CtIP -mediated alternative mRNA splicing finetunes the DNA damage response

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Abstract

In order to survive to the exposure of DNA damaging agents, cells activate a complex response that coordinates the cellular metabolism, cell cycle progression and DNA repair. Among many other events, recent evidence has described global changes in mRNA splicing in cells treated with genotoxic agents. Here, we explore further this DNA damage-dependent alternative splicing. Indeed, we show that both the splicing factor SF3B2 and the repair protein CtIP contribute to the global pattern of splicing both in cells treated or not to DNA damaging agents. Additionally, we focus on a specific DNA damage- and CtIP-dependent alternative splicing event of the helicase PIF1 and explore its relevance for the survival of cells upon exposure to ionizing radiation. Indeed, we described how the nuclear, active form of PIF1 is substituted by a splicing variant, named vPIF1, in a fashion that requires both the presence of DNA damage and CtIP. Interestingly, timely expression of vPIF1 is required for optimal survival to exposure to DNA damaging agents, but early expression of this isoform delays early events of the DNA damage response. On the contrary, expression of the full length PIF1 facilitates those early events, but increases the sensitivity to DNA damaging agents if the expression is maintained long-term.

INTRODUCTION

DNA is constantly threatened by endogenous and external sources that compromise its integrity. Thus, during evolution eukaryotes have developed a complex signaling network that finetunes the response to those threats. Generally referred as the DNA Damage Response (DDR), such network affects virtually every aspect of the cell metabolism (Ciccia and Elledge, 2010; Jackson and Bartek, 2009). In addition to those changes, the DDR activates the actual repair of damaged DNA. There are many different DNA lesions, thus several specific repair pathways coexist. DNA double strand breaks (DSBs) are the most cytotoxic form of DNA damage. Indeed, repair of DSBs can be achieved by different mechanisms, generally grouped in two categories, regarding the use or not of a template for repair. Whereas non-homologous end-joining (NHEJ) uses no homology to seal DSBs, homologous recombination will copy the information from a homologous sequence (Jasin and Rothstein, 2013; Lieber, 2010). The decision between those pathways is controlled by the DDR, and relies in the activation or not of the processing of the DNA ends, the so-called DNA end resection (Symington et al., 2014). This regulation is mostly achieved by controlling a single protein, CtIP, which integrates multiple signals in order to activate or not end processing (Makharashvili and Paull, 2015; Symington et al., 2014). CtIP works together with several other proteins that affect the processivity of the end resection, mainly the tumor suppressor gene BRCA1, in order to modulate resection and, as a consequence, homologous recombination (Cruz-García et al., 2014).

Recently, a crosstalk between the DDR and the RNA metabolism at different levels has been discovered. Indeed, the number of factors that participate in the DNA damage response and/or are regulated by it has expanded considerably in recent years to include many RNA-related proteins, notably splicing and alternative splicing factors (Jimeno et

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al., 2019). So, post-translational changes of splicing factors following DNA damage such as phosphorylation, ubiquitination, sumoylation, neddylation, PARylation, acetylation and methylation of splicing factors, have been documented. On the other hand, bona fide DDR factors also directly control splicing. For example, BRCA1 regulates alternative splicing in response to DSB formation through its DNA damagedependent interaction with several splicing factors such as SF3B1, one of the subunits of the Splicing Factor 3B (SF3B) complex (Savage et al., 2014). SF3B is a multiprotein complex essential for the accurate excision of introns from pre-messenger RNA (Golas et al., 2003). This complex consists of seven subunits: SF3B1 (also known as SF3b155), SFB2 (SF3b145), SF3B3 (SF3b130), SF3B4 (SF3b49), SF3B5 (SF3b10), SF3B6 (SF314a) and SF3B7 (PHF5a) (Spadaccini et al., 2006). SF3B plays an indispensable role during the assembly of the pre-spliceosome recognizing the intron's branch point (Teng et al., 2017). Interestingly, several subunits of this complex have been found in genome wide screens for factors involved in DNA repair, affecting homologous recombination (Adamson et al., 2012), controlling genome stability (Paulsen et al., 2009) or as substrates of the checkpoint kinases (Matsuoka et al., 2007). Moreover, we recently reported that the SF3B complex directly interacts with CtIP and regulates its activity in DNA end resection (Prados-Carvajal et al., 2018).

