Phosphoproteomics Reveals Distinct Modes of Mec1/ATR Signaling during DNA Replication

Graphical Abstract

Highlights
- A combined genetic-proteomic approach to monitor Mec1 and Tel1 signaling
- Mec1 is highly activated during normal DNA replication
- Replication-correlated Mec1 action is distinct from canonical checkpoint
- 9-1-1 clamp and Dna2 lagging-strand factor activate Mec1 during DNA replication

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In Brief
Mec1/ATR kinase activity is typically associated with checkpoint signaling in response to replication stress and S-phase DNA damage. Bastos de Oliveira et al. use a quantitative mass spectrometry approach (QMAPS) to demonstrate that Mec1 has “replication-correlated” activity that is distinguishable from its action during canonical checkpoint responses.

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Phosphoproteomics Reveals Distinct Modes of Mec1/ATR Signaling during DNA Replication

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SUMMARY

The Mec1/Tel1 kinases (human ATR/ATM) play numerous roles in the DNA replication stress response. Despite the multi-functionality of these kinases, studies of their in vivo action have mostly relied on a few well-established substrates. Here we employed a combined genetic-phosphoproteomic approach to monitor Mec1/Tel1 signaling in a systematic, unbiased, and quantitative manner. Unexpectedly, we find that Mec1 is highly active during normal DNA replication, at levels comparable or higher than Mec1’s activation state induced by replication stress. This “replication-correlated” mode of Mec1 action requires the 9-1-1 clamp and the Dna2 lagging-strand factor and is distinguishable from Mec1’s action in activating the downstream kinase Rad53. We propose that Mec1/ATR performs key functions during ongoing DNA synthesis that are distinct from their canonical checkpoint role during replication stress.

INTRODUCTION

During DNA replication, cells are prone to accumulate genomic instabilities. Progression of the replication machinery is often impeded by barriers such as DNA adducts, DNA-RNA hybrids, and protein-DNA complexes (Lambert and Carr, 2013). Replication forks often stall upon encountering these hard-to-replicate regions, leading to exposure of single-stranded DNA (ssDNA), which, in turn, is a major signal for the activation of the evolutionarily conserved PI3K-like sensor kinase ATR (yeast Mec1) (MacDougall et al., 2007). Once activated, ATR and Mec1 initiate a signaling response that induces key effects such as cell-cycle arrest, inhibition of origin firing, and stabilization of stalled replication forks (Branzei and Foiani, 2010; Santocanale and Diffley, 1996). The importance of ATR is highlighted by the fact that deleterion or mutations that affect its activity are associated with embryonic lethality, chromosomal fragmentation, and increased sensitivity to genotoxic drugs (Brown and Baltimore, 2000; Wright et al., 1998). In budding yeast, strains with mec1 mutations were shown to accumulate gross chromosomal rearrangements (GCRs) (Myung et al., 2001) and be exquisitely sensitive to genotoxic drugs that induce replication stress (Weinert et al., 1994). Like ATR, the PI3K-like sensor kinase ATM (yeast Tel1) is also important during DNA damage responses. Cells lacking ATM show sensitivity to DNA-damaging agents and elevated levels of mitotic recombination (Meyn, 1993), but differently from ATR, which is a sensor for ssDNA accumulation, ATM responds mainly to DNA double-strand breaks (DSBs) (Shiloh and Ziv, 2013). In yeast, tel1Δ mutants are viable and show no significant sensitivity to DNA-damaging agents. However, mec1Δtel1Δ double mutants are more sensitive to replication stress and display a more severe growth defect than the single deletion mutants, revealing functionally redundant roles for these kinases (Morrow et al., 1995).

