-competed by other cells over time). Indeed, treatment with an Mps1 inhibitor caused an increase in the number of cells that harbored micronuclei, and this increase remained apparent up to 14 days after the drug had been washed out (Figures S4D and S4E; and see below). We found that HCT116 cells displayed lower levels of Matrigel invasion, even 7 or 14 days after exposure to AZ3146 (Figure S4F). Under these same conditions, cells treated with a low dose of either Mps1 inhibitor proliferated at an indistinguishable rate compared to wild-type cells (Figure S4G). In total, these experiments indicate that cIN is capable of suppressing cellular invasion, and that these results are unlikely to reflect an off-target effect or acute toxicity caused by Mps1 inhibitors.

cGAS/STING Activity Is Insufficient to Trigger an EMT or Explain the Invasive Behavior of HCT116 Ts5 Cells

CIN has been reported to promote invasive behavior by activating non-canonical NFκB signaling via the cytosolic DNA sensor cGAS and its adaptor STING (Bakhoum et al., 2018). As we found that Mps1 inhibitors are capable of suppressing cellular invasion, we sought to test whether they similarly affected these pathways. First, we confirmed that both AZ3146 and BAY1217389 lead to a significant, 6- to 10-fold increase in micronuclei in both HCT116 and DLD1 cells 24 h after drug washout (Figures 6A and 6B). While the expression of STING was normally low in these cells, treatment with AZ3146 or BAY1217389 caused an increase in the appearance of cells that were strongly STING positive (Figures 6C and 6D). Mps1 inhibitor treatment also caused an increase of 5- to 11-fold in the translocation of the NFκB transcription factor RelB into the nucleus (Figures 6E and 6F). We also confirmed via qPCR that multiple NFκB targets were upregulated (Figure 6G). Thus, these results demonstrate that the cytosolic DNA sensing pathway is functional in these cells, and that Mps1 inhibition is sufficient to activate a RelB-related transcriptional response. However, in a variety of different cancer cell lines, neither Mps1 inhibitor was sufficient to drive cellular invasion (Figures 4 and 5) or to alter the expression of the canonical EMT proteins EpCAM, E-cadherin, N-cadherin, or vimentin (Figure S5). We conclude that an EMT is not an obligate consequence of cIN, and instead the previously reported association between CIN, EMT, and metastasis may be limited to certain cancer types or specific ways of generating CIN.

Despite these results, cGAS/STING signaling undoubtedly has a profound effect on multiple cancer-related phenotypes. Additionally, STING is encoded on chromosome 5, and multiple inflammatory genes are upregulated in trisomic cell lines (Dürbaum et al., 2014; Viganò et al., 2018), leading us to investigate whether cGAS/STING contributed to the metastatic behavior of the HCT116 Ts5 cells. To accomplish this, we stably expressed a dCAS9-KRAB CRISPRi vector in two independent Ts5 clones, which allowed us to trigger the specific downregulation of a gene of interest (Horlbeck et al., 2016). We then transduced each line with two guide RNAs (gRNAs) that targeted either cGAS or STING, and we verified via western blot that these guides strongly suppressed cGAS and STING expression, respectively (Figure 6H). However, we found that silencing cGAS or STING had no effect on either EpCAM expression or invasive behavior in Ts5 (Figures 6I–6K). Thus, whole-chromosome aneuploidy is capable of driving invasion in a cGAS/STING-independent manner.

We hypothesized that the over-expression of a gene or genes on chromosome 5 led to the silencing of epithelial genes and the increased invasive behavior in HCT116 Ts5. We selected 24 candidate genes encoded on chromosome 5 that had known roles in development, signal transduction, or gene regulation, and we decreased their expression with CRISPRi in two independent Ts5 clones. However, none of these knockdowns was sufficient to restore EpCAM expression (Figure S6; Tables S1B and S2). The metastatic behavior of HCT116 Ts5 may therefore result from the coordinated over-expression of several chromosome 5 genes, or from a gene not included among this panel.

Different Chromosomal Aneuploidies Are Associated with Distinct Clinical Outcomes

Our results indicate that whole-chromosome aneuploidies have the capability to either promote or inhibit invasive behavior. Additionally, our previous research has demonstrated that certain aneuploidies can directly suppress transformation (Sheltzer et al., 2017). Yet, clinically, aneuploidy is widely reported to correlate with cancer progression and aggressive disease (Merkel and McGuire, 1990). How can these disparate observations be reconciled?

To address this, we performed Cox proportional hazards survival modeling on 10,686 patients with 27 different types of cancer from The Cancer Genome Atlas (TCGA; other abbreviations are defined in Table S3A). In this approach, one or more clinical variables are regressed against patient outcome, allowing us to identify features that correlate with prognosis. Here, we report the Z scores from these models, which capture both the directionality and significance of a particular clinical association. If a particular chromosome gain is significantly associated with patient death, then a Z score >1.96 corresponds to a p value <0.05 for that relationship. In contrast, a Z score less than −1.96 indicates that a chromosome gain event is significantly associated with patient survival (or that a chromosome-loss event is associated with death). 10,133 of these patients’ tumors have been analyzed by SNP array, allowing their tumors’ bulk karyotype to be determined (Taylor et al., 2018). Tumors in the TCGA lack comprehensive annotation of metastases, so we instead used either “overall survival” or “progression-free survival” as clinical endpoints for this analysis (Table S3A, and see the STAR Methods).

