Combining Genetic Perturbations and Proteomics to Examine Kinase-Phosphatase Networks in *Drosophila* Embryos

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SUMMARY

Connecting phosphorylation events to kinases and phosphatases is key to understanding the molecular organization and signaling dynamics of networks. We have generated a validated set of transgenic RNA-interference reagents for knockdown and characterization of all protein kinases and phosphatases present during early *Drosophila melanogaster* development. These genetic tools enable collection of sufficient quantities of embryos depleted of single gene products for proteomics. As a demonstration of an application of the collection, we have used multiplexed isobaric labeling for quantitative proteomics to derive global phosphorylation signatures associated with kinase-depleted embryos to systematically link phosphosites with relevant kinases. We demonstrate how this strategy uncovers kinase consensus motifs and prioritizes phosphoproteins for kinase target validation. We validate this approach by providing auxiliary evidence for Wee kinase-directed regulation of the chromatin regulator Stonewall. Further, we show how correlative phosphorylation at the site level can indicate function, as exemplified by Sterile20-like kinase-dependent regulation of Stat92E.

INTRODUCTION

Despite the ease with which we can identify protein phosphorylation, in the vast majority of cases, the protein kinase(s) or phosphatase(s) responsible for controlling any particular phosphorylation event is unknown. We sought to develop a proteomic strategy to easily and systematically screen for candidate protein kinase and phosphatase substrates in *Drosophila melanogaster* embryos, with the goal of identifying specific residues that these enzymes target in the context of development. *D. melanogaster* is an ideal model for the dissection of signaling mechanisms, as the majority of transcription in the embryo occurs after the midblastula transition (MBT), and thus, transcriptional feedback has relatively no impact on the phosphoproteome in early embryos. Additionally, since the embryo is a syncytium prior to cellularization at the MBT, distortions in phosphosite measurements due to contributions from multiple cell types can be avoided. However, acquiring sufficient material from mutant embryos for proteomic studies is a challenge. The classical technique to generate maternally deficient embryos—relying on the production of germline clones using the flipase (FLP) recombinase-mediated dominant female sterile technique (Chou and Perrimon, 1996)—is labor intensive, as it involves the construction of complex genotypes. Moreover, background mutations on the FLP-recognition-target-bearing chromosome can confound phenotype interpretation, and the approach does not typically yield enough material for proteomic studies.

Here, we describe how we have used genetic manipulation by transgenic RNA interference (RNAi) to derive sufficient quantities of embryos for phosphoproteomic analyses. RNAi is a well-founded method to analyze gene function in *D. melanogaster* (Perrimon et al., 2010), but the efficacy of RNAi during early embryogenesis has only recently been improved to enable robust gene knockdown during this developmental stage (Ni et al., 2011). By using the Gal4/UAS system (Brand and Perrimon, 1993) to temporally and spatially restrict expression of RNAi reagents, we confined protein kinase and phosphatase knockdown specifically to the germline. Using this strategy, we were able to query maternal gene function without affecting the viability of the animal, since an intact germline is dispensable for organismal development. We generated and validated a transgenic RNAi library that targets all protein kinases and phosphatases expressed in the *D. melanogaster* germline. Through rigorous characterization of our collection, we uncovered maternal-effect genes and verified previously implicated kinases and phosphatases in early *D. melanogaster* development. Furthermore, we systematically monitored global phosphoproteome alterations in kinase-deficient embryos for the purpose of illustrating how the method can generate lists of candidate kinase substrates. The approach illuminated kinase-dependent signaling and permitted the unbiased prediction of kinase consensus motifs that match kinase specificities previously characterized in vitro. As anticipated, the strategy identified downregulated phosphoproteins that include bona fide kinase substrates of the depleted kinase and an extensive list of candidate kinase-targeted substrates and phosphosites. We further
establish that two phosphosites consistently responding in the same direction (positive correlation) or the opposite direction (negative correlation) in different genetic contexts can illuminate phosphosite functionality. Given the extensive similarity between human and \textit{D. melanogaster} kinases, and the conservation of functional phosphorylation (Gnad et al., 2010; Landry et al., 2012), we anticipate that insight gained from our data and analyses will inform future mammalian studies.

\textbf{RESULTS}

\textbf{Compilation of the Maternally Inherited Protein Kinome and Phosphatome}

The \textit{D. melanogaster} genome encodes 32 tyrosine kinases, 237 serine/threonine kinases, and 112 protein phosphatases (Manning et al., 2002; Morrison et al., 2000). To systematically link protein phosphorylation sites with their cognate kinases and phosphatases in \textit{D. melanogaster}, we first identified the complement of kinase and phosphatase messenger RNAs (mRNAs) that are deposited maternally and contribute to the early zygote by analyzing developmental time course RNA sequencing (RNA-seq) data (Graveley et al., 2011). Using an RPKM (reads per kilobase of exon model per million mapped reads) cutoff of 3, determined by comparison to real-time quantitative PCR (qPCR) analysis of staged embryos (Hu et al., 2013b), we determined that 201 protein-kinase-encoding transcripts and 76 protein phosphatase-encoding transcripts (Figure 1A; Table S1 available online) are present during the first 4 hr of embryogenesis (stages 1–8). This accounts for 75% and 68% of all protein kinases and phosphatases, respectively, encoded in the \textit{D. melanogaster} genome (Figure 1A). We independently verified the presence of these transcripts by real-time qPCR (Figure 2A) but detected only 172 kinases and 67 phosphatases in 2-hr-old embryos (stages 1–4) at the protein level based on peptide MS1 feature intensities from shotgun mass spectrometry for the same developmental time. A total of 172 kinases and 67 phosphatases were quantified. Median signal-to-noise ratios observed across all matching peptides ranged from high (156: Cks30A) to low (5: CG7156) for kinases and from high (107: Pp2B-14D) to low (6: CG8147 and Ptp4E) for phosphatases.

