### A genetic interaction map of cell cycle regulators

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ABSTRACT Cell-based RNA interference (RNAi) is a powerful approach to screen for modulators of many cellular processes. However, resulting candidate gene lists from cell-based assays comprise diverse effectors, both direct and indirect, and further dissecting their functions can be challenging. Here we screened a genome-wide RNAi library for modulators of mitosis and cytokinesis in Drosophila S2 cells. The screen identified many previously known genes as well as modulators that have previously not been connected to cell cycle control. We then characterized ~300 candidate modifiers further by genetic interaction analysis using double RNAi and a multiparametric, imaging-based assay. We found that analyzing cell cyclerelevant phenotypes increased the sensitivity for associating novel gene function. Genetic interaction maps based on mitotic index and nuclear size grouped candidates into known regulatory complexes of mitosis or cytokinesis, respectively, and predicted previously uncharacterized components of known processes. For example, we confirmed a role for the Drosophila CCR4 mRNA processing complex component *l*(2)NC136 during the mitotic exit. Our results show that the combination of genome-scale RNAi screening and genetic interaction analysis using process-directed phenotypes provides a powerful two-step approach to assigning components to specific pathways and complexes.

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#### INTRODUCTION

Large-scale genetic screens have identified components of many biological processes in a broad spectrum of organisms (Patton and Zon, 2001; Jorgensen and Mango, 2002; St Johnston, 2002; Boutros and Ahringer, 2008). Such experiments have considerably expanded our knowledge of regulatory complexes and pathways. In recent years, classical genetic screens have been complemented by cellbased, loss-of-function experiments using RNA interference (RNAi; Kiger *et al.*, 2003; Boutros *et al.*, 2004; Carpenter and Sabatini, 2004; Kittler *et al.*, 2004; DasGupta *et al.*, 2005). Although screening technologies have made significant advances, moving from candidate "hit" lists to precise delineation of functional relationships has remained challenging. Prioritization of candidates for follow-up experimentation often relies on prior knowledge, leaving uncharacterized genes untouched. Therefore systematic and scalable secondary lines of screens are necessary to group candidates into functional categories, pathways, or complexes.

In yeast, systematic double-perturbation experiments, termed synthetic genetic arrays (SGAs), dSLAM, or E-MAPs (Pan *et al.*, 2004; Tong *et al.*, 2004; Schuldiner *et al.*, 2005), have been used to explore diverse biological processes and successfully identified previously undiscovered functional relationships (Tong *et al.*, 2004; Schuldiner *et al.*, 2005; Collins *et al.*, 2007). In both *Saccharomyces cerevisiae* and *Schizosaccharomyces pombe*, large collections of mutant alleles are available to generate maps of biological processes and compare their phylogenetic relationships (Dixon *et al.*, 2008; Roguev *et al.*, 2008; Jonikas *et al.*, 2009). The largest-scale

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Abbreviations used: APC/C, anaphase-promoting complex/cyclosome; CCT, cytosolic chaperonin–containing t-complex; CSI, connection specificity index; CSN, COP9-signalosome; FDR, false-discovery rate; GI, genetic interaction; PCC, Pearson correlation coefficient; pH3, phospho-Histone H3; RNAi, RNA interference; SAC, spindle assembly checkpoint; SGI, synthetic genetic interaction.

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synthetic genetic interaction study to date has generated genetic interaction profiles for ~75% of the yeast genome (Costanzo *et al.*, 2010). Although full-genome genetic interaction analysis has not yet been approached in metazoan cells, the concept of systematic codepletion and quantitative analysis has been applied in *Caenorhabditis elegans* (Lehner *et al.*, 2006; Tischler *et al.*, 2008), *Drosophila* cells (Bakal *et al.*, 2008; Horn *et al.*, 2011; Fischer *et al.*, 2015), and mouse (Roguev *et al.*, 2013) and human (Laufer *et al.*, 2013; Wang *et al.*, 2014) cell lines.

Here we describe a two-step approach that first prioritizes genes by genome-wide screening for subsequent medium-scale, imagingbased synthetic genetic interaction (SGI) analysis. We screened for modulators of cell cycle regulation by genome-wide RNAi and highthroughput imaging of *Drosophila* cells. Then we used double RNAi to systematically map the functional relationships between these genes. This analysis grouped genes into functional modules and generated hypotheses for the function of genes not previously implicated in cell cycle regulation, including *CG11753* (*Drosophila SYS1*) and *I*(*2*)*NC136* (*Drosophila CNOT3*). The results of this study demonstrate the use of synthetic genetic interaction experiments in metazoan cells to refine functional predictions from large-scale perturbation experiments.

