RESEARCH ARTICLE

MRN- and 9-1-1-Independent Activation of the ATR-Chk1 Pathway during the Induction of the Virulence Program in the Phytopathogen *Ustilago maydis*

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Abstract

DNA damage response (DDR) leads to DNA repair, and depending on the extent of the damage, to further events, including cell death. Evidence suggests that cell differentiation may also be a consequence of the DDR. During the formation of the infective hypha in the phytopathogenic fungus *Ustilago maydis*, two DDR kinases, Atr1 and Chk1, are required to induce a G2 cell cycle arrest, which in turn is essential to display the virulence program. However, the triggering factor of DDR in this process has remained elusive. In this report we provide data suggesting that no DNA damage is associated with the activation of the DDR during the formation of the infective filament in *U. maydis*. We have analyzed bulk DNA replication during the formation of the infective filament, and we found no signs of impaired DNA replication. Furthermore, using RPA-GFP fusion as a surrogate marker of the presence of DNA damage, we were unable to detect any sign of DNA damage at the cellular level. In addition, neither MRN nor 9-1-1 complexes, both instrumental to transmit the DNA damage signal, are required for the induction of the above mentioned cell cycle arrest, as well as for virulence. In contrast, we have found that the claspin-like protein Mrc1, which in other systems serves as scaffold for Atr1 and Chk1, was required for both processes. We discuss possible alternative ways to trigger the DDR, independent of DNA damage, in *U. maydis* during virulence program activation.

Introduction

When faced with DNA damage, eukaryotic cells activate DNA damage response (DDR) pathways that help to preserve genome integrity and cell viability. Depending on the manner, extent, and cellular context of the DNA damage, the outcomes of DDR signaling range from transient cell cycle arrest coupled with DNA repair to apoptosis or senescence. In parallel, an increasing number of reports show the involvement of DDR programs in cell differentiation, aside from their more conservative role of protecting genome integrity [1]. The most obvious example is the vertebrate adaptive immune system, which requires the programmed induction and subsequent repair of double strand breaks (DSB) during antigen receptor gene
rearrangements [2]. A second well-known example, in fungi, is the switching of mating-type in budding and fission yeast, which involves a programmed DSB followed by DNA repair [3, 4]. In other occasions, the induction of DDR in response to non-programmed DNA damage can be used to activate alternative differentiation processes, such as neurite outgrowth in neuronal stem cells [5], or as the transition between yeast and hyphal growth in Schizosaccharomyces japonicus [6]. Perhaps the most extreme case supporting a role of DDR in developmental programs is the use of elements from the DDR cascade to control cell cycle regulation during a differentiation process in the absence of any observed DNA damage. In the embryos from the nematode Caenorhabditis elegans, differential cell cycle duration starts at the two-cell stage, when the larger anterior blastomere AB divides before the smaller posterior blastomere P1. Interestingly, the delay in P1 cells relies on two conserved elements from DDR cascade, Atl-1 (ATR) and Chk-1 (Chk1), and apparently there is no DNA damage associated with the activation of the DDR cascade during this process [7]. In the same way, during the differentiation of mammalian trophoblast stem (TS) cells, Chk1 prevents cell cycle exit and thereby premature differentiation of TS cells, in the absence of induced DNA damage [8].

Owing to the intricate connections between cell cycle regulation and differentiation processes [9], as well as to the ability of DDR cascade to regulate the cell cycle, it is tempting to hypothesize that DDR cascade may be recruited to modulate cell cycle during developmental processes, even if no DNA damage signal exists; most likely, this would occur through alternative ways of DDR activation. In the present work we aimed to test this hypothesis by studying it during the virulence program of the phytopathogenic fungus Ustilago maydis.