First, we regressed each tumor’s total aneuploidy burden against patient outcome. We determined that highly aneuploid cancers were associated with a significantly worse prognosis than tumors with low aneuploidy in nine of 27 different patient cohorts (Figures 7A and 7B). In the remaining 18 cancer types, no significant association was detected. We did not find any cohorts in which high aneuploidy portended a favorable outcome (Figure 7B). Some cancer types in which aneuploidy has previously been reported to be a prognostic biomarker were not found to exhibit a significant correlation in this analysis (e.g., lung cancer; Choma et al., 2001); this may be because of differences in the technology used to measure aneuploidy, differences between patient populations, or because of the limited number of patients included in some cohorts. The acquisition of aneuploidy
was linked to the loss of p53 function, as p53-mutant tumors tended to have higher levels of aneuploidy than p53-wild-type tumors (Figure S7A). However, the overlap between p53 mutations and highly aneuploid tumors was unable to fully account for the prognostic significance of aneuploidy: in multivariate Cox regression models that included both p53 status and aneuploidy, high aneuploidy burden was still significantly correlated with poor survival in six different cohorts (Figures S7B and S7C).

We next set out to determine how specific aneuploidies influenced patient prognosis. We constructed Cox models to interrogate the relationship between the alteration status of each chromosome arm (“loss,” “neutral,” or “gain”) and patient outcome.
across the 27 cohorts. This analysis revealed 160 arm-length aneuploidies that were associated with survival time (Figure 7C; Table S3B). In every cancer type, patients were subdivided into two groups: low aneuploidy (≤ 20th percentile aneuploidy score) and high aneuploidy (≥ 80th percentile aneuploidy score). Kaplan-Meier curves are shown for three cancer types. (B) Z scores from Cox-proportional hazards analysis based on the aneuploidy scores, as described above, are displayed. Note that Z > 1.96 indicates a significant association between higher aneuploidy and death, while Z < –1.96 indicates a significant association between higher aneuploidy and survival. (C) A heatmap comparing chromosome arm copy number versus patient outcome across 10,133 patients with 27 different types of cancer. The color bar indicates the Z score from the Cox analysis. For visualization purposes, Z scores were capped at 4 and –4. (D) A heatmap comparing dichotomized chromosome arm copy number versus patient outcome across 10,133 patients with 27 different types of cancer. In this analysis, survival was compared between patients in which a given arm was gained and patients in which a given arm was either lost or was copy neutral. The color bar indicates the Z score from the Cox analysis. For visualization purposes, Z scores were capped at 4 and –4. (E) Representative Kaplan-Meier curves of two instances in which chromosome gains are associated with improved patient prognosis. (F) A heatmap comparing dichotomized chromosome arm copy number versus patient outcome across 10,133 patients with 27 different types of cancer. In this analysis, survival was compared between patients in which a given arm was lost and patients in which a given arm was either gained or was copy neutral. The color bar indicates the Z score from the Cox analysis. For visualization purposes, Z scores were capped at 4 and –4. (G) Scatter plots comparing the average chromosome copy number versus the Z score obtained from Cox analysis for three different cancer types. (I) A bar graph displaying the average chromosome copy number across all 27 cancer types, binned based on the Z score from the Cox model.

Figure 7. A Pan-Cancer Analysis of Chromosome Copy-Number Changes Associated with Patient Survival
(A) The total number of aneuploid chromosomes in each tumor was calculated according to (Taylor et al., 2018). In each cancer type, patients were subdivided into two groups: low aneuploidy (≤ 20th percentile aneuploidy score) and high aneuploidy (≥ 80th percentile aneuploidy score). Kaplan-Meier curves are shown for three cancer types.

Throughout the text, there are references to cell-free DNA (cfDNA) as a source of biomarkers and various cancer types, such as breast, lung, and colorectal cancers. The analysis involves the use of bioinformatics tools, such as the Cancer Genome Atlas (TCGA) and the cBioPortal, to identify genetic alterations associated with cancer progression.

As our in cellulo and in vivo work had primarily focused on colon cancer, we next examined the clinical correlates of aneuploidy in this patient cohort in particular. We found that total aneuploidy tended to increase with colon cancer stage: stage types. Thus, there is no single “pro-metastatic” karyotype, and the effects of specific aneuploidies on cancer cell dissemination likely depend on a tumor’s genetic and epigenetic background.
IV tumors harbored a median of 16 arm-length alterations, nearly twice as many as the median stage I or II tumor (Figure S7D). Interestingly, colon cancer aneuploidy was not significantly associated with a worse response to therapy (Figure S7E). However, tumors that expressed high levels of an EMT-related gene signature displayed more aneuploidy than tumors that lacked this signature, suggesting a potential in vivo link between aneuploidy and cell state transitions (Figure S7F). Colon tumors commonly displayed a number of recurrent aneuploidies, including the gain of Chr. 20q, Chr. 7, and Chr. 13, and the loss of Chr. 18 and Chr. 17 (Figures S7G and S7H). We did not detect a significant correlation between the gain of chromosome 5 and patient outcome; as RPE1 Ts5 cells also did not display a significant increase in invasiveness, this is consistent with our hypothesis that aneuploidies interact with a cell’s genetic or epigenetic background to drive metastasis (Figure S7I). In fact, no whole-chromosome events were individually associated with colon cancer outcome, and only three arm-length alterations were (8p, 10p, and 17q; Figures S7I and S7J). Thus, as we observed in our cell line models, a majority of individual aneuploidies are not strong drivers of tumor metastasis.