Figure 1. Expression and Conservation of Protein Kinases and Protein Phosphatases during Early \textit{D. melanogaster} Embryogenesis

(A) Of 269 \textit{D. melanogaster} protein kinase-encoding genes, 201 were identified by RNA-seq between 0 and 4 hr of embryogenesis, while 76 of 112 protein phosphatase-encoding transcripts were identified for the same developmental window. Represented is an average RPKM value from two time points comprising stages 1–8. Undetected transcripts are those with an RPKM value less than 3. Average RPKM values ranged from high (257: polo) to low (3: btl, PVR, and CG43143) for kinases and from high (327: mts) to low (3: CG565 and CG16771) for phosphatases. Corresponding proteins, identified from MS2-based peptide fragmentation, were quantified based on label-free peptide MS1 feature intensities from shotgun mass spectrometry for the same developmental time. A total of 172 kinases and 67 phosphatases were quantified. Median signal-to-noise ratios observed across all matching peptides ranged from high (156: Cks30A) to low (5: CG7156) for kinases and from high (107: Pp2B-14D) to low (6: CG8147 and Ptp4E) for phosphatases.

(B) Conservation of expressed (outer ring) and undetected (inner pie) \textit{D. melanogaster} protein phosphatases during early embryogenesis (0–4 hr) to human and yeast. Conservation was considered when three or more ortholog predictions tools (DIOPT score > 3) predicted a high confidence ortholog. See also Figure S1.

\textbf{Generation and Validation of the Transgenic shRNA Collection Targeting Kinases and Phosphatases}

We previously demonstrated the utility of short hairpin RNAs (shRNAs) embedded in an endogenous microRNA scaffold to knock down maternal gene function in \textit{D. melanogaster} embryos. A side-by-side comparison of shRNA with long double-stranded
RNA (dsRNA) transgenic lines indicates that screening of shRNA lines triples the frequency of RNAi-derived germline phenotypes (Yan et al., 2014), generally due to higher expression of shRNAs in the germline (Ni et al., 2011). Having characterized the requirements for efficient gene knockdown during oogenesis, we sought to generate a complete and validated set of shRNA-expressing transgenic lines capable of targeting protein kinases and phosphatases that are contributed maternally to the developing embryo. To induce shRNA expression specifically in the female germline using the Gal4-UAS system, we crossed females heterozygous for a UAS shRNA and either MTD-Gal4 (a line bearing three copies of Gal4 expressed sequentially throughout oogenesis; Petrella et al., 2007) or tub-Gal4 (a line bearing two insertions of Gal4 expressed from a maternal tubulin promoter during mid- and late oogenesis; Staller et al., 2013) to shRNA-bearing males in order to recover fertilized eggs. We analyzed more than 450 transgenic lines expressing shRNAs targeting protein kinases and phosphatases (Table S2). We were unable to recover eggs from 12% of the lines crossed to MTD-Gal4, accounting for 46 kinases and 6 phosphatases implying that these genes are required for early oogenesis.

For those lines from which we could recover eggs, we determined by real-time qPCR, following the Minimum Information for Publication of Quantitative Real-Time PCR Experiments guidelines (Bustin et al., 2009), that more than half of the ~450 transgenic lines we analyzed generated greater than 60% knockdown of corresponding kinase or phosphatase mRNA levels in 0-4 hr embryos, relative to a control shRNA targeting enhanced green fluorescent protein (EGFP) (Figures 2A and S2A). We observed excellent correlation between knockdown at the mRNA and protein level, which was assessed by comparing mRNA levels assessed by real-time qPCR to immunoblots of a subset of proteins for which antibodies were available (Figure 2B). We were interested in determining the number of transgenic lines that would need to be considered to observe at least one achieving >60% knockdown of the targeted transcript. We found that, when considering two unique shRNAs targeting the same gene product, this occurs at a frequency of 86% (N = 81 pairs) (Figure S2B). These data suggest that generating two independent shRNA lines is usually sufficient for obtaining at least one line that confers adequate knockdown. Interestingly, many cases of poor knockdown can be attributed to shRNA targeting design. Specifically, our data indicate that shRNAs targeting the transcript coding sequence (CDS) are more effective at knockdown than those targeting 3’ untranslated regions (UTRs) (Figure S2C), possibly reflecting inaccuracies in UTR annotation (Hu et al., 2013b).

Notably, we found no correlation between the degree of knockdown and the level of corresponding transcript in untreated early embryos (Figure S2D). Furthermore, our data exhibit no bias toward the concentration of recovered RNA or the date of sample collection (Figures S2E and S2F). Taken together, our collection consists of at least one transgenic line that provides a minimum of 60% knockdown for each maternally inherited protein kinase and phosphatase.

Assessment of Transgenic shRNA Collection Phenotypes

Our shRNA-directed knockdown strategy recapitulated many documented maternal-effect phenotypes (Figure 3A). As
Figure 3. Embryonic Phenotypes Generated from shRNA-Mediated Knockdown of Maternally Contributed Protein Kinase and Phosphatases

(A) Cuticle phenotypes of embryos derived from maternal-Gal4>UAS-shRNA females crossed to UAS-shRNA males. Description of associated phenotypes can be found in Table S2.