The activation of the virulence program in the corn smut fungus U. maydis involves the mating of a pair of compatible haploid budding cells, which results in an infectious dikaryotic hypha that grows on the plant surface until finding a suitable place to penetrate the plant tissue [10]. A peculiar characteristic of the U. maydis dikaryotic filament is the sustained G2 cell cycle arrest while growing on the plant surface [11, 12]. This cell cycle arrest is a requisite for the virulence in U. maydis. Mutant strains unable to arrest the cell cycle cannot effectively infect plants, because their ability to differentiate specific infection structures—the appressoria—is severely impaired [13]. As described in other systems, the response to DNA damage in U. maydis is mediated by the DDR kinase Chk1, activated by the upstream kinase Atr1 [14]. Previous research from our group showed that Chk1 was necessary for the cell cycle arrest establishment observed during the formation of the infective filament [15]. Moreover, during this process, Chk1 is activated via phosphorylation by Atr1 at the same residues which need to be phosphorylated in response to DNA damage [16]. Our work strongly suggests that the differentiation process during the virulence program in U. maydis involves the Atr1-Chk1 axis. The formation of the infectious hypha in U. maydis and the induced cell cycle arrest are triggered by the expression of a transcriptional master regulator called b-factor [17]. How this transcription factor can induce the activation of the Atr1-Chk1 cascade in order to arrest the cell cycle is unknown. Here we report our attempts to determine whether DNA damage is associated with the induction of the virulence program in U. maydis, as well as to define additional elements of the Atr1-Chk1 cascade involved in the control of the cell cycle arrest occurring in the infective filament.

Materials and Methods

Strains and growth conditions

U. maydis strains were derived from FB1 and FB2 genetic backgrounds [18] and are listed in Table 1. Cells were grown in rich medium (YPD), complete medium (CMD) or minimal medium (MMD) [19]. FACS analyses were described previously [20].
Plasmid and strain constructions

Plasmid pGEM-T easy (Promega) was used for cloning, subcloning and sequencing of fragments generated by PCR. The oligonucleotides are described in Table 2. To construct the different strains, transformation of *U. maydis* protoplasts with the indicated constructions was performed following published procedures [21]. Deletion of *mre11*, *rec1* and *mrc1* genes was done by gene replacement [22]. Briefly, a pair of DNA fragments flanking the corresponding ORF were amplified and ligated to antibiotic resistance cassettes via SfiI sites. The 5' and 3' fragments were amplified using the oligonucleotide pairs respectively (Table 2). Each fragment was about 1 kbp in length. Integration of the disruption cassette into the corresponding loci was verified in each case by diagnostic PCR and subsequent Southern blot analysis.

For C-terminal fusion of proteins to fluorescent markers, the adaptation of the SfiI-dependent gene replacement strategy for C-terminal tag [23] was used. To produce Rfa1–3GFP, 5' and 3' fragments were digested with SfiI and ligated to a cassette carrying a triple GFP-encoding gene. Chk1-GFP fusion was already described [14].

RNA analysis

Total RNA was extracted with acidic phenol solution. After extraction, the RNA was cleaned using the High Pure RNA Isolation Kit (Roche Diagnostics GmbH). For qRT-PCR, cDNA was synthesised using the High Capacity cDNA Reverse Transcription Kit (Applied Biosystems) employing 1 µg total RNA per sample. qRT-PCR was performed using the SsoAdvanced

Table 1. *U. maydis* strains used in this study.

