A novel mechanism for the prevention of transcription replication conflicts

Berta Canal, Alba Duch, Francesc Posas & Eulàlia de Nadal

To cite this article: Berta Canal, Alba Duch, Francesc Posas & Eulàlia de Nadal (2018) A novel mechanism for the prevention of transcription replication conflicts, Molecular & Cellular Oncology, 5:3, e1451233, DOI: 10.1080/23723556.2018.1451233

To link to this article: https://doi.org/10.1080/23723556.2018.1451233

© 2018 The Author(s). Published with license by Taylor & Francis Group, LLC

Accepted author version posted online: 28 Mar 2018.
Published online: 28 Mar 2018.

Article views: 256
A novel mechanism for the prevention of transcription replication conflicts

Berta Canal*, Alba Duch*, Francesc Posas, and Eulàlia de Nadal

Cell Signaling Research Group, Departament de Ciències Experimentals i de la Salut, Universitat Pompeu Fabra (UPF), Barcelona, Spain

ABSTRACT

Transcription and replication complexes can coincide in space and time. Such coincidences may result in collisions that trigger genomic instability. The phosphorylation of Mrc1 by different signaling kinases is part of a general mechanism that serves to delay replication in response to different stresses that trigger a massive transcriptional response in S phase. This mechanism prevents Transcription-Replication Conflicts and maintains genomic integrity in response to unscheduled massive transcription during S phase.

Transcription-Replication Conflicts (TRCs) are a major cause of genomic instability. Cells have evolved a wide range of strategies to prevent TRCs, whereby they limit potential head-on collisions between the transcription and replication machineries, which are the events most likely to trigger recombination events and genomic instability. Eukaryotic cells focus their strategies for the protection of highly transcribed regions either by the use of non-nucleosomal DNA-bound protein structures that cause a polar block of DNA replication or by locating replication and transcription factories in temporarily and spatially separated chromatin domains. However, how cells cope with sudden and unscheduled outbursts of transcription during S phase is an unclear question.

Cells are continuously exposed to changing environmental conditions that challenge their survival. Specific signaling pathways are activated to cope with such cellular stresses by the induction of a global cellular response to maximize cell survival. Stress-Activated Protein Kinases (SAPKs) are key for this response to stress. Hog1, the yeast counterpart of the mammalian p38, is a prototypical SAPK. Hog1 is activated upon osmостress and leads to the regulation of many cellular functions including the transcription of hundreds of stress-responsive genes. Such massive activation of gene expression, although crucial for the osmoadaptive response, represents an obstacle to the replication machinery during duplication of the genome. Remarkably, the stress-activated Hog1 phosphorylates Mrc1, a protein of the replication complex, to temporarily block DNA replication. The Hog1-mediated phosphorylation sites in Mrc1 (Thr169, Ser215 and Ser229) are located in the N-terminal domain and are different to the Mrc1 sites phosphorylated by Mec1 in response to DNA damage. Cells carrying the Hog1-non-phosphorylatable allele of MRC1 (mrc13A) bypassed the replication block and displayed a dramatic increase in Transcription-Associated Recombination (TAR) and genomic instability. Therefore, when cells activate transcription upon osmостress, the concomitant transient block of DNA replication is critical in order to guarantee cell survival.

In addition to its role in osmостress, Mrc1 also plays a key role in maintaining genomic integrity in response to other stresses that trigger the Environmental Stress Response (ESR) transcriptional signature. The N-terminal phosphorylation of Mrc1 at the Hog1 sites is essential for blocking S phase upon heat, oxidative or low glucose stresses, which also involve the massive transcriptional response featured by the ESR. Correspondingly, mrc13A cells bypass the S phase block in response to these stresses and display TAR and genomic instability. An unbiased kinome screen identified Mpk1, Psk1 and Snf1 as the kinases that phosphorylate Mrc1 upon heat, oxidative and low glucose stresses respectively. Thus, cells use canonical stress-signaling pathways to phosphorylate a common substrate, Mrc1, to protect genomic integrity during unscheduled massive transcription in S phase. Interestingly, the role of Mrc1 is not restricted to environmental cues and it also responds to internal stresses, such as slow growth or genomic instability, that also trigger the ESR response. Overall, Mrc1 plays a key role in protecting genomic integrity in scenarios that compromise cell survival, either due to environmental insults or as a result of abnormal cell fitness. This “Mrc1 transcription-replication safeguard mechanism (MTR)” protects genomic integrity when outbursts of transcription occur during S phase (Fig. 1).