Over the last decade, others and we have identified many candidate substrates of Mec1/Tel1 and ATR/ATM using large-scale mass spectrometry (MS)-based approaches (Chen et al., 2010; Matsuoka et al., 2007; Smolka et al., 2007). However, our understanding of how these kinases promote a systemic cellular response that safeguards genomic integrity and allows cells to better cope with the effects of replication stress is still limited. A major limitation toward a more comprehensive characterization of Mec1/Tel1 and ATR/ATM action is posed by the difficulty of reproducibly and quantitatively monitoring the many substrates identified by MS. Consequently, the use of antibody-based approaches to monitor well-established substrates remains the method of choice. Substrates commonly monitored using western blotting techniques include the histone variant H2AX (yeast H2A) and the downstream checkpoint kinases CHK1 and CHK2 (yeast Rad53). Despite the biological relevance of these substrates, the use of their phosphorylation as readouts for the checkpoint response has introduced a marked bias in studies aiming at characterizing Mec1/Tel1 action. To address this problem, here we employed a combined genetic-proteomic approach (which we refer to as quantitative mass-spectrometry...
analysis of phospho-substrates [QMAPS] for identifying and monitoring multiple in vivo kinase substrates in a systematic, unbiased, and quantitative manner. Using QMAPS, we show that Mec1 is robustly activated during unperturbed DNA replication, in a manner that correlates with the extent of DNA replication and that is distinct from a canonical checkpoint. Collectively, our results demonstrate the importance of unbiased and quantitative analysis of kinase substrates to comprehensively characterize the in vivo action of multi-functional kinases.

**RESULTS**

**Unbiased Delineation of Mec1 and Tel1 Action Using a Genetic-Proteomic Approach**

Our current understanding of Mec1 and Tel1 action is biased toward the use of a few established substrates as reporters of the in vivo activity of these kinases. In particular, the activation state of the major downstream kinase Rad53 has been extensively used as a key indicator of Mec1 and Tel1 activation status. To circumvent this bias and be able to comprehensively characterize the action of Mec1 and Tel1, we used quantitative MS analysis of kinase mutant strains to identify and monitor as many candidate substrates of these kinases as possible. First, we performed a proteomic screen to globally define the set of Mec1 and Tel1 candidate targets. Building on our previously published work (Smolka et al., 2007), we used quantitative MS to compare the phosphoproteome of wild-type (WT) and mec1Δ tel1Δ cells treated with methyl methanesulfonate (MMS) or hydroxyurea (HU) to induce replication stress. To facilitate the classification of Mec1/Tel1-dependent phosphorylation sites into direct or indirect Mec1/Tel1 phosphorylation events, we also quantified the relative abundance of the phosphopeptides in cells lacking Rad53, the major kinase downstream of Mec1/Tel1. We were able to identify and quantify more than 6,000 phosphopeptides over distinct biological replicates (Figure 1A; Table S1). Of interest, the abundance of 232 of the identified phosphopeptides was significantly reduced in cells lacking Mec1 and Tel1, and we refer to them as Mec1/Tel1-dependent events. Among the 232 Mec1/Tel1-dependent targets, 115 were found to be dependent on Rad53, and thus considered as indirect Mec1/Tel1-dependent events (Figure 1B). In our strategy, direct targets of Mec1/Tel1 should be present in the group of phosphopeptides carrying a Mec1/Tel1-dependent and Rad53-independent phospho-site. As shown in Figure 1C (Table S1), analysis of the amino acid in the +1 position of Mec1/Tel1-dependent and Rad53-independent phospho-sites revealed a strong enrichment of the S/T-Q motif, consistent with previous work indicating this preferential motif for Mec1 and Tel1 (Kim et al., 1999; Smolka et al., 2007). Of the 117 Mec1/Tel1-dependent and Rad53-independent phosphorylation events, 97 are in the preferred S/T-Q motif, and we considered them as directly targeted by Mec1 or Tel1. On the other hand, Rad53 showed a bias toward the S/T-bulky amino acid (ø) motif (Figure 1B; Table S1). For more than 60% of the proteins found to have a Mec1/Tel1-dependent phosphorylation, we were able to also detect at least one Mec1/Tel1-independent phosphorylation event, supporting that most of the observed changes are not due to changes in protein abundance (Figure S1; Table S1).