#### **RESULTS AND DISCUSSION**

### A two-step RNAi screening approach for regulators of the cell cycle

To identify and map potentially novel regulators of the cell cycle, we used high-throughput imaging to measure the effects of single and double perturbations on cell cycle-relevant phenotypes in Drosophila S2 cells (Figure 1A and Materials and Methods). First, using a genome-wide RNAi library targeting ~98% of the protein coding transcripts (Horn et al., 2010; Horn and Boutros, 2013), we systematically interrogated the Drosophila genome for effects of single-gene knockdowns on the cell cycle. Fluorescence microscopy images of nuclei (DNA) and phospho-Histone H3-positive (pH3; phosphorylated from mitotic prophase to anaphase) cells were captured for each condition after 5 d of knockdown. We focused our analysis on three phenotypes extracted from the images: number of nuclei (cell number) as a measure of cell viability and average nuclear area and mitotic index (proportion of pH3-positive cells) as measurements directly related to cell cycle-relevant processes (Materials and Methods). After selecting a set of ~300 putative novel cell cycle regulators from the genome-wide data set, in a second experiment we tested double knockdowns of the candidates with 14 well-characterized cell cycle regulators using the same microscopic readouts and assessed genetic interactions. In that way, we aimed at mapping the putative novel regulators to known cell cycle processes.

### Results of the genome-wide screen

To identify RNAi reagents that had effects on the cells in the genome-wide screen, we computed z-scores for each of the three phenotypes. For nuclear area and mitotic index phenotypes, we adjusted for cell count effects using regression analysis (*Materials and Methods*). The z-scores of two replicates showed strong positive correlations for all three phenotypes (Supplemental Figure S1A). By comparing z-scores between the three phenotypes across the entire screen, we found that they provided complementary information about the effects of gene knockdowns as indicated by their low Pearson correlation (Supplemental Figure S1B). Figure 1B highlights the information gain by assessing nuclear area and mitotic index in addition to cell count: although genes implicated in different biological processes affected cell counts similarly (or had no effect at all), they showed differential effects in the other phenotypes. For example, knocking down genes encoding sperm-specific dynein intermediate chains (Sdic1, Sdic2, Sdic3, Sdic4) or genes involved in mRNA cleavage and polyadenylation (Cpsf100, Cpsf160, and Cpsf73) showed mild to no effects on cell counts, but the mitotic index was increased after perturbation of the dynein intermediate chains and decreased after perturbation of the mRNA-processing factors (Figure 1C). This was also in agreement with their known implications for the cell cycle, where dyneins are required for mitotic progression and mRNAprocessing factors are important during interphase (Manley, 1995; Nurminsky et al., 1998). In another example, we found that genes with a role in cytokinesis could have diverging effects on the mitotic index (increased for tum, pbl, zip, and scra; decreased for Incenp, borr, Det, and ial), while invariably decreasing cell counts and increasing nuclear area (Figure 1C). This confirmed the requirement of the components of the chromosomal passenger complex inner centromere protein (Incenp), borealin (Borr; Drosophila CDCA8), and deterin (Det, Drosophila survivin) for Histone H3 phosphorylation by ial (aurora B kinase [aurB]; Carmena et al., 2012), in contrast to the proteins tumbleweed (tum, Drosophila RACGAP1) and pebble (pbl, Drosophila ARHGEF5), which are essential for cytokinesis but do not alter phospho-Histone H3 phosphorylation (Zavortink et al., 2005). Overall, assessing the three complementary phenotypes provided a more detailed view of gene functions and highlighted genes with potential roles in the cell cycle.

To get a more comprehensive picture of factors affecting nuclear area and mitotic index as markers of cell cycle progression, we filtered the results of the genome-wide screen for all treatments with an absolute z-score  $\geq$ 3 for either of the phenotypes. Of ~15,000 genes assessed by RNAi (Supplemental Table S1), knockdowns of ~1000 (6.7%) resulted in such a phenotype (Supplemental Table S2). Approximately 17% of them were specific to nuclear area and 75% to mitotic index, and 8% scored in both phenotypes. We grouped the phenotypic profiles of the 1000 genes by hierarchical clustering of their z-scores (Figure 2A). This identified several clusters that were enriched for known cell cycle regulators (Figure 2, B-E). For example cluster I (Figure 2B) grouped proteins required for formation of the mitotic spindle (Teixido-Travesa et al., 2012), including tubulins, as well as components of the augmin and CCT chaperonin complexes (Supplemental Table S3). Loss of function of these proteins commonly increased the mitotic index, indicating a mitotic arrest. In contrast ,cluster IV (Figure 2E) grouped components of the COP9 signalosome (CSN; Supplemental Table S3) complex, which is required during the G1/S transition of the cell cycle (Doronkin et al., 2003). Knocking down CSN members commonly decreased the mitotic index, indicating a cell cycle arrest before mitosis. To address more systematically the question of which processes could be detected by our assay, we manually annotated a high-confidence list of 131 proteins with well-characterized roles during G1/S transition, the G2/M checkpoint, and the mitotic spindle, as well as regulators of mitotic progression and cytokinesis (Supplemental Table S3). We found that 96 of the known factors (73%), covering all manually annotated cell cycle processes, had an absolute z-score  $\geq 3$  in nuclear area and/or mitotic index (Supplemental Figure S2A), which validated our screening approach. Of interest, when comparing the phenotypes (nuclear area and mitotic index) across the different processes, we found that they affected the phenotypes differently (Supplemental Figure S2, B–D), which confirmed the previous observation that the measurement of different phenotypes was required to be able to assay different cell cycle processes.