Across all 27 cancer cohorts, we observed a positive relationship between chromosome copy number and survival (Z > 1.96) for 66 arm-length aneuploidies, while we found a negative relation (Z < −1.96) for 94 aneuploidies. These results could be consistent with two different models: a positive Z score could indicate that tumors that harbor an amplification of that chromosome have a worse prognosis than tumors that lack that amplification, or it could indicate that tumors that have lost a copy of that chromosome have a better prognosis than tumors that are copy neutral. Similarly, negative Z scores could indicate that chromosome losses accelerate patient death or that chromosome gains protect against it. To differentiate between these models, we reanalyzed the TCGA survival and karyotype data, binning “loss” and “neutral” calls together, or binning “gain” and “neutral” calls together (Figures 7D–7G; Tables S3C and S3D). This analysis revealed that in ~87% of cases, the aneuploidy event (either gain or loss) was associated with worse patient outcome. However, strikingly, in ~13% of cases, the gain or loss of a chromosome arm correlated with improved patient survival. For instance, glioblastomas that are copy neutral or have lost Chr. 10p have a median survival time of 422 days, while glioblastomas that harbor a gain of Chr. 10p have a median survival time of 1,987 days (Figure 7E). Thus, while most prognostic aneuploidies correlate with poor outcomes, aneuploidy of many different individual chromosomes can be associated with decreased tumor aggressiveness.

If aneuploidy is capable of either promoting or suppressing aggressive behavior, then why is bulk tumor aneuploidy a common hallmark of poor prognosis? To address this, we compared the arm-level Z scores obtained from our survival analysis with the overall frequency of each aneuploidy event. We found that, in general, chromosome arms that were associated with poor prognosis when amplified were significantly more likely to be gained than lost in primary tumors. Similarly, arms whose deletions correlated with dismal survival were more commonly lost than gained (Figures 7H and 7I). For instance, in head and neck squamous cell carcinoma, 8q amplifications are associated with poor prognosis, and this chromosome arm was gained in 54% of patients and lost in <1% of patients. In contrast, 3p deletions are a biomarker of poor prognosis, and this chromosome arm was lost in 61% of patients and gained in <1% (Figure 7H). This suggests that aneuploidies capable of suppressing aggressive behavior are selected against during tumor development, leading to an enrichment of metastasis-promoting aneuploidies, even though they represent only a fraction of all possible copy-number changes.

**DISCUSSION**

Stoichiometric imbalances in endogenous proteins caused by aneuploidy interfere with multiple cellular functions (Sheltzer and Amon, 2011). It is therefore conceivable that cell motility, matrix degradation, and the other processes necessary for invasive behavior might also be perturbed by aneuploidy-induced dosage imbalances. Thus, while aneuploidy is a hallmark of cancer, aneuploidy in itself might potentiate suppression of metastatic behavior. To investigate the relationship between aneuploidy and metastasis, we have generated and analyzed a series of modified cell lines that differ from each other by a single chromosome. Using this system, we discovered that single-chromosome gains can have multifaceted effects on several proxies for metastatic ability. Across 13 different trisomies, 12 inhibited or had a minor effect in assays designed to test invasiveness and cell motility.

HCT116 colon cancer cells harboring an extra copy of chromosome 5 behaved differently from every other trisomy studied and displayed a clear increase in metastatic capacity. We traced this phenotype to an aneuploidy-induced partial EMT, in which the specific amplification of chromosome 5 causes the silencing of epithelial cell-adhesion genes. This effect is independent of cGAS/STING signaling, and we posit that the over-expression of one or more genes on chromosome 5 cooperate to induce this phenotypic switch. Aneuploidies-induced cell state transitions may be an important and understudied cause of the strong correlation between aneuploidy and death from cancer. Additionally, recent evidence has underscored the importance of phenotypic plasticity in the metastatic cascade and has suggested that an EMT is neither a necessary nor an irreversible step in cancer cell dissemination (Chen et al., 2019; Padmanaban et al., 2019; Zheng et al., 2015; Fischer et al., 2015; Liu et al., 2019). Instead, cancer cells can lose certain epithelial characteristics to promote migration out of the primary tumor site and into circulation, but they can re-establish them (through a mesenchymal-epithelial transition) in order to colonize a distant location (Dongre and Weinberg, 2019). Aneuploidy, too, can be reversible: gaining and losing chromosomes through CIN could contribute to this plasticity and allow cancer cells to switch between different phenotypic states.

CIN has previously been reported to directly influence metastasis by triggering a cGAS/STING-dependent EMT and increase in cell motility (Bakhoum et al., 2018). By generating CIN with Mps1 inhibitors, we have verified the link between CIN and cGAS/STING activation, but we find that these inhibitors suppress rather than promote metastatic behavior. These results demonstrate that increased invasiveness is not an obligate consequence of either CIN or cGAS/STING signaling and may instead depend on the mechanism by which CIN arises. As
CIN causes the acquisition of aneuploidy, we believe that, as with the single-chromosome trisomic cell lines that we have characterized, CIN-induced proteome deregulation may interfere with the processes necessary to achieve a metastasis-competent state. Low levels of CIN, or CIN that allows the development of rare, metastasis-promoting aneuploidies, may be necessary to observe the conditions under which CIN enhances metastasis. Additionally, as several Mps1 inhibitors have entered clinical trials in human patients (Xie et al., 2017), our results further suggest that one potential benefit of Mps1 inhibition as a therapeutic strategy may be an overall decrease in metastatic dissemination.