| Strain | Relevant Genotype | Reference |
|--------|-------------------|-----------|
| FB1    | a1 b1             | [19]      |
| FB2    | a2 b2             | [19]      |
| AB33   | a2 Pnar1:bW2 Pnar1:bE1 | [23]      |
| AB34   | a2 Pnar1:bW2 Pnar1:bE2 | [23]      |
| UMC19  | a2 Pnar1:bW2 Pnar1:bE1 cbx1::Pnar:cdk1AF-myc-cbx | [16]      |
| UMC20  | a2 Pnar1:bW2 Pnar1:bE1 cbx1::Pnar:cdk1-myc-cbx | [16]      |
| UMT007 | a1b1 rfa1-GFP     | This study|
| UMT011 | a2 Pnar1:bW2 Pnar1:bE1 rfa1-GFP | This study|
| UMT012 | a2 Pnar1:bW2 Pnar1:bE2 rfa1-GFP | This study|
| UMP210 | a1b1 Δrec1       | [39]      |
| UMP211 | a2b2 Δrec1       | This study|
| UMP219 | a1b1 Δmre11      | [39]      |
| UMP220 | a2b2 Δmre11      | This study|
| UMT010 | a2b2 mrc11-914-HA | This study|
| UMT009 | a2b2 mrc11-914-HA | This study|
| UMP111 | a1b1 chk1-3GFP   | [15]      |
| UMT002 | a1b1 chk1-3GFP Δrec1 | This study|
| UMT019 | a1b1 chk1-3GFP Δmre11 | This study|
| UMT014 | a1b1 chk1-3GFP mrc11-914 | This study|
| UMP121 | a2 Pnar1:bW2 Pnar1:bE1 Pdik6:NLS-GFP | [16]      |
| UMT005 | a2 Pnar1:bW2 Pnar1:bE1 Pdik6:NLS-GFP Δrec1 | This study|
| UMT015 | a2 Pnar1:bW2 Pnar1:bE1 Pdik6:NLS-GFP Δmre11 | This study|
| UMT016 | a2 Pnar1:bW2 Pnar1:bE1 Pdik6:NLS-GFP mrc11-914 | This study|

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Universal SYBR Green Supermix (BioRad) in a CFX96 Real-Time PCR system (BioRad). Reaction conditions were as follows: 3 min 95°C followed by 40 cycles of 10 sec 95°C/10 sec 60°C/30 sec 72°C.

**Plant infections**

Pathogenic development of wild type and mutant strains was assayed by plant infections of the maize (Zea mays) variety Early Golden Bantam (Olds seeds) as described before [24].

**Microscopy**

Images were obtained using a Nikon Eclipse 90i fluorescence microscope with a Hamamatsu Orca-ER camera driven by Metamorph (Universal Imaging, Downingtown, PA). Images were further processed with Adobe Photoshop CS software.
Results

ATR-Chk1 activation is not triggered by impaired bulk DNA replication

The formation of the infectious dikaryotic hypha in *U. maydis* depends on an intricate transcriptional program that primarily involves a transcriptional regulator called b-factor [17]. The production of this master regulator is linked to the mating process that, after cell fusion, leads to the interaction of the two subunits composing the b-factor (bW and bE), each subunit provided by each mating partner. In the laboratory it is possible to bypass the requirement for a mating process for production of the infective filament by using the specialized strain AB33 and its control strain AB34. These strains harbor the compatible bE1 and bW2 and non-compatible bE2 and bW2 genes under the control of the nitrate-inducible *nar1* promoter, respectively [25] (Fig 1A). Induction of bE1/bW2 in the AB33 strain growing in medium with nitrate results in the formation of monokaryotic infective filaments that resemble the infectious hypha formed after fusion of compatible haploid cells, including the cell cycle arrest in G2 phase [15].

Previous studies have analyzed the *U. maydis* transcriptome in AB33 cells in conditions of expression of b-factor (i.e. forming the infective filament) and found that mRNA levels of several genes involved in DNA replication decrease upon *b*-expression [26]. Among these genes were those encoding components of the replication machinery such as um04529 (*pol1*, DNA-directed DNA polymerase α); um03501 (*top2*, DNA topoisomerase II); um01008 (*pol2*, a subunit of the DNA polymerase ε); and um06402 (*mcm7*). In addition, the expression of two of the components of the ribonucleotide reductase (um06368 and um11750, encoding for the small and large subunits, *rrn2* and *rrn1* respectively), which is required for the synthesis of dNTPs, appears severely down-regulated. The decrease in the levels of any of these proteins would affect the ability of the cells to replicate their DNA, and it could be a source of signals to activate DDR in *U. maydis* during the formation of the infective filament. Therefore we decided to address whether cells accumulate in the S-phase at some point during the induction of the infective filament. For that we monitored the DNA content using FACS analysis of cultures of AB33 and AB34 strains for each hour after the induction of the expression of b-factor (Fig 1B). No accumulation of S-phase cells was detectable at these times just before the arrest, suggesting that DNA replication was not compromised during the formation of the b-dependent filament.