The genome-wide screen identified many components with phenotypes similar to those of known cell cycle regulators





FIGURE 1: Setup of the combined genome-wide RNAi and synthetic genetic interaction analysis using multiple phenotypic readouts. (A) Genome-wide RNAi screening and high-throughput imaging to quantify the number of cells, nuclear area, and mitotic index are used to identify cell cycle regulators, which are further characterized by genetic interaction analysis. See the text for details. (B) Comparison of the three phenotypes across the genome-wide data set. The knockdown conditions (*x*-axis) are sorted according to their cell count phenotype (top). Nuclear area (middle) and mitotic index (bottom) phenotypes are shown in the same order. Highlighted in red are proteins that show differential effects across the phenotypes. Dashed lines indicate *z*-score cut-offs of 3 and –3 (points below the cut-offs are light gray, and points above are dark gray). (C) Bar plots comparing the phenotypes (per functional group) for the knockdowns highlighted in B. Knockdowns of dynein IC proteins and mRNA-processing factors show opposing effects on mitotic index and have no effects on the other phenotypes. Knockdowns of proteins required for cytokinesis all increase the nuclear area, whereas they have opposing effects on mitotic index. Dashed lines indicate *z*-score cut-offs of 3 and –3.

(Figure 2, A–E). We attempted to use genetic interaction analysis to elucidate their functional relationship with the known cell cycle machinery. To this end, we selected genes from the genome-wide screen that were expressed and displayed mitotic index and nuclear area phenotypes, prioritizing genes that had not been linked to cell cycle regulation previously (Bettencourt-Dias *et al.*, 2004; Bjorklund *et al.*, 2006; Kondo and Perrimon, 2011). We also excluded genes that were not conserved in humans, as well as

genes with strong viability defects (for details of candidate selection, see the Supplemental Methods). This left us with 275 potential novel modulators from the genome-wide screen to be subjected to genetic interaction analysis: 238 with mitotic index phenotypes and 37 with nuclear area phenotypes (Supplemental Table S4). In addition we included several controls: genes with known roles in cell cycle regulation that also scored in the genome-wide screen, such as components of the anaphase-promoting



FIGURE 2: Genome-wide RNAi screening identifies known and potential novel regulators of cell cycle. (A) Heatmap of z-scores for 1018 genes with an absolute z-score  $\geq$  3 in nuclear area and/or mitotic index after hierarchical clustering. Black bars to the left of the heatmap highlight genes that were selected for the SGI analysis. Bars on the right of the heatmap highlight clusters of genes that are enriched for factors with known roles in cell cycle and are shown in greater detail in B–E. (B–E) Detailed heatmaps of the clusters highlight din A. Colors of gene names indicate their membership in certain groups of cell cycle regulators (see box). Arrows highlight genes that were selected for SGI analysis. See the text for details. (F) Scatter plot of nuclear area and mitotic index phenotypes for genes selected for SGI analysis (black) on top of the genome-wide data (gray). The distribution of candidates and controls selected for the SGI analysis shows that it is representative of the effect range observed in the genome-wide screen. Different types of candidates and controls are indicated by shape (see legend and Supplemental Tables S4 and S5). APC/C, anaphase-promoting complex/cyclosome; CCT, cytosolic chaperonin–containing t-complex; CSN, COP9 signalosome; SAC, spindle assembly checkpoint.