To sort out the relative contribution of Mec1 or Tel1 in the response, we performed similar analyses as described above, but comparing WT cells to cells lacking either Mec1 or Tel1 (Figure 1D). Of the Mec1 and Tel1 direct phospho-events identified above, 67% were found to heavily depend mostly on Mec1 (Figure 1E; Table S1). Only four phospho-sites were found to heavily depend exclusively on Tel1, consistent with the fact that cells lacking Tel1 don’t exhibit significant sensitivity to replication stress-inducing agents (Morrow et al., 1995). Importantly, about 29% of Mec1/Tel1-dependent sites were found to remain robustly phosphorylated in cells lacking either Mec1 or Tel1 and represent a set of common candidate substrates of these kinases (Figures 1D–1F; Table S1). These results establish a large set of Mec1 and Tel1 targets and define their relative level of dependency for each of these kinases. This defined set of phosphorylation sites targeted by Mec1 and/or Tel1 forms the basis of our unbiased strategy to characterize the action of these kinases in different growth conditions and genetic backgrounds. The output of this analysis of substrates is a quantitative map, herein named QMAPS, revealing the relative levels of phosphorylation of identified phosphopeptides in two different conditions being tested (see Figure 2A).

**QMAPS Reveals Robust Activation of Mec1 during Normal DNA Replication**

It is currently accepted that activation of Mec1 is strongly induced by replication stress. This notion is mainly based on the fact that HU-induced replication fork stalling leads to a robust activation of Rad53 (Tercero et al., 2003). To test if our unbiased QMAPS approach could reveal new insights into the action of Mec1 or Tel1, we compared the phosphorylation level of Mec1/Tel1 candidate substrates in cells undergoing normal S-phase with cells treated with HU. In both cases, cells were arrested in G1 with α-factor and then released from the arrest in media containing HU or not for 45 min. As shown in Figure 2A and Table S2, nearly all phosphopeptides carrying a Rad53-dependent phosphorylation site were induced by HU. Unexpectedly, only a minor fraction of Mec1 and/or Tel1 candidate substrates was induced by HU treatment. This fraction included a phosphorylation site in Rad53 (serine 24) and a phosphorylation in the Mrc1 protein (serine 189), the adaptor known to transduce signals from Mec1 to Rad53 in response to HU. Most phosphorylation events in Mec1 and/or Tel1 targets were either only slightly induced by HU or did not change at all when comparing cells going through normal replication with cells experiencing HU-induced replication stress. Remarkably, a Mec1 autophosphorylation site (serine 38) and phosphorylation of Rfa1 and Rfa2 (serines 178 and 122, respectively), which are highly dependent on Mec1, were in fact inhibited by HU. Targeted analysis of purified Mec1 complexes further confirmed that the Mec1 auto-phosphorylation site and phosphorylation of Rfa1 are indeed induced during normal S-phase and accumulate as more DNA is replicated, following a similar trend observed for the acetylation of H3K56, which is a well-established replication mark (Figure 2B) (Masumoto et al., 2005). To test if Mec1 activation in normal S-phase is dependent on DNA replication, we used QMAPS to compare the phosphorylation levels of its targets in WT cells as well as in cells lacking the S-phase cyclins Clb5.
and Clb6, which display delayed replication initiation due to delayed CDK activation but undergo normal budding dynamics as they progress through S-phase (Figures 2C and 2D) (Donaldson et al., 1998). As shown in Figure 2D, several Mec1 candidate substrates are highly induced during S-phase in WT cells but are not induced in clb5Δclb6Δ cells at the 35 min time point, when only limited DNA replication had occurred in the mutant (Figures 2C and 2D; Table S2). Taken together, these results show that Mec1 action in normal S-phase depends, at least partially, on DNA replication. While the MS analysis could detect many Mec1/Tel1-dependent phosphopeptides in G1 in clb5Δclb6Δ cells (Figure 2D), we attributed this basal phosphorylation level to the potential accumulation of these phosphopeptides in the extended and deregulated S-phase from the previous cell cycle. Very few Rad53-dependent phosphopeptides were detected in the absence of drug-induced replication stress (data not shown). Even in cells lacking three phosphatases known to act on Rad53, namely Ptc2, Ptc3, and Pph3 (Heideker et al., 2007), we were not able to detect robust Rad53 action during a normal S-phase as we still identified a very limited set of
targets (Figure S2). Nonetheless, we were able to observe an increase in the level of phosphorylation of the detected Rad53 targets in \( ptc2\Delta ptc3\Delta pph3\Delta \) triple mutant cells compared to WT cells, suggesting that phosphatases play a role in counteracting Rad53 activation during normal DNA replication.