(B) Frequency of observed embryonic phenotypes derived from maternal-Gal4/UAS-shRNA females crossed to UAS-shRNA males, from a total of 450 examined lines.

(C) Twenty-four pairs of shRNAs targeting the same gene and generating >60% knockdown were compared for qualitatively similar embryonic phenotypes. Four of the six cases of a differential phenotype can be explained by degree of knockdown.

See also Table S2.
Figure 4. Phosphoproteomes of Kinase-Deficient *D. melanogaster* Embryos Generated by Quantitative Mass Spectrometry and Isobaric Labeling with Tandem Mass Tags

(A) Strategy followed to identify differential phosphorylation between kinase shRNA and control shRNA embryos (see Supplemental Experimental Procedures for details).

(B) Relative phosphosite levels between kinase shRNA and control shRNA embryos. Plotted is the fold change relative to a control shRNA (white) for phosphosites found among all experiments. These 1,139 unique phosphopeptides meet stringent criteria in terms of isolation specificity and phosphosite assignment (see Experimental Procedures). The hierarchical 2D matrix is clustered based on a correlation distance metric using average linkage. Knockdown efficiencies are as follows: Cdk8, 87%; Cks30A, 85%; slik, 92%; wee, 81%; Tao, 91%; mei-41, 84%; tefu, 68%; Ikb1, 86%; Afg1, 92%; Bub1, 99%; grp, 79%; cg3608, 89%; Gprk2, 82%; cdc2r, 85%; gish, 58%; mos, 90%; Csk, 90%; Pak, 95%; Eip63E, 71%.

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expected, we observed anterior-posterior terminal defects following the disruption of terminal signaling, such as that resulting from knockdown of the receptor tyrosine kinase torso, the SHP2 phosphatase ortholog corkscrew, or the downstream kinase suppressor of ras, ksr (Figure S3A). Altogether, we observed maternal-effect phenotypes for ~18% of lines that achieved 60% or greater knockdown (Figure 3B; Table S2), representing approximately 33% and 18%, respectively, of protein kinases and phosphatases expressed during early embryogenesis. Of those protein kinases and phosphatases for which a maternal-effect phenotype has been reported, we observed the same qualitative phenotype as that described in the literature approximately 74% of the time (26/35 genes considering germline clone-derived embryos; 2/3 considering embryos derived from homozygous mutant mothers; Table S3A). Anomalies can likely be attributed to: (1) weak hypomorphic alleles generating a less severe phenotype than extensive knockdown; (2) insufficient knockdown by an shRNA to produce phenotypes generated by strong or null mutant alleles; or, (3) in the case of embryos derived from mutant mothers, an effect resulting from mutant somatic follicular cells. Despite the fact that protein kinases and phosphatases are among the best characterized classes of genes, we uncovered unappreciated phenotypes for approximately 40 of these enzymes, implying roles in oogenesis and early embryogenesis. Further, knockdown of an additional 12 predicted kinases and phosphatases resulted in oogenesis and maternal-effect phenotypes, warranting more extensive characterization. A searchable interface to query genes for individual transgenic lines, a description of their knockdown and embryonic phenotypes, and photos of cuticle preparations for those with maternal-effect phenotypes can be found at http://www.flyrnai.org/RSVP.html.

We addressed the possibility and frequency of shRNAs generating phenotypes as a result of off-target effects (OTEs) in two ways. First, we compared pairs of unique shRNAs targeting the same gene for similar phenotypes. Comparing 24 efficient targeting pairs, we found that 80% produced the same qualitative phenotype as that described in the literature approximately 74% of the time (26/35 genes considering germline clone-derived embryos; 2/3 considering embryos derived from homozygous mutant mothers; Table S3A). Anomalies can likely be attributed to: (1) weak hypomorphic alleles generating a less severe phenotype than extensive knockdown; (2) insufficient knockdown by an shRNA to produce phenotypes generated by strong or null mutant alleles; or, (3) in the case of embryos derived from mutant mothers, an effect resulting from mutant somatic follicular cells. Despite the fact that protein kinases and phosphatases are among the best characterized classes of genes, we uncovered unappreciated phenotypes for approximately 40 of these enzymes, implying roles in oogenesis and early embryogenesis. Further, knockdown of an additional 12 predicted kinases and phosphatases resulted in oogenesis and maternal-effect phenotypes, warranting more extensive characterization. A searchable interface to query genes for individual transgenic lines, a description of their knockdown and embryonic phenotypes, and photos of cuticle preparations for those with maternal-effect phenotypes can be found at http://www.flyrnai.org/RSVP.html.

We initially assessed the reproducibility of phosphoproteomic profiles generated from analysis of separate populations of control shRNA embryos. We utilized mass spectrometry and an isobaric labeling strategy (see the Protocol) that enables multiplexing and relative quantification between samples (see Supplemental Experimental Procedures). Since ~700 embryos constitute the amount of material (~1 mg protein) we chemically label, the phosphoproteomic profile is a representative average of phosphorylation in this population. Amine-reactive TMT Isobaric Mass Tags, identical in mass but differing in their isotopic distribution of atoms, permit the simultaneous spectral identification of unique reporter ions generated from MS2-based fragmentation of each tag from labeled peptides. We compared TMT reporter ion intensities and phosphopeptide identities from three TMT-labeled control shRNA embryo populations in two independent experiments (Table S4). When considering those phosphopeptides in the same multiplex experiment (10,166 phosphopeptides for one experiment and 8,032 for the other; see Supplemental Experimental Procedures for normalization and specific criteria), we generally identify the same phosphopeptide in all three biological replicates (~99% of the time). In the two independent experiments, we observed phosphopeptide levels deviating an average of 7% (Figure S3A) and 29% (Figure S3B) between three biological replicates. This indicates that variability in factors such as peptide labeling and embryo collection has little influence on our ability to consistently detect the majority of phosphopeptides.