Interestingly, while Ts5 enhanced metastasis in HCT116 cells, this same chromosome mildly suppressed metastatic behavior when added to retinal-pigment epithelial cells. We speculate that the gain of chromosome 5 has a specific effect on HCT116 cells based on their genetic and epigenetic background, and our pan-cancer analysis of aneuploidy-associated patient mortality supports this hypothesis. Many aneuploidies exhibit distinct clinical associations in different cancer types: for instance, Chr. 10p losses are associated with poor survival in gliomas and kidney cancers, but Chr. 10p amplifications are associated with poor survival in leukemia and hepatocellular carcinoma (Table S3B). We further observed that, among amplified chromosomes, Chr. 8q gains showed the strongest correlation with patient death, but this association was only present in six of 27 cancer types (Table S3B). We find no evidence to support the hypothesis that specific aneuploidies are universal metastasis promoters (Duesberg et al., 2006). Instead, the consequences of each aneuploidy are closely tied to the original tumor type. Yet, chromosome copy-number changes can provide raw fodder for tumor evolution. While a subset of aneuploidies do in fact correlate with decreased aggressiveness in patient samples, these aneuploidies are rarely observed. The copy-number changes that instead drive malignant behavior are selected through evolution, and thus “deadly” aneuploidies appear to be more common than they actually are. These results reconcile the apparent contradiction between the behavior of artificially constructed aneuploid cells, which almost always display a pronounced growth defect and the aggressive behavior of aneuploid human tumors (Stingele et al., 2012; Sheltzer et al., 2017).
Chromosomal instability drives metastasis through a cytosolic DNA response. Nature 553, 467–472.


Maciejowski, J., George, K.A., Terret, M.E., Zhang, C., Shokat, K.M., and Jallepalli, P.V. (2010). Mps1 directs the assembly of Cdc20 inhibitory...


# STAR METHODS

## KEY RESOURCES TABLE

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LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Jason Sheltzer (sheltzer@cshl.edu).

Materials Availability Statement

Trisomic cell lines (HCT116 and RPE1) generated in this study were obtained and are available from Dr. Zuzana Storchova. DLD1 trisomic cell lines were generated by the Sheltzer lab for this paper and are available from Dr. Sheltzer.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cell Lines

The identity of each human cell line was verified by STR profiling (University of Arizona Genetics Core, Tucson, AZ). The karyotype of every aneuploid cell line used in this manuscript was verified with the SMASH technique (Wang et al., 2016) or had been previously analyzed with a similar low-pass whole-genome sequencing method (Sheltzer et al., 2017). The names of the clones and their karyotypes are summarized in Table S4. HCT116 (sex: male), DLD1 (sex: male), RPE1 (sex: female), Cal51 (sex: female), SW480 (sex: male), and A375 (sex: female) were grown in DMEM supplemented with 10% FBS, 2 mM glutamine, and 100 U/mL penicillin and streptomycin. T47D (sex: female) cells were grown in RPMI supplemented with 10% FBS, 6.94 μg/ml insulin (Thermo Fisher, Waltham, MA; BN00226), 2 mM glutamine, and 100 U/ml penicillin and streptomycin. A549 (sex: male) cells were grown in Ham F12 supplemented with 10% FBS, 100 U/mL penicillin and streptomycin and 2 mM glucose. All cell lines were grown in a humidified environment at 37°C and 5% CO2.

In Vivo

Nude mice (NU/J (Foxn1nu) female, 6-week old; 20-30g; JAX stock #002019) were obtained from the Jackson Laboratory. The Cold Spring Harbor Laboratory IACUC reviewed and approved all procedures. The mice were housed four per cage (369 x 156 x 132 mm; 1145T, Tecniplast, UK) and maintained on a 12 hours light-dark cycle, with temperature and relative humidity ranging between 20–25 degree Celsius and 70–80%, respectively. Food and water were available ad libitum.

METHOD DETAILS

Sample Sizes and Statistical Methodology

For the Matrigel invasion assays (Figures 1B, 2A, 2F, 3E, 4N, 5A, 5C, 5E, 6J, S2B, S3D, S3G, and S4F) we analyzed 15-20 independent fields of view, collected from 2 to 4 independent biological replicates of the experiment. For the migration assays (Figures 1E, 2C, 2H, 4G, and 4O) we analyzed 15-20 independent fields of view, collected from 2 to 4 independent biological replicates of the experiment. For the scratch assays (Figures 1G, 2D, 2I, 4H, and 4P) we analyzed 8-10 fields along the scratch, collected from 2-4 independent biological replicates of the experiment. For the spleen-liver metastasis assays (Figures 1N and 3M), 16 NU/J mice were injected with HCT116 luciferase-expressing cells, 10 mice were injected with HCT116 Ts5 c1 luciferase-expressing cells, 8 mice were injected with HCT116 Ts3 luciferase-expressing cells, and 8 mice were injected with HCT116 luciferase-expressing cells treated with 2 mM AZ3146. For Mps1i-induced micronuclei formation (Figures 6A and S4D), we analyzed a minimum of 500 cells per trial collected from 3 independent biological replicates of the experiment. For the STING and Rel-B immunofluorescence assays (Figures 6C and 6E), we analyzed a minimum of 500 cells per trial collected from 3 independent biological replicates of the experiment. For the qPCR assays (Figures 6G, S2A, and S6B), we performed 3 technical replicates collected from 3 independent biological replicates of the experiment. For live cell mitotic imaging (Figures 4A, S3B, and S3F), we analyzed a minimum of 100 cells from 1-2 independent biological replicates of the experiment.