Strikingly, our conclusion seems to be contradictory with the published observation of a down-regulation of genes involved in DNA replication upon *b*-expression [27]. However, down-regulation of these genes could be merely a consequence of the b-induced cell cycle arrest. Therefore, we analyzed the expression of the down-regulated genes in conditions of *b*-expression and non-arrested cell cycle. For that we used a *U. maydis* strain simultaneously expressing the genes encoding the b-factor as well as an ectopic Cdk1 allele refractory to inhibitory phosphorylation at Tyr15 (cdk1AF), the ultimate cause of the b-induced G2 cell cycle arrest [15, 28]. In this strain, the cdk1AF allele (and a control wild-type cdk1 allele) was expressed under the *nar1* promoter. As a consequence, in spite of the activation of the b-dependent transcriptional program, the cell cycle was not arrested [15, 28]. We have found, in agreement with a previous report [27], that for all the analyzed genes, the levels of mRNA dramatically decreased upon *b*-expression. However, this decrease seems to be a consequence of the cell cycle arrest: interference with the b-induced cell cycle arrest upon expression of the cdk1AF allele prevented the decrease in the mRNA levels in all cases (Fig 1C).

ATR-Chk1 activation seems not to be due to massive DNA damage

Having no evidence supporting the idea of a global replication collapse as responsible for activating the Atr1-Chk1 axis during the induction of the b-dependent filament, we searched for
any signal of DNA damage during this process. In a previous report [15], we had tried to use the presence of Rad51-GFP foci as a surrogate marker for the presence of DNA damage [29]. Using this approach we found no evidence of DNA damage during the induction of the b-dependent filament. However, since Rad51 acts in only a subset of responses to DNA damage, we sought to use an alternative way to detect any DNA damage signal. The appropriate DNA substrates for checkpoint initiation can be generated by several pathways, but all of them have in common the production of various types of single-stranded DNA (ssDNA) regions that are bound by the single-strand binding protein RPA [30]. RPA-coated ssDNA is instrumental in the recruitment of checkpoint complexes, and can be detected as foci using either immunofluorescence or fluorophore-coupled alleles of components of this protein complex [31]. For this purpose, we constructed a GFP-tagged allele of rfa1, encoding one of the subunits from the RPA complex in U. maydis [32]. A wild-type strain carrying the endogenous rfa1 gene tagged with GFP showed the presence of nuclear foci in response to DNA damage agents, such as

![Diagram of gene cassettes expressing compatible (AB33) or non-compatible (AB34) b-factor genes.](image)

![Graph showing FACS analysis of DNA content of AB34 and AB33 strains.](image)

![Graph showing quantitative real-time PCR for indicated genes.](image)
hydroxyurea (HU) and methyl methanesulfonate (MMS) treatment (Fig 2A). We introduced the \textit{rfa1-GFP} allele into the AB33 strain and checked for the presence of nuclear foci during the induction of \textit{b-dependent} filaments. The activation of Chk1 during the formation of the infective filament is transient, with its maximum achieved within 4 hours from \textit{b-factor} induction [15]. Therefore, we analyzed the presence of RPA foci during this period. Strikingly, we did not observe a higher frequency of RPA foci in cell nuclei with respect to AB34 control strain (Fig 2B and 2C). These results claim against the presence of massive DNA damage during the formation of the \textit{b-dependent} filament as the trigger that activates the Atr1-Chk1 pathway in \textit{U. maydis}.

**Characterization of elements upstream of Atr1-Chk1 axis in \textit{U. maydis}**

The absence of RPA foci does not necessarily preclude the presence of some specific damage in the DNA during the induction of the \textit{b-dependent} filament. It is plausible that in response to \textit{b}-induction, some limited DNA damage could be enough to induce the Atr1-Chk1 cascade, even when this is not detected by the presence of RPA foci. Therefore, we reasoned that the characterization of elements acting upstream of the Atr1-Chk1 cascade would help to define which kind of DNA damage signal, if any, could be involved in the \textit{b-dependent} activation of Chk1 in \textit{U. maydis}.