A Resource Providing Accessibility for Proteomic Analyses and Kinase Characterization

Our current reagents for germline-specific RNAi make it relatively easy to obtain large numbers of eggs depleted of a single gene product. This allowed us to perform quantitative proteomic experiments to measure the global effect of each perturbation on the phosphoproteome (Figure 4A). We anticipated that a relative quantitative and global assessment of altered phosphorylation in protein kinase-deficient embryonic extracts could provide a list of putative protein kinase-substrate (KS) and phosphosite matches. We also reasoned that phosphorylation signatures could also be used to predict roles for protein kinases and phosphatases in specific biological processes and reveal functional redundancy.

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Given the reproducibility between control shRNA replicates, we extended our phosphoproteomic examination to embryos derived from females expressing efficient shRNAs (as determined by real-time qPCR) targeting 19 different protein kinases (Figure 4B). We were able to quantify nearly 8,500 unique phosphosites among 19 deficient kinase samples (Tables S5A–S5D). The number of unique phosphosites we quantified between experiments, ranging from 6,331 to 2,448, was based on the number of unique phosphopeptides identified per experiment, ranging between 22,942 and 12,201. Notably, 1,140 phosphosites were quantified in all 19 kinase knockdown conditions, 1,343 in ten kinase knockdown conditions, and 4,358 in five kinase knockdown conditions. The majority of phosphopeptides (C) Phosphoproteins with two or more downregulated phosphosites (>1.5-fold) were classified into four types based on the directionality of change of the majority of identified phosphosites for the same protein: type 1, most identified phosphosites do not change; type 2, most identified phosphosites are downregulated; type 3, all identified phosphosites are downregulated; and type 4, most identified phosphosites are upregulated. See also Figures S3 and S4.
in each kinase-deficient sample were unchanging in abundance experiment. In terms of candidate kinase-targeted phospho-
sites, either direct or indirect, we consider those downregulated sites with changes of 1.5-fold or greater in kinase-deficient em-
bryos relative to control embryos, since observed phosphopep-
tides for seven shRNA-targeted kinases—Wee, Tao, Atg1, Gilgamesh (Gish), Lkb1, Grapes (Grp), and Sterile20-like kinase (Silk)—minimally met this criterion (2.27-fold, 1.95-fold, 1.69-fold, 2.24-fold, 2.68-fold, 1.98-fold, and 2.1-fold, respec-
tively) in corresponding kinase-deficient embryos (Figure S4A). We did not detect phosphopeptides for the other 12 shRNA-tar-
geted kinases. Moreover, changes in the phosphorylation of known substrates of shRNA-targeted kinases approach this value; for instance, Histone H3, Med13, and Stat92E (downregu-
lation 2.2-fold, 1.8-fold, and 2.4-fold, respectively) in Cdk8-defi-
cient embryos and Cdk1, Klp61F, and Hsp83 (downregulated 1.4-fold, 2.1-fold, and 1.7-fold, respectively) in wee-deficient embryos. Indeed, for a third of the D. melanogaster orthologs of literate-curated Cdk8 substrates in yeast (Sharifpoor et al., 2011) we identified, one or more respective phosphopeptides were downregulated >1.5-fold in Cdk8-deficient embryos (Table S5E). Using this criterion, the number of downregulated phos-
sites in kinase-deficient profiles ranged from 22 (Bub1-defi-
cient embryos) to 752 (Cdk8-deficient embryos) (Table S5). Of note, while our Bub1-targeting shRNA generated efficient knock-
down (99% knockdown), Bub1-deficient embryos exhibited no morphological or hatch rate defects (Table S2), consistent with minimal effects on the phosphoproteome. Conversely, knock-
down of Cdk8, a cyclin-dependent kinase influencing transcription and cell cycle progression (Szilagyi and Gustafsson, 2013), resulted in dramatic and penetrant morphological and hatch rate phenotypes, consistent with extensive modulation of the observed phosphoproteome. We speculate that among those phosphosites downregulated >1.5-fold in the 19 kinase-deficient contexts we surveyed are sites directly targeted by the corre-
sponding depleted kinase(s), as well as indirect targets altered downstream of the manipulated kinase. For instance, in the case of gish-deficient embryos, we observed enrichment of downregulated phosphorylation of proteins involved in Hedge-
hog (Hh) and Wnt/Wingless (Wg) pathways (Table S5B), consist-
tent with a role for Gish in mediating Hh and Wg signaling (Davidson et al., 2005; Hummel et al., 2002). These data indicate that, by monitoring changes in the phosphoproteome, one can effectively screen for candidate substrates and alterations in signaling downstream of the targeted kinase. However, further scrutiny of any altered phosphosite is required to prove a KS rela-
tionship, as we demonstrate later.