For the invasion, migration, scratch, immunofluorescence, and live-cell imaging experiments, the samples were analyzed while blinded to cell identity. No outliers were excluded from analysis. Bar graphs display sample means +/- SEM. Unpaired t-tests were used to measure statistical significance.

Cell Lines and Tissue Culture Conditions

The identity of each human cell line was verified by STR profiling (University of Arizona Genetics Core, Tucson, AZ). The karyotype of every aneuploid cell line used in this manuscript was verified with the SMASH technique (Wang et al., 2016) or had been previously analyzed with a similar low-pass whole-genome sequencing method (Sheltzer et al., 2017). The names of the clones and their karyotypes are summarized in Table S4. HCT116, DLD1, RPE1, Cal51, SW480, and A375 were grown in DMEM supplemented with 10% FBS, 2 mM glutamine, and 100 U/mL penicillin and streptomycin. T47D cells were grown in RPMI supplemented with 10% FBS, 6.94 μg/ml insulin (Thermo Fisher, Waltham, MA; BN00226), 2 mM glutamine, and 100 U/ml penicillin and streptomycin. A549 cells were grown in Ham F12 supplemented with 10% FBS, 100 U/mL penicillin and streptomycin and 2 mM glucose. All cell lines were grown in a humidified environment at 37°C and 5% CO2.
Karyotype Analysis with SMASH
Cells were trypsinized and resuspended in PBS, centrifuged at 1000 rpm for 5 minutes, and then the pellets were collected. Total cellular genomic DNA was isolated using the Qiagen QIAamp kit (Cat. No. 51036). SMASH karyotyping was performed as described (Wang et al., 2016). In brief, total cell genomic DNA was enzymatically fragmented to a mean size of ~40bp and joined to create chimeric fragments of DNA suitable for creating NGS libraries (300-700bp). The fragment size selection was done with Agencourt AMPure XP beads (Beckman Coulter, Cat. No. A63881). Illumina-compatible NEBNext Multiplex Dual Index Primer Pairs and adapters (New England Biolabs, Cat. No. E6440S) were ligated to the selected chimeric DNA fragments. These barcoded DNA fragments were then sequenced using an Illumina MiSeq. Bin boundaries were determined empirically by apportioning valid SMASH mappings from 56 CHD trios sequenced at high depth to a fixed number of bins such that the minimum bin count is maximized and the remaining bins are populated as evenly as possible (Andrews et al., 2016; Andrews, 2017). Approximately 2% of the resulting bins showed significant variation in count across the CHD population and were thus excluded from further analysis and display using both automated means and human review. The generated reads were demultiplexed and mapped using custom scripts, and plots were generated with G-Graph (Andrews, 2017).

Single-Cell Sequencing
This protocol was adapted from (Baslan et al., 2012). In brief, HCT116 cells were plated into 6 well cell culture plates and 24hr later the cells were treated with 1 μM AZ3146. The next day, the media was changed. 24hr later, the cells were trypsinized and single-cell sorted into 96 well PCR plates (ThermoFisher Scientific, Cat. No. AB0731) containing cell lysis buffer (0.1% SDS, 2% Triton) and incubated at 65 °C for 1 hour. Genomic DNA was enzymatically digested with NlaIII (NEB, Cat. No. R0125). Both ends of fragments were tagged by using oligonucleotides that contain cell-barcode, universal primer and several random nucleotides (varential tags) through ligation and extension reactions. Cell barcodes allow us to multiplex samples, and the tags allow unique counting of initial DNA fragments. After amplification by universal primer, fragment size selection was performed with Agencourt AMPure XP beads (Beckman Coulter, A63881) to obtain fragments appropriate for sequencing. Barcoded sequencing adapters were ligated to the DNA fragments which allows us to further multiplex samples for sequencing. Next generation sequencing was done on an Illumina MiSeq. Reads were demultiplexed using customized scripts and karyotype plots were generated with G-Graph (Andrews, 2017).