Collectively, the QMAPS results shown in Figure 2 reveal that Mec1 is robustly activated during normal DNA replication and that this mode of Mec1 signaling is partially uncoupled from Rad53 activation. On the other hand, HU-induced replication stress leads to an increase in the phosphorylation of most targets.
Rad53 targets but to minor changes in the phosphorylation of a large fraction of Mec1 targets, or even inhibition of some of them. We therefore propose that Mec1 can operate in two distinct modes of signaling during DNA replication, one correlated with ongoing DNA synthesis ("replication-correlated") and another correlated with the extent of replication stress that involves strong Rad53 activation (canonical checkpoint response).

The 9-1-1 Clamp and the Lagging-Strand Factor Dna2 Are Important for "Replication-Correlated" Mec1 Activation

Recent work revealed that activation of the Mec1 kinase in response to replication stress or DNA damage requires the action of factors such as Ddc1, Dna2, and Dpb11, all of which possess an unstructured region that can tether Mec1 for activation (Kumar and Burgers, 2013; Navadgi-Patil and Burgers, 2009; Puddu et al., 2008). To test if the replication-correlated mode of Mec1 action also requires these factors for activation, we used QMAPS to compare phosphorylation of substrates in WT and mutants of Mec1-activating factors. As shown in Figure 3A and Table S3, mutation of two residues (W128A and Y130A) in Dna2 previously shown to be required for the ability of Dna2 to activate Mec1 has mild effects on the ability of Mec1 to target some of its specific targets, such as Rfa1, Spt7, and Dad1. Deletion of DDC1 had almost no effect in most targets (Figure 3A; Table S3), suggesting that Dna2 has a more prominent role in activating Mec1 during normal DNA replication. Importantly, deletion of DDC1 also prevents the recruitment of Dpb11 and its ability to activate Mec1 (Navadgi-Patil and Burgers, 2009). Finally, combination of the dna2 WY-AA mutation (herein referred as dna2-AA) with DDC1 deletion had a significant impact on the phosphorylation levels of targets that highly depend on Mec1, suggesting that Dna2 and Ddc1 function redundantly to activate Mec1 during normal DNA replication (Figure 3A; Table S3). This is consistent with the fact that these proteins are known to localize and function on the lagging strand of the replication fork. These results suggest that Mec1 may be activated mostly at the lagging strand of a moving replication fork during normal DNA replication.

Tel1 Phosphorylates a Specific Group of Mec1 Targets to Prevent GCR and Support Robust DNA Replication in the Absence of Mec1

Analysis of GCRs revealed that activation of Mec1 via Dna2 or Ddc1 during replication becomes particularly important in the absence of Tel1, as shown by the dramatic increase in GCR in tel1Δddc1Δdna2-AA cells (Figure 3B). This result highlights the key role of Tel1 in compensating for the loss of Mec1 during normal DNA replication. Consistent with this data, while...
**Dna2 and Ddc1 Are Not Essential for Activation of the Canonical Mec1-Rad53 Signaling Response Following Replication Stress**

To determine the extent in which Dna2 and Ddc1 are necessary for activation of the canonical Mec1-Rad53 response during replication stress, we performed QMAPS analysis comparing WT cells versus *ddc1Δdna2-2-AA* cells treated with MMS, which leads to robust Rad53 activation. As shown in Figure 4A (Table S4), *ddc1Δdna2-2-AA* cells exhibit strong defects in Mec1 activation during MMS treatment, but unexpectedly, activation of Rad53 under this condition does not seem to be greatly affected. On the other hand, similar QMAPS analysis comparing WT and *mec1Δ* cells revealed a strong impact in the phosphorylation of Rad53 targets in the absence of Mec1. These results show that *ddc1Δdna2-2-AA* cells do not phenocopy *mec1Δ* cells regarding Rad53 activation and suggest the existence of additional factors that may activate Mec1 to specifically activate Rad53, consistent with a recent paper (Bandhu et al., 2014). In support of the idea of additional Mec1 activator(s), *ddc1Δdna2-2-AA* cells are not as sensitive to MMS or HU as *mec1Δ* cells (Figure 4B). Also, while *mec1Δ* and rad53Δ cells are well known to require deletion of the ribonucleotide reductase inhibitor SML1 for viability (Zhao et al., 1998), we found that *ddc1Δdna2-2-AA* cells do not require SML1 deletion for viability (Figure 4C). Of note, even *tel1Δddc1Δdna2-2-AA* cells do not require SML1 deletion for viability, despite these cells showing the dramatic increase in GCR rates that is characteristic of *mec1Δtel1Δ* cells. We could exclude the possibility of a Ddc1-independent role for Dpb11 in the activation of Mec1 under MMS in *ddc1Δdna2-2-AA* cells as removal of the C-terminal region of Dpb11, which is required for its ability to activate Mec1 (Navadgi-Patil and Burgers, 2008), did not cause loss of viability or major growth defect (Figure 4D). As shown in the working model in Figure 4E, we propose two distinct modes of Mec1 action during DNA replication, one correlated with DNA replication and another correlated with the extent of replication stress as part of a canonical checkpoint signaling. In our model, the replication-correlated mode of Mec1 action functions redundantly with Tel1 to ensure robust DNA replication and prevent GCR. On the other hand, the canonical checkpoint mode leads to the well-established effects of inhibition of DNA replication and increased production of dNTPs.