In order to distinguish genuine kinase targets from phos-
phosite alterations due to protein instability, we classified phospho-
proteins with two or more downregulated phosphosites (>1.5-fold) into five categories based on the directionality of change of the majority of identified phosphosites for each indi-
vidual protein: type 1, the majority of phosphosites do not change; type 2, the majority of phosphosites are downregulated; type 3, all phosphosites are downregulated; type 4, most phos-
phosites are upregulated; and type 5, indistinguishable due to an equal distribution of unchanged, upregulated, or downregulated phosphosites (Figure 4G). In considering at least two phos-
sites, we increase the probability that the corresponding phospho-
proteins is indeed subjected to degradation and not merely reduced in phosphorylation at a single site. Type 1 and type 4 phosphoproteins are those for which we can reasonably discount the possibility of protein degradation as a mechanism of downregulated phosphorylation and, thus, are considered high-priority candidates for phosphorylation by the respective kinase. The observed downregulation of type 3 phosphoprote-
sins, on the other hand, can be explained by indirect mecha-
isms leading to protein degradation, such as altered protein-
protein interactions or phosphorylation-mediated degradation. Although most phosphoproteins in our data set are of type 1 (Fig-
ure 4G), type 3 phosphoproteins account for ~20%, on average, of those proteins with two or more downregulated phosphosites in each kinase depletion condition. This percentage is in line with previous reports that protein expression changes account for less than 25% of differential phosphorylation (Bodenmiller et al., 2010; Wu et al., 2011). We scrutinized respective trans-
cripts for type 2 and type 3 phosphoproteins in order to identify and filter our data set of potential OTEs due to partial comple-
mentarity of the targeting shRNA to unintended transcripts. By comparison to the frequency of partial complementarity of each targeting shRNA (seven-nucleotide match to seed) to the early embryonic transcriptome, we find a relatively weak proba-
bility for partial complementarity of our targeting shRNAs to 3’ UTRs or transcripts of type 2 phosphoproteins (see Supple-
mental Information). This probability declines when considering type 3 phosphoproteins, indicating that off-targets are not en-
riched in our data set and are therefore unlikely to explain alter-
ations observed in our analyses. To further substantiate this assumption, we proceeded to knock down respective trans-
scripts for type 3 phosphoproteins with the best matches to each corresponding kinase shRNA seed. Germline-specific knockdown of ten candidate off-targets predicted for six ki-
nase-targeting shRNAs failed to generate phenotypes that could explain specific phenotypes attributed to the corresponding kinase shRNA (see Supplemental Information).

Extracting Patterns in Phosphorylation Data Sets to Find KS Relationships

We speculated that we might be able to extract KS relationships and insight into signaling pathway connectivity from our phos-
phorylation data set as a whole by examining patterns in phos-
phoalterations among kinase-deficient contexts. For instance, since most kinases are activated by phosphorylation, correlative phosphorylation events observed between kinases and other proteins could be indicative of KS relationships. On the other hand, anticorrelatorative phosphorylation could be additionally informative; inhibitory phosphorylation of a kinase would always be out of phase with phosphorylation of that kinase’s respective tar-
gets. To explore such possibilities, we surveyed correlations in phosphorylation changes (>1.5-fold cutoff relative to a control shRNA) between identified phosphosite pairs among our kinase-deficient conditions (447,585 correlative pairs involving 2,058 phosphosites; Table S6A). When considering phosphosite pairs exhibiting positive or negative correlation in at least four kinase-deficient conditions (25,077 correlative pairs), we find enrichment for authentic KS pairs (Figure 5A), derived from 133,051 D. melanogaster KS pairs (Table S6B) predicted from
179 conserved human kinase phosphorylation motifs by NetPhorest (Miller et al., 2008) and from mapping of 517 gold standard KS pairs in yeast (Sharifpoor et al., 2011) to D. melanogaster. Enrichment for authentic KS pairs still exists when considering phosphosite pairs correlating in only two or three kinase-deficient conditions (Figure S5A). Strikingly, we also find enrichment for correlative phosphorylation among components of the same protein complex (p = 7.5 \times 10^{-157}), further substantiating how this phenomenon can be exploited to identify functionally relevant phosphosites.

While correlative analysis can clearly illuminate direct KS relationships in large-scale phosphorylation data, it can also provide functional information if one has a priori knowledge of the consequence of phosphorylation of one of the participating phosphosites. We exemplify this with the case of Silk and Stat92E. Phosphorylation of the Stat92E transcription factor at Tyr711 promotes DNA binding (Yan et al., 1996). We found that phosphorylation at this particular site positively correlates with phosphorylation of Silk at Ser1376 (Figure 5B), suggestive of a relationship between Silk and Stat92E; the probability of observing two phosphosites correlating among six kinase-deficient profiles is rare (p = 1.4 \times 10^{-5}). We predicted that Silk activates Stat92E given that reduced Stat92E phosphorylation in silk-deficient embryos (Figure 5C) cannot be explained by instability of Stat92E protein (Figure 5B). Indeed, Stat92E target gene expression was downregulated in silk dsRNA-treated embryos (Figure 5D). Insulin has been reported to enhance growth hormone-induced Stat activation in mature adipose cells (Zhang et al., 2013), and Stat may be a direct target of the insulin receptor (Sawka-Verhelle et al., 1997). We confirmed an increase in the activating phosphorylation of Stat92E in cells treated with insulin (Figure 5C). Remarkably, we observed that more than a quarter of phosphoproteins downregulated in silk-deficient embryos are upregulated in cells in response to insulin, including Silk (Figure 5C; Figures S5E and S5F). Moreover, 30% of phosphoproteins downregulated >1.3-fold in silk-deficient embryos (Table S7) were found to physically interact with components of the insulin-signaling network (Glatter et al., 2011). These observations suggest that Silk could be activating Stat92E via insulin signaling. Consistent with this, we observed a reduction in activated Akt1 in silk-deficient embryos, despite elevated total Akt1 protein (Figure 5E). A reduction in insulin signaling may, in fact, explain the longevity of silk^−^ mutant larvae (Hipfner and Cohen, 2003). Raf interaction has been suggested to bridge Silk to the MAPK proliferation branch of cell survival signaling (Hipfner and Cohen, 2003), which our data support, as we find that...
slik-deficient embryos exhibit defects in ERK activation (Figure SSD). Despite a nonessential role for slik in embryogenesis, our examination of coregulatory phosphorylation during this early stage illuminated Silk function, highlighting the power of our approach.