Invasion and Migration Assays
For transwell migration assays, cells were plated in the upper chamber of the transwell insert (Corning Cat. No. 3464, 24-well insert, pore size: 8 μm). For DLD1 and HCT116 cells, 3 × 10^4 cells were seeded, while for RPE1 cells 1 × 10^6 cells were seeded. For transwell invasion assays, cells were plated in the top chamber with Matrigel-coated membranes (Corning Cat. No. 354480, 24-well insert, pore size: 8 μm). For DLD1, 1 × 10^5 cells were seeded, for HCT116, 2 × 10^5 cells were seeded, and for RPE1, 1.5 × 10^6 cells were seeded. Cells in the upper chamber were plated in serum-free DMEM, while media with 10% FBS was added to the lower chamber. The plates were incubated for 24 hours at 37 °C, and then the cells on the upper surface were removed using cotton swabs. The membranes were then fixed in methanol and stained with crystal violet dye. The membranes were cut out and mounted on to slides. They were imaged at 40x (15-20 images) and counted to obtain the average number of cells per field that migrated or invaded. Two to three chambers were used per cell line and/or condition. For experiments using Mps1 inhibitors, the cells were either treated for 24 hours with the indicated drug, followed by 24 hours of drug-free washout prior to seeding, or the cells were treated with the indicated drug for 24 hours and then seeded immediately.

Scratch Assays
For scratch assays, 1-1.5 × 10^6 cells were seeded on a 6-well dish and 24 hours later (at around 90% confluency) the monolayer was scratched with a 200 μL pipette tip. The cell culture plate was then placed in an inverted Zeiss Observer for live-cell imaging (5% CO2, 37 °C). Phase contrast images were taken every 30 mins for 22 h at 8-10 positions along the scratch and 2-3 trials were repeated for each cell line and condition. Data was analyzed with ImageJ to determine the scratch area remaining using the Wound Healing Tool Macro (Collins, 2007). For experiments with Mps1 inhibitors, the cells were either treated for 24 hours with the indicated drug followed by 24 hours of drug-free recovery prior to seeding, or the cells were treated with the indicated drug for 24 hours immediately after seeding. A minimum of 2 trials per cell line and treatment were performed.

Western Blot Analysis
Whole cell lysates were harvested and resuspended in RIPA buffer [25 mM Tris, pH 7.4, 150 mM NaCl, 1% Triton X 100, 0.5% sodium deoxycholate, 0.1% sodium dodecyl sulfate, protease inhibitor cocktail (Sigma, Cat. No. 4693159001), and phosphatase inhibitor cocktail (Sigma, Cat. No. 4906845001)]. Quantification of protein concentration was done using the RC DC Protein Assay (Bio-Rad, Hercules, CA; Cat. No. 500–0119). Equal amounts of lysate were denatured and loaded onto a 10% SDS-PAGE gel. The Trans-Blot Turbo Transfer System (Bio-Rad) and polyvinylidene difluoride membranes were used for protein transfer. Antibody blocking was done with 5% milk in TBST (19 mM Tris base, NaCl 137 mM, KCl 2.7 mM and 0.1% Tween-20) for 1 hour at room temperature except for E-cadherin, cGAS, N-cadherin and Vimentin which used 5% BSA in TBST. The following antibodies and dilutions were used: E-cadherin (Cell Signal, Danvers, MA; Cat. No. 3195) at a dilution of 1:1000 (5% BSA), N-cadherin (Cell Signal; Cat. No. 13116) at a dilution of 1:1000 (5% BSA), Vimentin (Cell Signal; Cat. No. 5741) at a dilution of 1:1000 (5% BSA), EpCAM (Abcam, Cambridge, MA; Cat. No. ab124825) at a dilution of 1:1000 (5% milk), Claudin-7 (Abcam; Cat. No. ab27487) at a dilution of 1:1000 (5% milk), Claudin-7 (Abcam; Cat. No. 5741) at a dilution of 1:1000 (5% BSA), and the remaining bins are populated as evenly as possible (Andrews et al., 2016; Andrews, 2017). Approximately 2% of the resulting bins showed significant variation in count across the CHD population and were thus excluded from further analysis and display using both automated means and human review. The generated reads were demultiplexed and mapped using custom scripts, and plots were generated with G-Graph (Andrews, 2017).
Overexpression of E-Cadherin and EpCAM

The following plasmids were obtained from Applied Biological Materials (Richmond, BC, Canada): E-cadherin (Cat. No. LV704934), EpCAM (Cat. No. LV149412) and an empty control plasmid (Cat. No. LV591). The plasmids were transfected into HCT116 Trisomy 5 cells using Lipofectamine 3000 (ThermoFisher, Cat. No. L3000001). The cells were then selected with puromycin for 2 weeks and then bulk-sorted with FACS for RFP-expressing cells. Invasion assays and western blot analysis was performed on cells that were expanded following FACS purification.

Senescence Staining

To test for senescence, cells were stained for beta-galactosidase using the Cell Signaling Technology kit (#9860) as per the manufacturer’s protocol. As a positive control, cells were treated with etoposide (20 μM) for 24 hours followed by 4 days of recovery. A minimum of 500 cells were counted per condition.

CRISPRi Plasmid Construction and Virus Generation

Guide RNAs for CRISPRi experiments were chosen from (Horlbeck et al., 2016). Guides were cloned into the LRG2.1 mCherry vector (Addgene; Cat. No. 108099) using a BsmBI digestion as previously described (Giuliano et al., 2019). Plasmids were transformed in Stbl3 E. coli (Thermo Fisher; Cat. No. C737303) and sequenced to confirm the presence of the correct gRNA sequence. A dCas9-KRAB construct (Addgene; Cat. No. 85969) was used to knock down target gene expression. Guide RNA sequences are listed in Table S1B.