Elements upstream of the Atr1-Chk1 cascade have extensively been investigated in other organisms. While ATR recruitment to the damaged DNA seems to rely just on the presence of RPA-coated ssDNA, its activation needs action from other proteins. This is the case of the PCNA-like complex 9-1-1 (Rad9-Rad1-Hus1), which acts at two distinct levels depending on the species and the cell cycle phase [33]: directly activating ATR by DNA-bound 9-1-1; or indirectly, by recruiting the TopBP1/Dpb11 protein to the damaged sites, which in turn directly activates ATR [30]. Nevertheless not only 9-1-1 is able to assemble TopBP1 to these areas, and there are cases in which this function has also been found in the MRN complex (Mre11-Rad50-Nbs1) [34]. In addition to these complexes, adaptor proteins working as scaffolds are required for the appropriate transmission of the DNA damage signal. For instance, in budding yeast the adaptor proteins Rad9 and Mrc1 are required for checkpoint kinases activation in response to different kind of damages [35–37].

Components from 9-1-1 complex, like Rec1 (the Rad1 ortholog), and from MRN complex, like Mre11, have been previously described for \textit{U. maydis} [38, 39]. We queried the NCBI and Broad Institute databases for \textit{U. maydis} homologues of TopBP1/Dpb11, \textit{S. cerevisiae} Rad9 (\textit{S. pombe} Crb2) and Mrc1. With the exception of Rad9/Crb2, we were able to identify in the genome of \textit{U. maydis} the putative homologues of TopBP1/Dpb11 (um00290, renamed as Dpb11) and Mrc1 (um06299, renamed as Mrc1).

Neither \textit{rec1} nor \textit{mre11} are essential genes in \textit{U. maydis}, and therefore it was possible to construct the respective loss-of-function mutants (See below). In contrast, both Dpb11 and Mrc1 turned out to be essential proteins in \textit{U. maydis}, and we were unable to disrupt their respective genes in haploid cells. In other organisms, both Dpb11/TopBP1 and Mrc1/Claspin maintain additional functions during DNA replication [40, 41] that could explain their essential role. In \textit{S. cerevisiae}, Dpb11 is also essential but it is possible to separate the critical replication function from the checkpoint activation function by using specific mutants. One of these mutants is \textit{dpb1-11}, which carries a truncated C-terminus immediately after the fourth BRCT domain (W583STOP) [42]. We tried to recreate this mutant in haploid cells introducing a stop codon at the equivalent residue in \textit{U. maydis} \textit{dpb11} (Q846STOP, S1 Fig) with no success, suggesting that this kind of mutation does not recapitulate the same phenotype in \textit{U. maydis}. In the case of Mrc1, separation-of-function mutants have also been described for \textit{S. cerevisiae} \textit{MRC1},
Damage-Independent DDR Activation in Smut Fungus

(A) 

- DIC
- Control
- 5 mM HU
- 0.01% MMS

(B) 

- AB33 Rfa1-GFP
- 0 h
- 2 h
- 4 h

(C) 

Bar graph showing cells with RPA foci (%) over time of induction (hours) for AB33 and AB34.
consisting in site-specific mutations in all SQ/TQ residues [43]. *U. maydis* Mrcl carries 32 SQ/TQ residues scattered along the entire protein. However, we have found that, in *U. maydis*, the *mrc11-914* allele (S2 Fig), carrying a C-terminal end truncation of Mrcl resulted into viable cells that do not respond to checkpoint activation (see below).

To establish whether 9-1-1, MRN, and Mrcl are required in *U. maydis* for Chk1 activation, first we confronted the respective mutants to different genotoxic insults: UV irradiation, which induces pyrimidine dimers in DNA; HU, which inhibits ribonucleotide reductase and, therefore, affects replication by depletion of deoxynucleotide triphosphates, causing replication fork stalling and collapse; MMS, which induces DNA alkylation; phleomycin, a radiomimetic drug that causes DSB in DNA; and ionizing radiation (IR), which also generates DSB. We have found that all the mutants were sensitive to MMS, to phleomycin and to both UV and gamma irradiation, but only *rec1Δ* and *mrc11-914* mutants were sensitive to HU (Fig 3A). These results were consistent with different complexes devoted to signaling different forms of DNA damage —MRN, which senses DSBs, and 9-1-1, which senses ssDNA that is produced as a consequence of DNA replication stress or resection [44].