**An Examination of Wee-Dependent Phosphorylation**

We chose to examine more closely the phosphoproteomic profile of RNAi-derived wee kinase-deficient embryos, since their phenotype mirrored that reported for mutant wee embryos (Price et al., 2000). Wee, Cdk1, and Aurora operate in a regulatory kinase cascade to control nuclear divisions in the early embryo. Phosphorylation and activation of Aurora by Cdk1 is inhibited by Wee and delays entry into mitosis. Wee inhibits Cdk1 by phosphorylating a conserved tyrosine (Tyr15) located in the ATP binding pocket (Campbell et al., 1995; Stumpf et al., 2004). Therefore, we expected Cdk1 and Aurora to be hyperactive in the absence of Wee. Indeed, we find motif enrichment (Figure 6A) among phosphate-enriched phosphoproteins that are enriched among phosphoproteins altered >1.5-fold in wee-deficient embryonic lysates relative to control. Motif-X was used to identify motifs (Chou and Schwartz, 2011). The PLogo tool was used to generate motif logos. Favoring amino acids at corresponding positions are indicated above the black line, while disfavored amino acids are below. "0" indicates the site of phosphorylation.

**Figure 6. Phosphoproteomic Characterization of wee-Deficient Embryos**

(A) Indicated are motifs encompassing phosphosites that are enriched among phosphosites altered >1.5-fold in wee-deficient embryonic lysates relative to control. Motif-X was used to identify motifs (Chou and Schwartz, 2011). The PLogo tool was used to generate motif logos. Favoring amino acids at corresponding positions are indicated above the black line, while disfavored amino acids are below. "0" indicates the site of phosphorylation.

(B) Levels of a Cdk1 Tyr15 encompassing phosphopeptide in wee-deficient embryos relative to control embryos (w, white) as determined by TMT reporter ion signal (right) from the corresponding peptide identified by MS2 fragmentation (left, MS2 spectra). The hashtag indicates the localized site of phosphorylation (p < 0.05). Indicated is a representative peptide.

(C) Of 308 phosphoproteins identified as Cdk1 substrates in yeast (Holt et al., 2009), we mapped 120 to fly with a DIOPT score ≥ 1. Half of the orthologous D. melanogaster counterparts exhibit altered phosphorylation (>1.3-fold) in wee-deficient embryos.

(D) Approximately half of those phosphosites upregulated >1.3-fold in wee-deficient kinases can be attributed to Cdk and the downstream kinase Aurora based on kinase consensus motif matching.

(E) Gene Ontology Consortium term enrichment among altered phosphoproteins (>1.5-fold) in wee shRNA embryos relative to control embryos, identified using the DAVID Functional Annotation Tool.

(F) Levels of a Stwi Tyr305 encompassing phosphopeptide in wee-deficient embryos relative to control embryos (w, white) as determined by TMT reporter ion signal (right) from the corresponding peptides identified by MS2 fragmentation (left, MS2 spectra). The hashtag indicates the site of phosphorylation (p < 0.05).
upregulated phosphosites in wee-deficient embryos that resembles Cdk and Aurora kinase consensus motifs (Cdk1: pS/T-P-X-K/R; pS/T-P-X-X-K) and Aurora: R-R/K-pS/T; R/K-X-pS/T; R/R- K-X-pS/T) (Alexander et al., 2011). Accordingly, we consistently observed less TMT reporter ion signal proportionate to levels of K-X-pS/T) (Alexander et al., 2011). Wee kinase in these wee-deficient embryos. Of these, we observed hemizygous embryos (Stumpff et al., 2004). Aurora is also hyperactive in wee-deficient embryos, reflected by the upregulation in phosphorylation of characterized targets: kinesin-like protein at 10A (Klp10A pSer210: 2.5-fold), inner centromere protein (Incenp pSer163: 1.5-fold and pSer164: 3-fold), and histone H3 (HH3 pSer10: 15-fold; pSer28: 7-fold) (Adams et al., 2001; Jang et al., 2009; Kang et al., 2001). We verified HH3 phosphoalterations in wee-deficient embryos by immunoblotting (Figure S6A). Surprisingly, half of the upregulated phosphosites we identified in wee kinase-deficient embryos reside within sequence recognized by Cdk1 or downstream Aurora kinase (Figure 6D). This observation highlights the utility of phosphoproteomic signatures to reveal genetic epistasis. We also find enrichment for specific Gene Ontology Consortium categories for those phosphoproteins regulated by Wee (Figure 6E). As anticipated, we observed enrichment for cell cycle classified factors, particularly those with mitosis-specific functions such as nuclear migration, spindle organization, and chromosome segregation. Intriguingly, proteins with roles in chromatin assembly are overrepresented in our list of upregulated phosphoproteins in wee-deficient embryos. This is interesting, given the reported hypocondensation of mitotic chromatin in wee null embryos (Stumpff et al., 2004). Motif enrichment among downregulated phosphorylations is illustrated (Figure S6B). Another indirect consequence of wee knockdown is the upregulation of Stat92E phosphorylation at Tyr711 in wee-deficient embryos (Table S5A). Cdk1 has been shown to regulate Stat92E phosphorylation at Tyr711 in cells (Baeg et al., 2005), and indeed, we detect elevated Y711-encompassing Stat92E phosphopeptides in wee-deficient embryos that cannot be attributed to increased Stat92E levels (Figure S5B). Wee functions as a preserved tyrosine kinase (Campbell et al., 1995; McGowan and Russell, 1993); therefore, we inquired as to whether any phosphoproteins for which tyrosine phosphorylation was reduced in wee-deficient embryos were attributable to increased Stat92E levels (Figure S5B). Wee Kinase