**DISCUSSION**

The ATR and ATM kinases, and their yeast orthologs, regulate hundreds of substrates, but our ability to fully capture their multi-functional action in vivo has been hampered by the common use of one or a few classical substrates as readouts of their activity. Here we used a quantitative MS approach to monitor in vivo Mec1/Tel1 kinase action in a systematic, unbiased, and quantitative manner. Our analysis revealed surprising insights into how Mec1 functions during DNA replication and provided evidence of a non-canonical mode of Mec1 action, which we propose is distinct from Mec1’s established role in checkpoint signaling (see model in Figure 4E).

By quantitatively monitoring the phospho-status of dozens of Mec1 candidate substrates, we found that Mec1 is highly active during normal DNA replication. In fact, genetic data support the idea that Mec1 functions during normal DNA replication. For example, cells lacking *MEC1* and *TEL1* exhibit high rates of GCR in an assay performed in the absence of any exogenously-induced DNA damage (Myung et al., 2001). But the prevailing hypothesis has been that the ability of Mec1 to suppress spontaneous GCR accumulation is attributed to a residual action of Mec1 in response to spontaneous DNA damage generated during DNA replication. Distinct from the notion of residual Mec1 activation during normal replication, our work supports a model in which Mec1 is highly engaged onto sites of ongoing DNA synthesis to become activated in a “replication-correlated” manner. Also, distinct from the established role of Mec1 in checkpoint signaling, our results reveal that the action of Mec1 during normal DNA replication is partially uncoupled from the action of the downstream kinase Rad53. Our results are consistent with the idea that Mec1 is either continuously activated during ongoing DNA synthesis or is activated at many sites in the genome that pose moderate level of difficulty for replication forks to pass. At these sites, forks would only dynamically pause, allowing sufficient ssDNA exposure for Mec1 recruitment and activation but not for robust Rad53 activation, which requires further recruitment and/or phosphorylation of mediator proteins to mount a full checkpoint response. Nonetheless, it is important to mention that Rad53 also needs to be activated during normal DNA replication. Cells lacking Rad53 are not viable, unless the RNR inhibitor SML1 is also deleted (Zhao et al., 1998). But contrary to Mec1’s action, our quantitative analysis reveals that the activity of Rad53 in normal DNA replication is significantly lower...
than drug-induced Rad53 activity (Figure 2A). We speculate that during normal DNA replication, Rad53 becomes preferentially activated at specific genomic sites that pose major challenges for replication, such as hard-to-replicate transcriptional barriers. Interestingly, our results suggest that phosphatases such as Pph3, Ptc2, and Ptc3 may also function during normal S-phase to prevent excess Rad53 activation, consistent with a recent report showing a constitutive Mec1-Pph3 interaction (Hustedt et al., 2015).