Activation of the DDR in *U. maydis* is marked by phosphorylation of Chk1 and by its relocation into the nucleus [14]. We examined the subcellular localization of GFP-tagged Chk1 in the presence of sub-lethal concentrations of either HU (producing mainly the presence of unreplicated forks) or phleomycin (inducing DSBs) (Fig 3B and 3C). While control cells showed a clear nuclear accumulation of the fluorescent signal in the presence of these DNA damaging agents, the different mutant strains failed to accumulate the fluorescent signal into the nucleus, with the exception of *mre11Δ* mutant in the presence of HU, coherently with the lack of sensitivity of this mutant to HU observed in the plate assay showed above.

These observations strongly suggested that 9-1-1 and MRN complexes, as well as the claspin-like Mrcl, are required for the activation of Chk1 under different types of genotoxic stress.

**Mrc1**, but neither 9-1-1 nor MRN, is required for the G2 arrest after the b-factor induction, as well as for full infection symptoms

Having demonstrated that *rec1*, *mre11* and *mrc1* genes were required for Chk1 activation upon induced genotoxic stress, we sought to address the question of whether they were also required for the cell cycle arrest that takes place upon b-induction. To test this possibility we introduced the *rec1Δ*, *mre11Δ* and *mrc11-914* alleles into the UMP112 strain, which is derived from AB33 and carries under the control of the dik6 promoter, a GFP fused to a nuclear localization signal. The expression of dik6 promoter is dependent on an active b heterodimer [25], and therefore it allowed us the use of the nuclear fluorescence as a marker of the release of cell cycle arrest (counting the nuclear content of the filaments) as well as a surrogate marker of the ability of the different mutants to respond to the b-program. We have found no difference in the proportion of cells responding to b-factor in the different mutant backgrounds. Even at short times upon induction of b expression (4 h), we have observed that almost the whole cell population shows nuclear fluorescence, indicating that there might be no interferences with the b-induced transcriptional program. Strikingly, we have found that both Rec1 and Mre11 were dispensable for b-dependent cell cycle arrest: No differences between control (UMP112) filaments and
filaments carrying the rec1Δ and mre11Δ alleles were found regarding nuclear content (Fig 4A and 4B). In contrast, in filaments carrying the mrc11-914 allele, it was possible to observe two and, less frequently, three nuclei, indicating that they are able to divide at least once, similarly as described for chk1 and atr1 mutants [15, 16].

Cells defective in Atr1 and Chk1 functions are affected in their pathogenic capability [15, 16]. Therefore we also investigated whether rec1Δ, mre11Δ and mrc11-914 mutants are able to infect maize. In U. maydis, virulence and sexual development are intricately interconnected. A prerequisite for generating the infectious stage is the mating of two compatible budding haploid cells to generate, after cell fusion, the infective dikaryotic filament. We constructed compatible haploid strains (i.e. a1b1 and a2b2 mating types) carrying the distinct mutant alleles. Mixtures of the respective mutant compatible strains, as well as wild-type controls, were used to infect seven-day-old maize seedlings by stem injection. The severity of disease symptoms
was then scored 14 days after infection [24] (Fig 4C). We have found that the infection with strains carrying either rec1Δ or mre11Δ alleles were indistinguishable from those with wild-type cells, while infection with strains carrying the mrc1<sup>1-914</sup> allele was less efficient and never produced large tumors, as previously described for chk1 and atr1 mutants [15, 16].

Altogether, these results indicated that while the scaffold Claspin-like Mrc1 protein may be required for b-induced cell cycle arrest and virulence of <i>U. maydis</i>, the DDR upstream regulators Rec1 (9-1-1 complex) and Mre11 (MRN complex) seem dispensable for these functions.