FIGURE 3: Genetic interactions across different phenotypes. (A) Comparing the number of genetic interactions affecting cell count, mitotic index, or nuclear area shows that the majority of interactions are found for the mitotic index phenotype. Interactions were estimated from a set of 14 "query" genes and 350 "candidate" genes/controls. (B–D) Node degree distributions showing number of positive (yellow bars) and negative (blue bars) genetic interactions per phenotype and on a per-gene basis. (E) A Venn diagram shows that there are overlapping, but also many exclusive, genetic interactions between the phenotypes. (F, G) Genetic interactions within the CSN largely overlap between the phenotypes (F), whereas genetic interactions within the APC/C are exclusive to the mitotic index (G). In F and G. all genetic interactions are alleviating, that is, were less severe than expected according to the reference model. All genetic interactions were called at FDR of 1%.

complex/cyclosome (APC/C) and the CSN complex, as well as genes that had no or only mild effects on the phenotypes of interest, in order to prevent biases during the analysis of genetic interactions. The list of candidates covered a broad range of mitotic index and nuclear area phenotypes, representative of the range of effects observed in the genome-wide screen (Figure 2F).

# Different phenotypes identify nonredundant genetic interactions

To test the set of candidates from the genome-wide screen for genetic interactions, we systematically codepleted them with 14 known cell cycle regulators ("query" genes, Supplemental Table S5) in all pairwise "candidate  $\times$  query" combinations. To reduce the influence of potential off-target effects on the interaction analysis, we used two sequence-independent RNAi reagents to target each candidate and each query gene, resulting in 20,216 co-RNAi experiments with four independent measurements per gene pair. This experimental setup was shown previously to allow robust estimation of single- and double-RNAi phenotypes (Horn *et al.*, 2011). Genetic interactions and their significance were measured independently for each phenotype as deviations from the multiplicative model, which describes the expected combinatorial effect as the product of the single-knockdown effects, and are summarized in  $\pi$ -scores (Supplemental Table S6).

For cell count, mitotic index, and nuclear area phenotypes we observed 1419 genetic interactions at an experiment-wide falsediscovery rate (FDR) of 1%. The mitotic index yielded a ~10- and 3-fold higher rate of genetic interactions than cell count and nuclear area, respectively (Figure 3A), which might be due to the phenotypes of the candidates that were mostly mitotic index effects (Supplemental Table S4). We assessed the node degree distribution for each phenotype (Figure 3, B-D). The majority of genes had only few interactions (Figure 3, B-D, 1-3) independent of the phenotype. The number of positive and negative interactions tended to be unbalanced, and the strongest differences were seen for the mitotic index phenotype. Here many genes had a few positive interactions (Figure 3C, 1-3), and a few genes had many negative interactions (Figure 3C, 7-9). By comparing the genetic interactions between phenotypes, we found that only 31 gene pairs interacted in all three phenotypes. In contrast, 1183 gene pairs (83.4%) displayed interactions specific for a single phenotype (Figure 3E). Whereas genetic interactions between some known cell cycle regulators, such as the members of the CSN complex, were observed in all three phenotypes (Figure 3F), other regulators, including the APC/C subunits cdc23, ida, Apc10, and fzy, showed genetic interactions for the mitotic index phenotype only (Figure 3G). This indicated that measuring different cell cycle-relevant phenotypes might be required to connect novel genes to known processes.

#### Phenotype-specific interaction networks functionally connect candidate genes

To systematically predict functional associations of candidate genes, we assessed the similarity of their genetic interaction profiles. This type of analysis was previously shown to provide robust approximations of functional associations (Costanzo *et al.*, 2010; Fuxman Bass *et al.*, 2013). We first calculated the Pearson correlation coefficient (PCC) between the interaction profiles of all candidates for each phenotype. To correct for genes that shared a high PCC with many genes, we transformed the PCC data into the connection specificity index (CSI; Figure 4A), which reduces potential nonspecific similarities between genes by ranking the similarities according to the connectivity of their interaction partners (Green *et al.*, 2011; Fuxman Bass *et al.*, 2013).

We assessed whether the CSI improved functional associations between candidate genes compared with their single-knockdown phenotypes. As expected, using the mitotic index, all APC/C components tested (*ida*, *cdc23*, and *fzy*) had a high mitotic index–based CSI with Apc10 (Figure 4B). In contrast, components of the Augmin complex (*wac*, *dgt2*) and the DREAM complex (*mip120*, *mip130*) shared a lower CSI with Apc10, despite their similar single-knockdown effect on mitotic index (Figure 4B). Several other genes, such as the known modulator of mitosis pontin (pont, a member of the INO80/SWR1 chromatin-remodeling complexes; Ducat *et al.*, 2008),