Figure 7. Identification of Stwl as a Target of Wee Kinase

(A) Lysates from Drosophila cells expressing HA-tagged Wee together with 3xFLAG-tagged candidate Wee substrates were subjected to immunoprecipitation with anti-FLAG antibody and analyzed by immunoblotting with the indicated antibodies. (B) Lysates from Drosophila cells expressing HA-tagged Wee together with 3xFLAG-tagged Stwl were subjected to immunoprecipitation with anti-phosphotyrosine antibody and analyzed by immunoblotting with the indicated antibodies. (C) Recombinant GST-Stwl fusion proteins were incubated with human WEE1 kinase and radio-labeled ATP and analyzed by SDS-PAGE and autoradiography. Histone H2B serves as a positive control (lane 4). The migration of input proteins is indicated with asterisks. Autophosphorylated WEE1 migrates at 120 kDa. (D) Lysates from 0–2 hr embryos derived from females expressing shRNAs targeting wee, stwl, or an EGFP control shRNA were analyzed by immunoblotting with anti-Stwl and anti-Wee antibodies. Immunoblotting with anti-tubulin serves as a loading control. (E) Lysates from 0–2 hr embryos derived from females expressing shRNAs targeting wee, stwl, or an EGFP control shRNA were analyzed by immunoblotting with antibodies recognizing different histone H3 posttranslational modifications.
We generated a validated collection of transgenic *D. melanogaster* shRNA lines targeting protein kinases and phosphatases maternally deposited in embryos. The collection permits the examination of zygotic lethal gene perturbations, without the effort of germline clone derivation. Multiple lines of evidence support that the embryonic phenotypes generated by our collection are, indeed, a result of shRNA on-targeting: (1) near-identical qualitative phenotypes generated by two unique shRNAs targeting the same gene for the 15% of the collection we tested; (2) abolition of shRNA-induced phenotypes by substitution of three nucleotides (C911s) precluding on-target binding; (3) the high degree of overlap between our shRNA-derived phenotypes and literature-reported mutant embryo and germline clone-derived embryo phenotypes; and (4) our general inability to accredit specific shRNA phenotypes to candidate OTEs derived from proteomics and partial complementarity matching.

**A General Method to Predict Kinase Motifs and Targets**

Using our shRNA collection, we performed quantitative phosphoproteome assessments of genetically compromised animals. An advantage of our gene knockdown strategy over gene knockout is that we restrict RNAI to the germline: since germline development is dispensable for organismal development, our RNAI method likely avoids major adaptation and compensation due to effects on the viability of the animal, such as that seen, for example, with yeast deletion mutants (Bodemiller et al., 2010; Teng et al., 2013). Additionally, the modest amount of transcription in early-stage embryos further minimizes the possibility of compensation at the transcriptional level, although nontranscriptional compensation is a possibility. Conceivably, by comparing genetic knockout to incomplete depletion by RNAI-mediated knockdown, one could identify compensatory rewiring events. From phosphoproteomic profiling of kinase-deficient embryos, we identified altered phosphorylation of characterized substrates of depleted kinases and generated an extensive list of candidate substrates of the depleted kinase and altered phosphoproteins targeted by downstream kinases. A challenge will be to distinguish between primary and secondary targets. It is difficult to evaluate the number of primary targets per kinase since this will depend on multiple factors, including the function of the kinase and its expression level, localization, and connectivity with other proteins. Indeed, studies from yeast and mammalian kinases have illustrated that the number of substrates for any one kinase can range from hundreds to only a few (Übersax and Ferrell, 2007). Thus, we expect variability in the number of substrates depending on the analyzed kinase. Furthermore, biologically meaningful alterations in phosphorylation may have been missed in our analyses, given the limitations of current mass spectrometry technology. We illustrate, however, that current instrumentation can be used to identify known and predicted targets relevant to the function of the perturbed kinase (e.g., Stwl is a direct target of Wee kinase) and so, despite perhaps only scratching the surface, we have generated biologically pertinent information. Additional information (e.g., in vitro kinase activity toward a substrate, protein-protein interaction, and functional assays) is necessary, of course, to infer a direct KS relationship (Sopko and Andrews, 2008). Undoubtedly, extension of our methodology will be effective for systematically mapping substrates to culpable kinases and for pinpointing critical phosphosites important for substrate function.

**Correlation and Anticorrelation: An Application for Network Analysis**

Our correlative analysis examining coordination between alterations in phosphosite pairs among kinase-deficient profiles...
uncovered signaling mechanisms. We demonstrate how a role for Slik kinase in regulating the transcription factor Stat92E could be predicted from correlative phosphorylation of these two proteins. The predictive power of this approach could be extended by knowing if specific phosphorylation events serve activating or inhibitory functions and by superimposing kinase consensus motifs. Our analysis demonstrates how functional phosphorylation might be uncovered in any phosphoproteomic data using simple correlative principles. Notably, predictions for any particular kinase can be made indirectly from its detection in varying genetic contexts, with no requirement for direct modulation of the queried kinase. The data we generated from embryos will complement orthogonal data sets such as kinase consensus motif and protein-protein interaction data derived from, for example, peptide and protein chip assays, coaffinity purifications, and yeast two-hybrid assays. Furthermore, phosphosite correlation information could be integrated with large-scale RNAi phenotype data in order to predict whether phosphorylation of a target by a specific kinase serves an activating or inhibiting function.