**Discussion**

During the infection of corn plants by <i>U. maydis</i>, the fungal cell cycle is arrested at G2 phase while it is growing on plant surface. This cell cycle arrest is required for the infective process, as mutants unable to stop the cell cycle progression are severely impaired in virulence [13, 15, 16]. Importantly, cell cycle arrest is dependent on the activation of the Atr1/Chk1 cascade, which would imply the presence or induction of some sort of DNA damage during this process. The aim of this work was to find any evidence of the DNA damage signal that would be feeding the ATR1/Chk1 cascade during the infective filament formation upon the b-factor expression in <i>U. maydis</i>. However, our results do not support the presence of the claimed DNA damage signal. We did not observe any signal of DNA damage in the form of RPA-GFP foci. Likewise, no obvious difficulties were found during S phase. Furthermore, the fact that neither 9-1-1 nor MRN, the two main sensors and transducers of DNA damage signals, were required for the b-induced cell cycle arrest as well as for virulence, also supports that no DNA damage signal would be feeding the Atr1/Chk1 cascade during the formation of the infective filament. It might also be plausible that induced and very localized DNA lesions could be recognized by alternative unknown sensors able to activate the checkpoint response. However, at this stage, we are much in favour of the hypothesis that Atr1/Chk1 activation during the development of the infective filament is independent of the presence of DNA damage as signaling cascade inducer.

Our analyses indicate that, although no upstream complexes such as MRN and 9-1-1 may be required, the adaptor protein Mrc1/Claspin seems necessary. Studies in yeast cells have shown that colocalization of Mec1 (Atr1) and Mrc1 is sufficient to induce the phosphorylation of the downstream kinase Rad53<sup>in vivo</sup> [45]. More importantly, in this report, authors showed that this phosphorylation is independent of upstream signaling components. It could be possible that in <i>U. maydis</i>, upon activation of the genetic program responsible for the infective filament formation, Atr1 would interact with Mrc1 in a DNA-damage independent manner and thereby the complex would activate Chk1.

If no DNA damage signal is associated with the b-induced activation of the Atr1/Chk1 cascade, how might this activation take place? We envisioned several possibilities based on recent reports that indicate the possibility of activation of this pathway in a DNA-damage independent manner in other systems.

The first of these possibilities could be related with some altered chromatin conformation associated with the transcriptional program induced by the b-factor. Recent work proved that
chromatin compaction was able to induce DDR in mammalian cells in the absence of any DNA damage [46]. So far, no studies about the chromatin state of *U. maydis* nuclei during the formation of the infective filament have been carried out, and therefore we cannot assure that such a chromatin compaction is taking place.

An alternative possibility is related to the stretch the nuclear envelope could be suffering during the formation of the infective filament. At this stage, the *U. maydis* cell experiments a strong induction of the polar growth, which is also dependent on the activation of the b-factor. During this elongation phase, the nuclei maintain a central position in the filament and for that it has to travel along the cytoplasm using microtubule-based motors, anchored to the nuclear pores [47]. Most likely, throughout this process the nuclear envelope is submitted to strong tensional forces and in fact nuclei inside the filament adopt a stretched appearance. In mammalian cells it has been reported that ATR activity at the nuclear envelope responds to mechanical stress. This ATR-mediated mechanical response occurs within the range of physiological forces, it is reversible, and it is independent of DNA damage signaling [48]. It could well be that in the case of *U. maydis*, the nuclear envelope deformation activated the Atr1/Chk1 cascade.

Although further work will be required to address these possibilities, our results reinforce the emerging idea that along evolution, the DDR cascade has been recruited to modulate developmental processes, most likely through its interaction with cell cycle machinery, even if no DNA damage signal exists.

**Supporting Information**

S1 Fig. (TIF)
S2 Fig. (TIF)

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**Author Contributions**

Conceived and designed the experiments: MTG CST JPM. Performed the experiments: MTG CST JPM. Analyzed the data: MTG CST JPM. Contributed reagents/materials/analysis tools: MTG CST JPM. Wrote the paper: MTG JPM.

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