FIGURE 4: Genetic interaction profiles associate genes with similar functions. (A) Processing of genetic interaction (GI) profiles to predict functional similarity between genes. The profiles (14  $\pi$ -scores) of each candidate gene pair were compared per phenotype by calculating the PCC. The example shows the PCC between the mitotic index-based profiles of the APC/C components *Apc10* and *ida*. The PCC was corrected by calculating the CSI with a correction factor/constant of 0.01. The example shows the relation between the PCC and CSI of Apc10. (B) Comparison of the mitotic index-based CSI connecting *Apc10* to all tested genes with the single-knockdown effect of all genes shows additional APC/C components (and cell cycle regulators) in proximity to *Apc10*. (C) Comparison of the nuclear areabased CSI connecting *Rho1* to all tested genes with the single-knockdown effect of all genes shows additional cytokinesis components (and cell cycle factors) in proximity to *Rho1*. See the text for details. The point size is proportional to the CSI in B and C.

also had a high CSI with Apc10 without increasing the mitotic index when depleted alone (Figure 4B), drawing a potential connection to the APC/C. We also compared the nuclear area-based CSI of the cytokinesis regulator *Rho1* to the single-knockdown effects of all other genes. This highlighted many known cytokinesis factors (e.g., *tum*, *zip*, *Det*) but deprioritized other, likely functionally independent genes that also showed an increased nuclear area, such as *twinstar* (*tsr*, *Drosophila* cofilin), a component required for mitotic telophase and cytokinesis (Gunsalus *et al.*, 1995; Figure 4C). These results show that the CSI could highlight genes with similar functions that could not be differentiated based on their single-knockdown effects.

To identify association more systematically for each phenotype, we placed all candidate genes in a force-directed network according to their functional similarity (CSI; Figure 5, A and B, and Supplemental Figure S5A). The mitotic index–based network associated the APC/C with components facilitating mitotic spindle organization and regulation, including dynactin subunits (Arp87C) and the CCT chaperonin complex member Cct5 (Figure 5A). These close associations with the APC/C were not seen for the other phenotypes (Figure 5B and Supplemental Figure S5, A and B). In the proximity of APC/C, we observed additional interesting connections. We already highlighted the high CSI of the INO80/ SWR1 complex member pont with Apc10. We found more members of different chromatin-remodeling complexes in the proximity of the APC/C, including domino (dom) and Yeti (both members of SWR1), Nurf-38 (a member of the NURF complex), and female sterile homeotic (fs(1)h, a BET-family protein). Further, several genes involved in regulation of transcription, such as pointed (pnt, ETS transcription factor) and I(2)NC136 (Not3, a member of the CCR4-NOT deadenylation complex), were connected to the APC/C. These results suggested a specific role for regulation of chromatin and transcription during mitosis (Morrison and Shen, 2009; Tanenbaum et al., 2015). Other processes/complexes in the proximity of the APC/C included the DREAM complex (mip120, mip130), which was previously shown to regulate mitotic events



FIGURE 5: Networks based on cell cycle–relevant phenotypes. (A) Mitotic index–based and (B) nuclear area–based functional associations of candidate genes show known cell cycle processes (different colors) linked to each other and to uncharacterized genes/processes. Genes were placed in a force-directed network based on the genetic interactions for each phenotype. Only gene pairs (connections) with CSI > 0.95 are shown. (C) *l*(*2*)*NC136* and *CG11753* are required for mitotic progression in *Drosophila* cells. Each gene was knocked down by two independent dsRNA designs, *ida* was knocked down as positive control, and firefly luciferase was used as negative control. At 96 h past dsRNA transfection, the G2/M checkpoint was triggered by doxorubicin or etoposide, and cells were allowed to leave M phase for 6 h before assessment of the mitotic index. The data show the mean of 32 (control) or 4 (*ida*, *l*(*2*)*NC136* and *CG11753*) replicates, and the error bars indicate SEM. (D) Functional depletion of the human orthologues of *l*(*2*)*NC136* and *CG11753* (*CNOT3* and *SYS1*, respectively) using siRNAs for 72 h shows a delayed mitotic exit after release from prometaphase arrest in HeLa cells. Prometaphase arrest was induced by nocodazole treatment for 18 h, and the fraction of G2/M cells was assessed at different time points after release by fluorescence-activated cell sorting analysis. The data show the mean of three replicates; the error bars indicate SEM.

(Beall et al., 2004; Georlette et al., 2007), and the regulators of nuclear import *Nup107*, *Nup160*, and *Kap-alpha3*. The latter finding suggested a role for nuclear pore complex components during mitosis that has been described before (Loiodice et al., 2004). Other subnetworks highlighted known cell cycle–regulatory complexes, such as the CSN, but also processes that are less well understood in the context of the cell cycle, such as vesicle trafficking.