Perspective
Given that key signaling pathways and kinases implicated in human disease are conserved in Drosophila (Rubin et al., 2000), the insight gained from our kinase-deficient phosphoproteomic signatures constitutes an important step toward understanding the kinome network. Going forward, we anticipate that phosphoproteomic assessment of other posttranslational modifications and more complex genotypes, using combinatorial knockdown (two shRNAs) or knockdown in combination with transgene overexpression or gain-of-function mutations, will appreciably illuminate our ability to decipher signaling mechanisms. In this way, global proteomic analyses could map pathways but also reveal critical nodes in signaling that may partially or completely overcome mutations resulting in pathway hyperactivity. Alternatively, phosphoproteomic assessment of a sensitized kinase mutant in the context of a substrate gain of function could expose altered signaling mechanisms contributing to compromised viability (Sopko et al., 2006). Finally, genetic combinations would mimic more natural scenarios in terms of genetic heterogeneity contributing to susceptibility to disease and, by mapping contextual phosphorylation, would improve on our ability to predict and target essential signaling nodes.

EXPERIMENTAL PROCEDURES
Detailed methods are available in the Supplemental Experimental Procedures. Mass spectrometric sample preparation is further described in the Protocol.

Transgenic shRNA Line Generation
shRNAs (21 base pairs) were cloned into VALIUM series vectors and injected into embryos for targeted phiC31-mediated integration at genomic attP landing sites on the second or third chromosome as described elsewhere (Ni et al., 2011). All transgenic lines were sequenced to confirm the identity of the shRNA and miR-1 scaffold.

Protein Kinase or Phosphatase-Deficient Embryo Derivation
Females heterozygous for the UAS-shRNA and either MTD-Gal4 (Petrella et al., 2007), expressing three versions of Gal4 sequentially throughout oogenesis, or tub-Gal4, a line expressing Gal4 from a maternal tubulin promoter at two insertion sites during mid- and late oogenesis (Stailer et al., 2013), were crossed to UAS-shRNA males to recover fertilized embryos.

RNA Isolation, Reverse Transcription, and Real-Time qPCR
RNA was isolated using guanidinium thiocyanate-phenol-chloroform extraction using TRIzol (Life Technologies) and glass-bead-based cell disruption. Genomic DNA was eliminated by incubation with DNase (QIAGEN), and samples were processed for cleanup with an RNaseasy MiniElute Cleanup Kit (QIAGEN). One microgram of purified RNA was incubated with a mix of oligo(dT) and random hexamer primers and with iScript RT (iScript cDNA Synthesis Kit, Bio-Rad) for complementary DNA (cDNA) synthesis. cDNA was used as the template for amplification, using validated primers in iQ SYBR Green Supermix with a CFX96 Real-Time PCR detection system (Bio-Rad). Query gene expression was relative to a control sample, normalized to the expression of three reference genes: ribosomal protein L32, alpha-tubulin, and either nuclear fallout or Gapdh1, using the ΔΔC(t) analysis method.

Maternal Phenotype Derivation
Hatch rate was calculated from counting embryos 24 hr after deposition. For genotypes with defective hatching, cuticles prepared in Hoyer’s mounting media were imaged with a Zeiss AxioCam HRC Camera mounted on a Zeiss Axiophot microscope.

Commmunoprecipitation and Immunoblotting
Lysates were subjected to immunoprecipitation using the indicated antibodies, and samples were subjected to SDS-PAGE followed by immunoblotting.

Quantitative Phosphoproteomics
Embryos lysed in 8 M urea were digested with trypsin, and peptides were chemically labeled with one of six TMT Isobaric Mass Tags (Thermo Fisher Scientific), separated into 12 fractions by strong cation exchange chromatography, purified with TiO2 microspheres, and analyzed via liquid chromatography-tandem mass spectrometry on an Orbitrap Velos Pro mass spectrometer (Thermo Fisher Scientific). Peptides were identified by Sequest and filtered to a 1% peptide false discovery rate (FDR). Proteins were filtered to achieve a 2% final protein FDR (final peptide FDR near 0.15%). TMT reporter ion intensities for individual phosphopeptides were normalized to the summed reporter ion intensity for each TMT label. The localizations of phosphorylations were assigned using the AScore algorithm.

In Vitro Kinase Assay
In vitro kinase assays were carried out as described elsewhere (Sopko et al., 2006).

Correlative Analysis
A phosphosite matrix was constructed where rows correspond to identified phosphosites and columns correspond to kinase-deficient data sets. Only phosphosites with ≥0.58 log2 fold change were distinguished, by values +1 and −1, based on an increase or decrease, respectively, in levels relative to an shRNA control. All pairwise combinations of phosphosites were classified as positive or negative correlating based on their change in the same or opposite direction, respectively, for each kinase-deficient condition. A correlation sign score was determined, considering the number of positive and negative correlations and the total number of kinase-deficient phosphorylation profiles where both phosphosites change.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Supplemental Experimental Procedures, six figures, seven tables, and a Protocol and can be found with this article online at http://dx.doi.org/10.1016/j.devcel.2014.07.027.

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