Identification of the MTR highlights the necessity for cells to possess a dedicated mechanism for the prevention of TRCs that can eventually occur in the genome when there is an unscheduled massive induction of transcription. Such unscheduled transcription differs from transcriptions that operate at particular sites in the genome, or at particular genes that are scheduled...
to be induced during the S phase of the cell cycle. Future work will allow further exploration of whether the function of Mrcl is relevant beyond stress and whether it plays a pivotal role in preventing collisions and genomic instability when regular transcription takes place during S phase.

Of note, Mrcl has two independent functions that are necessary for maintaining genomic integrity. First, Mrcl is a protein of the Replication Complex that is essential for achieving efficient fork speed during DNA replication. Second, Mrcl also has a crucial function in activation of the DNA damage checkpoint in response to replication stress. Protection of genomic integrity by N-terminally phosphorylated Mrcl is a third independent function of Mrcl since mrcl1Δ cells replicate efficiently under normal conditions and are proficient in activating the DNA damage checkpoint pathway in response to DNA damage and replication stress.

If the MTR safeguard mechanism were to be conserved in higher eukaryotes, it could be crucial not only for avoiding collisions and genomic instability in cells exposed to environmental changes (e.g., epithelial cells that suffer temperature changes; renal cells that support high osmolarity, etc.) but it might also play an important function in pre-cancerous cells, which display genomic instability that is sensed as an internal stress. Furthermore, a high percentage of tumorigenic cells display mutations in the DNA damage checkpoint, resulting in an important obstacle to chemotherapy treatment. Thus, the discovery of a novel regulatory pathway that blocks the cell cycle independently of known checkpoint pathways opens a new landscape of possible new compounds that might block cell cycle progression.

Disclosure of potential conflicts of interest
No potential conflicts of interest were disclosed.

Funding
BC is a recipient of an FPI fellowship. The FP and EdN’s laboratory is supported by grants from the Spanish Ministry of Economy and Competitiveness (BFU2015-64437-P and FEDER, BFU2014-52125-REDT, and BFU2014-51672-REDC to FP, BFU2017-85152-P and FEDER to EdN), the Catalan Government (2017 SGR 799), the Fundación Botín, and by the Banco Santander through its Santander Universities Global Division to FP. FP is a recipient of an ICREA Acadèmia (Generalitat de Catalunya). The department is supported by Unidad de Excelencia María de Maeztu, MDM-2014-0370.

ORCID
Francesc Posas http://orcid.org/0000-0002-4164-7076
Eulàlia de Nadal http://orcid.org/0000-0003-0039-5607

References

Figure 1. The “MTR safeguard mechanism” protects genomic integrity in response to stress-induced outbursts of transcription during S phase. Cells induce a conserved transcriptional signature known as the ESR upon environmental stresses such as osmo (NaCl), heat (37°C), oxidative (H2O2) or low glucose as well as to mutations that reduce cell fitness. (A) Mrcl is phosphorylated at the N-terminus by multiple signaling kinases (e.g. Hog1, Mpk1, Psk1 or Sfn1), which delays replication to maintain genomic integrity upon stress. (B) The unphosphorylatable mutant of Mrcl (mrc13A) fails to delay replication and accumulates TAR and genomic instability. (A) Mrc1 is phosphorylated at the N-terminus by Hog1, Mpk1, Psk1 or Sfn1, which delays replication to maintain genomic integrity upon stress. (B) The unphosphorylatable mutant of Mrcl (mrc13A) fails to delay replication and accumulates TAR and genomic instability due to collisions between RNA and DNA polymerases. The role of Mrcl in the MTR is fully independent of the DNA damage checkpoint pathway.

Figure 2. Mrc1 is a fork barrier component whose phosphorylation in response to DNA replication stress activates Rad53. Genes Dev. 2003;17:1755–67. doi:10.1101/gad.1098303. PMID:12865299