The mitotic index-based network failed to identify known functional associations between cytokinesis regulators (Figure 5A and Supplemental Figure S5C). In contrast, the nuclear area-based network connected Rho1 with other components of cytokinesis (tum, ial, Det, lin19, and zip; Figure 5B). Of interest, several regulators of vesicle trafficking and exocytosis were found in subnetworks connected to cytokinesis, suggesting a potential role of exocytosis during cytokinesis that has been previously described (Skop et al., 2001). Although vesicle components grouped in the mitotic index-based network as well (e.g., Arf72A, CG11753, Syx1A), they were not connected directly to any of the other known processes, which highlights the information gain from generating phenotype-specific networks. Despite the conservation of some subnetworks across all three phenotypes, such as the CSN or the APC/C, the connections between the processes and uncharacterized genes were phenotype specific. With mitotic index-based and nuclear area-based networks being stronger indicators of cell cycle-specific processes (Figure 5, B and C), this provided us with testable hypotheses for potentially novel regulators of the cell cycle.

## Golgi-resident and mRNA processing factors modulate mitotic progression

To assess the validity of our network-based prediction of gene functions, we tested the role of two genes in cell cycle-relevant assays. The mitotic index-based network connected the APC/C and other regulators of mitosis to chromatin-remodeling complexes and transcriptional regulators. We followed up on one of the genes in proximity to the APC/C, I(2)NC136, which is a member of the conserved deadenylation complex CCR4-NOT. Its human orthologue, CNOT3, regulates mitotic progression by destabilizing the mRNA of the spindle assembly checkpoint (SAC) component MAD1 (Takahashi et al., 2012). To confirm the predicted role for I(2)NC136, we knocked down the gene in Drosophila S2 cells with double-stranded RNAs (dsRNAs) for 4 d and triggered a G<sub>2</sub>/M checkpoint arrest by doxorubicin or etoposide treatment for 6 h to allow cells with intact mitotic progression machinery to exit M phase (Figure 5C). Cells depleted of I(2)NC136 displayed approximately fourfold higher frequency of pH3-positive nuclei in both treatments than with a control knockdown, indicating a delay in mitotic exit. To validate the conserved function of I(2)NC136 in human cells, we knocked down its orthologue, CNOT3, in HeLa cells using siRNAs and assessed the cycle after release from a nocodazole-induced prometaphase arrest. Similarly to Drosophila cells, HeLa cells displayed a significant mitotic delay (assessed by quantifying  $G_2/M$ -phase contents; p <0.019, paired Student's t test) after knockdown of CNOT3 (Figure 5D and Supplemental Figure S5D) compared with a control siRNA. These results confirmed a role of l(2)NC136 during mitotic exit, as suggested by the network-based predictions.

Another process highlighted in both the mitotic index-based and nuclear area-based networks is vesicular trafficking. Several components of this network were previously shown to have cell cycle-specific functions. For example, *syntaxin1A* (*syx1A*), which encodes for a target-soluble *N*-ethylmaleimide-sensitive factor attachment protein receptor, was shown to be required for mitotic telophase (Somma *et al.*, 2002). *Gmap* and *Arf72A* are crucial for Golgi inheritance during cell division (Rios *et al.*, 2004; Eisman *et al.*, 2006). These genes formed a subgroup in the mitotic index-based network also containing the largely uncharacterized gene *CG11753* (Figure 5B). Its human orthologue, *SYS1*, has been shown to have a function during Golgi-targeted vesicular transport (Behnia *et al.*, 2004; Setty *et al.*, 2004). In agreement with its predicted role, depletion of CG11753 in *Drosophila* S2 cells and SYS1 in HeLa cells delayed mitotic progression (Figure 5, C and D).

The mitosis phenotypes seen for *l*(2)*NC136/CNOT3* and *CG11753/SYS1* in independent assays in *Drosophila* and human cells showed that hypothesis generated from mitotic index– or nuclear area–based interaction networks could be validated. Future studies could address in a more detailed way the manner in which regulation of transcription and vesicular traffic connect to cell cycle/mitosis. Further, additional subnetworks could be subjected to

similar follow-up analysis to understand their role during the cell cycle.

### Conclusion

This study demonstrates how unbiased, genome-wide screening followed by genetic interaction analysis can fine-tune predictions of gene function and guide follow-up experimentation by providing more specific and testable hypotheses. Of importance, predictions of gene functions from interactions based on phenotypes relevant to the biology examined, such as mitotic index and nuclear area to assay cell cycle, were superior to predictions of gene functions from viability/cell count-based interactions. Genetic interaction networks based on relevant phenotypes reconstructed known functional relations during mitotic progression and cytokinesis and suggested potentially novel regulators. We validated novel roles for an mRNAprocessing component and a Golgi-vesicular transport protein during mitotic exit.