The identification of a replication-correlated mode of Mec1 action leads to a paradox, as Rad53 has established roles in inhibiting DNA synthesis as part of a canonical checkpoint response to replication stress (Santocanale and Diffley, 1998). We hypothesize that Mec1 positively regulates DNA replication when functioning uncoupled from Rad53 activation in the replication-correlated mode (Figure 4E). Consistent with this hypothesis, the Bell lab has shown that Mec1 phosphorylates the MCM complex to prime it for activation (Randell et al., 2010). We further speculate that the replication-correlated mode of Mec1 signaling plays a major role in facilitating the movement of replication forks by preemptively opening chromatin and/or removing RNA and transcriptional machineries from DNA.
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Consistent with this notion, we found that Mec1 targets several proteins involved in transcription, RNA processing, and chromatin remodeling during unchallenged DNA replication. Also, we showed that during normal DNA replication Tel1 partially compensates for the lack of Mec1 by targeting substrates involved in transcription and chromatin regulation. The fact that cells lacking both Mec1 and Tel1 are extremely slow growing further strengthens the idea that the set of Mec1 substrates that can also be phosphorylated by Tel1 comprise a critical set of proteins involved in promoting robust DNA replication. Previous reports have functionally connected Mec1 to chromatin and transcription regulation (Rodriguez and Tsukiyama, 2013; Seeber et al., 2013). Our work suggests that regulation of these processes by Mec1 is actually part of the normal replication program that positively controls ongoing DNA synthesis. The delineation of which substrates are common to Mec1 and Tel1 should provide the framework of targets that will help better understand the mechanisms by which Mec1 and Tel1 positively impact DNA synthesis. Finally, the observation that replication-correlated mode of Mec1 and Tel1 action does not efficiently relay signaling to Rad53 activation is consistent with these ideas, as it is well known that Rad53 activation leads to outputs that would antagonize the potential role of Mec1 as a positive regulator of DNA replication.

EXPERIMENTAL PROCEDURES

Cell Culture

Yeast strains used in this study are listed in Table S5. For stable isolate labeling of amino acids in cell culture (SILAC), auxotrophic yeast strains for lysine and arginine were grown in -Arg -Lys synthetic dropout media supplemented with either normal L-arginine and L-lysine (light culture) or L-lysine13C6, 15N4 and arginine were grown in -Arg -Lys synthetic dropout media supplemented with either normal L-arginine and L-lysine (light culture) or L-lysine13C6, 15N4 (heavy culture) as described in Ohouo et al. (2013). Our work suggests that regulation of these processes by Mec1 is actually part of the normal replication program that positively controls ongoing DNA synthesis. The delineation of which substrates are common to Mec1 and Tel1 should provide the framework of targets that will help better understand the mechanisms by which Mec1 and Tel1 positively impact DNA synthesis. Finally, the observation that replication-correlated mode of Mec1 and Tel1 action does not efficiently relay signaling to Rad53 activation is consistent with these ideas, as it is well known that Rad53 activation leads to outputs that would antagonize the potential role of Mec1 as a positive regulator of DNA replication.

Phosphopeptide Enrichment

Phosphopeptide enrichment was performed as described in (Ohouo et al., 2013). See Supplemental Experimental Procedures for further details.

MS Analysis

Phosphopeptides were subjected to LC-MS/MS analysis using a Q-Exactive Orbitrap or an Orbitrap XL mass spectrometer. See Supplemental Experimental Procedures for further details.

ACCESSION NUMBERS

The MS proteomics data have been deposited to the Peptide Atlas database (http://www.peptideatlas.org/) with the data set identifier PASS00651 and PASS00652.

SUPPLEMENTAL INFORMATION

Supplemental Information includes three figures, five tables, and Supplemental Experimental Procedures and can be found with this article online at http://dx.doi.org/10.1016/j.molcel.2015.01.043.

AUTHOR CONTRIBUTIONS

F.M.B.d.O., D.K., and J.R.C. performed experiments. K.H.S. and L.D. provided reagents and performed the GCR analysis. J.D., M.C.J., and H.Y. designed the bioinformatics tools for the phosphoproteomics data analysis. F.M.B.d.O., D.K., and M.B.S. designed the study, performed data analysis, and wrote the paper.


In the original published version of the above article, the indication of the sites mutated in the dna2-AA mutant is incorrect. Instead of “mutation of two residues (W352A and Y544A),” the authors should have written “mutation of two residues (W128A and Y130A).” A similar error occurred in the article’s corresponding Supplemental Information file. The article and its Supplemental Information file have now been corrected. The authors apologize for any inconvenience this error has caused.