Lentivirus was generated using calcium phosphate transfection as previously described (Chang et al., 2013). Supernatant was harvested at 2 to 3 intervals 48 to 72 hr post-transfection by filtering through a 0.45 μm syringe, and then frozen at –80 °C for later use or supplied directly to cells with 4 μg/mL polybrene.

RNA Isolation and Quantitative Real-Time PCR

Total cellular RNA was extracted and isolated using TRIzol (Life Technologies; Cat. No. 15596018) and a Qiagen RNeasy Mini Kit (Cat. No. 74106). Total RNA was converted to cDNA using SuperScript™ III First-Strand System kit (ThermoFisher Scientific; Cat. No. 18080051). Quantitative PCR was performed for the target genes using SYBR Premier Ex Taq (Takara; Cat No. RR420L) and quantified using the QuantStudio 6 Flex Real-Time PCR system (Applied Biosystems). Primers are listed in Table S2.

Immunofluorescence and Microscopy

Cells were placed on autoclaved 12mm x 12mm glass coverslips in twelve well cell culture plates. The next day, the cells were treated with the indicated drug in separate cell culture wells. 24 hours later, fresh culture media was added for a post-drug recovery period of 24 hours. Later they were fixed with either 4% paraformaldehyde for 15 min at RT (when staining for STING) or cold (-20 °C) methanol for 5min at RT (when staining for Rel-B). For STING, selective plasma membrane permeabilization was done using 0.2% Saponin in TBS for 2 min. For Rel-B, nuclear permeabilization was done using 0.15% of Triton X-100 in TBS for 10 min. TBS + 1% BSA + 22.52 mg/ml of glycine was used as a blocking agent for 45 minutes. TBS + 1%BSA was used as blocking agent during primary antibody staining (STING at a dilution of 1:1000; Abcam Cat. No. ab181125; RelB at a dilution of 1:500; Abcam Cat. No. ab180127) overnight at 4 °C. After washing the cells, they were incubated with goat anti-rabbit Alexa Fluor 647 (at a dilution of 1:1000; Abcam Cat. No. ab150083). The coverslips were treated with DAPI (0.1μg/ml in TBS) for 2 min. The coverslips were then mounted with Prolong Antifade Mountant (ThermoFisher Scientific; Cat No., P36965). The cells were viewed on a spinning disk confocal microscope (UltraVIEW Vox; PerkinElmer) and quantified using Volocity version 6.3.

Live Cell Imaging

HCT116 cells expressing H2B-GFP were seeded and treated with Mps1 inhibitors as indicated. Live-cell imaging was performed at room temperature using spinning-disc confocal microscopy system (UltraVIEW Vox; PerkinElmer) and a charged-coupled device camera (ORCA-R2; Hamamatsu Photonics) fitted to an inverted microscope (DMi6000 B; Leica) equipped with a motorized piezo-electric stage (Applied Scientific Instrumentation). Overnight imaging was performed using a Plan Apochromat 20X 0.7 NA objective with camera binning set to 2x2. Image acquisition and analysis was performed using Volocity version 6.3 (PerkinElmer). A minimum of 100 cells per condition were analyzed for mitotic errors. Cells were tracked form nuclear envelope breakdown to anaphase exit and were analyzed for errors such as lagging chromosomes, polar chromosomes, anaphase bridges, and multipolar mitoses.
Quantification of Micronuclei

Cells were seeded onto twelve well-cell culture plates. The next day, the cells were treated with Mps1 inhibitors as indicated. Cells were then allowed to recover in drug-free media for 24 hours. Following this period, the cells were stained with 5 ng/ml Hoechst 33342 (Invitrogen; Cat. No. H3570) for 20 min at 37 °C. The cells were viewed under a Nikon Eclipse Ti-S microscope and quantified using NIS elements BR version 4.40. A minimum of 400 cells per condition were analyzed for micronuclei. In every field of view imaged, the total number of cells and the total number of cells with micronuclei were counted to determine the percent of cells with micronuclei.

Knocking-in AZ3146-Resistance Mutations with CRISPR-Mediated HDR

Mps1-targeting guides and single-stranded donor templates to introduce the AZ3146-resistance mutations were designed using Benchling (www.benchling.com). In addition to the C604W and S611G alterations, multiple silent mutations were included in the donor oligo to prevent re-cutting following template-mediated repair (Table S1A). Guide RNAs were cloned into the Lenti-Cas9-gRNA-GFP vector (Addgene # 124770) as previously described (Giuliano et al., 2019). To perform CRISPR-mediated HDR, 2 μg of Mps1 gRNA plasmid was transfected along with 100 pmol of ssODN into Cas9-expressing cell lines using Lipofectamine 3000 (Thermo Fischer Scientific, Cat. No. L3000015). Successful knock-in was confirmed using the primers listed in Table S1A.

TCGA Data Analysis

Survival data for TCGA patients was acquired from (Liu et al., 2018). P53 mutation data for TCGA patients was acquired from (Bailey et al., 2018). The EMT gene signature was acquired from (Gibbons and Creighton, 2018). Across the 33 available TCGA cohorts, we first eliminated the six cohorts with fewer than 100 patients. Next, we chose “overall survival time” as a default endpoint, as it reflects an objective and unambiguous event. However, we noted that five of the remaining cohorts had fewer than 15 deaths; for those five cohorts we used “progression-free survival” as a clinical endpoint rather than “overall survival” (Table S3A).