With automated imaging being a versatile tool to simultaneously record multiple quantitative phenotypes (Perlman *et al.*, 2005; Fuchs *et al.*, 2010; Fischer *et al.*, 2015), this approach could be expanded to other biological processes by assaying relevant markers. Established reporter-based readouts to monitor the activity of signaling pathways could also be used to perform genetic interaction analysis, and differential networks could be built from experiments using different ways to activate or suppress the pathway.

### MATERIALS AND METHODS

#### Drosophila tissue culture

We cultured Schneider S2 cells adjusted to serum-free growth medium (D.Mel-2; ThermoFisher, Waltham, MA) in Express Five SFM supplemented with 20 mM GlutaMAX and 1% penicillin-streptomycin (all from Invitrogen by ThermoFisher).

### **RNAi** library

We used a genome-wide RNAi library targeting ~98% of all coding genes in the *Drosophila* genome designed against FlyBase annotations for the *Drosophila melanogaster* Berkeley *Drosophila* Genome Project genome, releases 4 and 5, using NEXT-RNAi software (Horn et al., 2010). In addition, the library covered 1254 strongly expressed regions from the Heidelberg predictions (Heidelberg Collection) of *Drosophila* genes not covered by FlyBase (Hild et al., 2003). Primer and dsRNA sequence information, target mappings, and the analysis of specificity and additional features of the dsRNA designs are available through GenomeRNAi (www.genomernai.org; Schmidt et al., 2013). For screening, the library was aliquoted in 384-well plates (black/clear; BD Falcon) with an average of 250 ng of dsRNA/ well in 5 µl of water.

#### Genome-wide RNAi screening

In each biological replicate, we seeded 6500 cells in 40  $\mu$ l of culture medium with 0.2  $\mu$ l of 0.4 mg/ml dimethyldioctadecylammonium bromide (DDAB) per well (in black/clear 384-well plates; BD Falcon) and incubated the plates for 5 d at 25°C before fixation, staining, and imaging of cells.

#### Cell staining and imaging

Cell stainings were done using a Beckman Biomek FX robot with 384-well tip head. First, of the 40  $\mu$ l of assay volume, 15  $\mu$ l was removed, and cells were fixed/permeabilized for 60 min at room temperature by addition of 40  $\mu$ l of a 6% paraformaldehyde (Sigma-Aldrich, St. Louis, MO)/0.3% Triton X-100 (AppliChem, Darmstadt, Germany) solution in phosphate-buffered saline (PBS; Sigma-Aldrich).

Cells were washed by removing 40 µl of supernatant and adding of 50  $\mu$ l of PBS. Next 50  $\mu$ l of the supernatant was removed, and 10  $\mu$ l of a monoclonal phospho-Histone H3 antibody (conjugated to Alexa Fluor 647; 3458; Cell Signaling) diluted 1:750 in 4% bovine serum albumin (GERBU, Heidelberg, Germany)/0.1% Triton X-100 (in PBS) was added and incubated overnight at 8°C (protected from light). On the next day, 10 µl of supernatant was removed, and DNA was stained by addition of 40 µl of Hoechst 33342 (Invitrogen) diluted 1:2000 in PBS with incubation for 30 min at room temperature. Finally, cells were washed once by removing 40 µl of supernatant and adding 50 µl of PBS and washed two more times by removing 50 µl of supernatant and adding 50 µl of PBS. Plates were sealed with aluminum sealing tape (Corning, Corning, NY) and imaged directly or stored at 8°C until imaging. Fluorescence images were acquired on an IN Cell Analyzer 2000 (GE Healthcare, Little Chalfont, UK) using a 4× objective, which enables capture of the entire well in one image. One image was acquired for each of the two channels (Hoechst and phospho-Histone H3 antibody).

### Image analysis

Image analysis and feature extraction from the 4× images (16 bit, 2048  $\times$  2048 pixels) were performed using the R/Bioconductor package EBImage (Pau et al., 2010). The nuclei were segmented and identified separately for the Hoechst and phospho-H3 channels. Adaptive thresholding (width of moving window, 4 pixels) was used to separate nuclei areas from the image background. Nuclei were identified by local-maximum search on the fluorescence images. The nuclei areas were extended from the local maxima by a propagation algorithm (Jones et al., 2005). Nearest-neighbor search was used to match phospho-H3 objects to nuclei in the Hoechst channel (only mitotic nuclei were visible in the phospho-H3 channel). Matches were accepted if the distance between the centers of the objects in both channels was <5 pixels. Eighty-four quantitative features were extracted, of which we focused on three: nuclei count (a correlate of cell count), mitotic index (the ratio of nuclei in the phospho-H3 channel and the Hoechst channel), and nuclear area.

#### Computational analysis of the genome-wide RNAi screen

Raw data of each feature were first log<sub>2</sub>-transformed. To account for row and/or column effects, a local polynomial regression (loess) on the spatial coordinates was fitted to each plate and subtracted. Loess-normalized values were further adjusted for the plate variance, generating z-scores. Replicates were summarized for each feature by taking the median. We focused on three of the features: the number of cells per well, the mitotic index, and the average nuclear area. Overall a nonlinear trend was observed between cell count and the other two phenotypes (nuclear area, mitotic index). We calculated a model (using loess regression) to adjust for cell count effect on these phenotypes and computed z-scores from the residuals (Supplemental Table S1). The heatmap of z-scores in Figure 2A was generated using hierarchical clustering based on Euclidean distances.

#### **Combinatorial RNAi**

For the double-RNAi screen, we used a candidate-sensitizer design. We pipetted 125 ng (2.5  $\mu$ l) of each candidate dsRNA into a 384well clear-bottom microscopy plate (BD Falcon) as described. Then we added 125 ng (2.5  $\mu$ l) of one of the sensitizer dsRNAs using a NanoDrop II dispenser (Innovadyne), creating ~21,000 combinations (28 sensitizer vs. 750 candidate dsRNAs). Each pairwise combination of genes was assayed through four pairs of independent dsRNA designs: if we denote by A and A' the two independent designs targeting one gene and by B and B' the two independent dsRNA reagents targeting the other, then each biological replicate of the experiment contained the candidate-sensitizer combinations A-B, A-B', A'-B, and A'-B'. In each well, 6500 cells were seeded in 30  $\mu$ I of culture medium with 0.15  $\mu$ I of 0.4 mg/ml DDAB per well and incubated for 5 d at 25°C before fixation, staining, and imaging of cells (using the methods described for the genome-wide screen).

#### Mathematical modeling of synthetic genetic interactions

As described previously (Axelsson et al., 2011; Horn et al., 2011), we used a multiplicative model as the reference model (null model), which assumes that the double-RNAi phenotypic effect of noninteracting genes is equal to the product of the single-RNAi phenotypic effects. The single-RNAi phenotypic effects (main effects) for the 14 "query" genes were estimated by taking the median effect over all candidate genes (per phenotype). To prevent biases for the main-effect estimates of candidate genes, their main effects were estimated using measurements after the gene was codepleted with control dsR-NAs (targeting nonexpressed firefly luciferase). Pairwise interaction scores were then computed as the log ratio of the measured phenotype and the predicted phenotype (from the reference model), which is the product of the two single-knockdown effects. The significance of an interaction was assessed by calculating the deviation from zero from four interaction scores per gene pair (four independent co-RNAi experiments per gene pair). To this end, a moderated t test (R/Bioconductor package limma) was used, which first estimates the standard errors (SEM) by fitting a linear model through the four values, followed by empirical Bayes smoothing of the SEM (Smyth, 2004). The p values were adjusted for multiple testing by the method of Benjamini and Hochberg (1995) controlling the FDR.

# Calculation of the connection specificity index on the similarity between genetic interaction profiles

The CSI is based on a correlation matrix. For each pair of target genes, the PCC of the two genetic interaction profiles along all query genes was computed. The CSI of a gene pair A-B was then defined as the fraction of genes connected to A and B that have a PCC smaller than the PCC of A and B. A constant of 0.01 was applied in the CSI definition of Green *et al.* (2011).

#### Induction of G<sub>2</sub>/M checkpoint arrest in Drosophila S2 cells

To test the mitotic arrest after RNAi against genes selected from the primary screen, we triggered G<sub>2</sub>/M checkpoint arrest in *Drosophila* S2 cells 96 h past dsRNA transfection using 4  $\mu$ M doxorubicin or 20  $\mu$ M etoposide. After 6 h, cells were fixed and stained for their DNA content (Hoechst 33342) and for pH3, and their mitotic index phenotype was determined (for details see the Supplemental Methods).

### Fluorescence-activated cell sorting-based analysis of mitotic exit in human HeLa cells

Human HeLa cells were transfected with siRNA pools targeting the gene of interest and cultured for 3 d under standard conditions. A prometaphase arrest was induced by applying 40 ng/ml nocodazole for 18 h under standard conditions, and cells were released from this arrest and fixed at different time points up to 8 h postrelease. After propidium iodide staining, cell cycle profiles were determined using fluorescence-activated cell sorting analysis, counting 10,000 events/ sample (for details, see the Supplemental Methods).

#### Data access

Supplemental Tables S1–S7, Figures S1–S5, and Supplemental Material and Methods provide all interaction ( $\pi$ ) scores and p values.

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