Patient survival was analyzed using Cox proportional hazards modeling. The Cox model is given by the following function:

\[ h(t, X) = h_0(t) \exp(\sum_{i=1}^{n} \beta_i X_i) \]

In this model, \( t \) is the survival time, \( h(t, X) \) is the hazard function, \( h_0(t) \) is the baseline hazard, \( X_i \) is a prognostic variable, and \( \beta_i \) indicates the strength of the association between a variable and survival. In this model, patients have a baseline, time-dependent risk of death (\( h_0(t) \)), and this risk is modified by time-independent prognostic features that either increase (\( \beta_i > 0 \)) or decrease (\( \beta_i < 0 \)) the patient’s risk of death. In this paper, we report Z scores, which are calculated by dividing the regression coefficient (\( \beta_i \)) by its standard error. If a chromosome gain is significantly associated with patient death, then a Z score >1.96 corresponds to a P value < 0.05. In contrast, a Z score less than −1.96 indicates that a chromosome gain is significantly associated with patient survival (or, alternately, that a chromosome-loss event is associated with death). Cox proportional hazards models and their application to TCGA data is discussed in more detail in (Smith and Sheltzer, 2018).

Karyotype data for TCGA patients was acquired from (Taylor et al., 2018). Aneuploidy calls were made based on Affymetrix SNP 6.0 profiling of tumor samples. Tumor ploidy was determined using ABSOLUTE and was used to determine the baseline copy number for each patient (Carter et al., 2012). Arms or chromosomes in which more than 80% of the region was affected by a CNA were called positive for an arm or whole-chromosome alteration, while arms or chromosomes in which less than 20% of the region was affected by a CNA were considered to be negative. Aneuploidy scores were calculated by summing the total number of arm-length alterations detected in a tumor sample.

For analysis including p53, only tumors that harbored a non-synonymous mutation were considered p53-mutant (missense, nonsense, or frameshift). To assess the link between bulk aneuploidy and patient outcome, we compared survival between “low aneuploidy” tumors (≤ 20th% aneuploidy score) and “high aneuploidy” tumors (≥ 80th% aneuploidy score). Cox proportional hazards modeling was performed using Python, pandas, rpy2 and the R survival package. Code written for this analysis is available at https://github.com/joan-smith/aneuploidy-survival. This analysis also relied on packaged code from (Smith and Sheltzer, 2018), now also available at https://github.com/joan-smith/biomarker-survival for ease of reuse.

Animals and Splenic Injections

Nude mice (NU/J female, 6-week old; JAX stock #002019) were obtained from the Jackson Laboratory. The Cold Spring Harbor Laboratory IACUC reviewed and approved all procedures. HCT116 and trisomic cells were injected via splenic injection to generate a preclinical model of hepatic metastasis. The cell lines were transduced to express luciferase (Addgene #75020) for bioluminescence imaging.

Anaesthetized mice were swabbed with an alcoholic solution of iodine or chlorhexidine before a small incision (10-15mm) was made. The spleen was identified and carefully manipulated through the incision to sit outside the mouse on a moist gauze swab. 10^6 cells in 100 μl PBS suspensions were injected into the organ. For the spleen, effective injection was monitored by bleeding of the organ and lack of bleeding. In some cases, splenectomy can be performed by clamping and cauterizing the splenic arteries and venous supply, thus effectively removing the spleen. The process of splenectomy carries a significant risk of bleeding, which typically occurs if the blood supply is not cauterized correctly. Successful splenic injections were verified by visual assessment for minimal bleeding. The abdominal muscle wall was then closed using an absorbable suture. The incision was closed in the top two layers using a continuous vicryl suture (Matric 1.5, Ethicon) for the peritoneum and a wound clip. Analgesia was performed...
with meloxicam (5-10 mg/kg PO or SQ) or Ketoprofen (5mg/kg, SC) at daily intervals for 3 days as necessary. Animals were allowed to recover from anesthesia in pre-warmed clean cages for a period of 20 minutes.

Following the injections, mice were imaged regularly for 7 weeks to monitor for metastatic growth. To perform BLI imaging, mice were injected intra-peritoneally with XenoLight D-Luciferin - K+ Salt Bioluminescent Substrate (15 mg/ml, 100ul/10g; Perkin Elmer). The mice were imaged with Xenogen IVUS Spectrum (Perkin Elmer). 42 days post-injection, the mice were euthanized and metastatic nodules were visually scored. In total, four cohorts of mouse injections were performed. Note that in the second cohort, three mice injected with HCT116 Ts5 cells and one mouse injected with HCT116 cells died prior to day 42. The metastatic nodule counts in these mice are therefore from their time of death, and not day 42.

QUANTIFICATION AND STATISTICAL ANALYSIS

Graphpad Prism was used for all statistical work, excluding the survival analysis. Unpaired t-test were used to determine statistical significance. Statistical details can be found in the figure legends which describe the statistical tests used, exact value of n, what n represents, and dispersion and precision measures.

DATA AND CODE AVAILABILITY

Code written for this analysis is available at https://github.com/joan-smith/aneuploidy-survival. Survival data for TCGA patients was acquired from (Liu et al., 2018). P53 mutation data for TCGA patients was acquired from (Bailey et al., 2018). The EMT gene signature was acquired from (Gibbons and Creighton, 2018). Processed Z scores are included in Table S3.