Interestingly, in addition to its well defined role in DSB repair by regulating DNA end resection, CtIP seems to perform many additional tasks in the cell, affecting DNA repair, cell cycle progression, checkpoint activation, replication and transcription (Duquette et al., 2012; Liu and Lee, 2006; Makharashvili and Paull, 2015; Moiola et al., 2012; Wu and Lee, 2006). CtIP promotes the expression of several genes, such as Cyclin D1, and also activates its own promoter (Liu and Lee, 2006). The role of CtIP in regulating gene expression is confirmed by its interaction with other transcriptional

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factors like IIKZF1, TRIB3 and LMO4 (Koipally and Georgopoulos, 2002; Sum et al., 2002; Xu et al., 2007). CtIP contributes to DNA damage-dependent cell cycle arrest in S and G2 phases promoting p21 transcription (Li et al., 1999; Liu et al., 2014) and upregulating the expression of GADD45A (Liao et al., 2010). Additionally, CtIP has been reported to regulate R-loop biology. CtIP deficiency has been shown to promote the accumulation of stalled RNA polymerase and DNA:RNA hybrids at sites of highly expressed genes (Makharashvili et al., 2018). On the contrary, CtIP depletion reduces DNA:RNA hybrid accumulation dependent on *de novo* transcription of dilncRNA (damage-induced long non-coding RNAs) starting at DSBs (D'Alessandro et al., 2018). Hence, CtIP depletion seems to increase R-loops that are produced as a consequence of previous transcription and appears to decrease *de novo* production of diRNAs (DSB-induced small RNA), thus reducing the DNA:RNA hybrids formed after DNA damage.

Hence, CtIP has a central role in the DDR and DNA repair, but plays additional roles in RNA biology. Indeed, it physically interacts with the SF3B splicing complex (Prados-Carvajal et al., 2018). Thus, we wondered whether CtIP-SF3B functional relationship might extend to controlling mRNA splicing and, more specifically, DNA damage-induced alternative splicing. Here we show that both SF3B and CtIP, albeit in a more modest manner, influence expression and splicing of hundreds of genes. This effect is visible in unchallenged cells, but more evident when cells have been exposed to a DNA damaging agent. Then, we analyzed in detail the effect of a DNA damage- and CtIP-dependent alternative splicing event of the helicase PIF1. Although PIF1 and CtIP also interact directly and are involved in DNA end resection, we observed that such event is not involved in DNA end processing. However, it affects the cell survival upon exposure to DNA damaging agents. Interestingly, this alternative PIF1 form, when expressed constitutively, hampers the recruitment of DSB repair proteins at early time

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points but makes cells hyper-resistant to treatments with camptothecin.

RESULTS

SF3B controls DNA damage-induced alternative splicing

As mentioned in the introduction, SF3B controls HR and DNA end resection (Prados-Carvajal et al., 2018). Whereas the resection phenotype was completely dependent on regulation of CtIP, our data suggested other, CtIP-independent, roles of SF3B in DNA repair (Prados-Carvajal et al., 2018). Due to the well stablished role of SF3B in splicing and, particularly its implication in DNA damage-dependent alternative splicing (Savage et al., 2014), we decided to analyze this role in more detail. Thus, we carried out a splicing microarray Transcriptome Arrays HTA & MTA using both damaged (6 hours after 10 Gy of irradiation) or untreated cells that were depleted or not for SF3B2 (Figure 1A; see Materials and Methods for details). As previously published, SF3B2 depletion affects CtIP protein levels slightly (Prados-Carvajal et al., 2018). Such array allows the genome-wide study of RNA expression and RNA splicing simultaneously. As SF3B2 controls the levels of CtIP and BRCA1 mRNA (Prados-Carvajal et al., 2018), we first focused on total RNA levels genome-wide. Changes were considered significant when the fold change (FC) was 2 or more and the p-value less than 0.05. Indeed, SF3B2 depletion leads to the specific upregulation of 52 genes solely in undamaged conditions when compared with control cells (Figure 1B). Moreover, 26 genes were exclusively overexpressed in SF3B2 downregulated cells upon exposure to DNA damage (Figure 1B). Additionally, mRNAs abundance from 27 genes was increased in cells with reduced SF3B2 levels in both damaged and undamaged cells (Figure 1B). A list of those genes could be found in Table 1. On the other hand, SF3B2 depletion also reduced expression of 97 genes (Figure 1C and Table 2). Only 10 of those genes were downregulated specifically in unperturbed conditions, 82 in cells that

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were exposed to ionizing radiation and 5 in both conditions, including SF3B2 itself as expected due to the shRNA-induced downregulation (Figure 1C, Table 2).

In terms of mRNA splicing, we confirmed that both depletion of SF3B2 and DNA damage induction affect such RNA processing globally. We first calculated the splicing index of exons on all four conditions and compared them *in silico* (FC>2, p-value<0.05; see Materials and Methods for additional details). The results are summarized in Figure 1C and a list of genes can be found in Supplementary Table S1. More than 4500 genes were differentially spliced when SF3B2 was absent, compared with a non-targeted shRNA (Figure 1D). Almost 25% did so regardless of the presence or absence of an exogenous source of DNA damage (yellow), but almost 45% showed splicing events that were both DNA damage- and SF3B-dependent (red) and only 30% of the genes were spliced by SF3B in undamaged conditions (green). Thus, most of the splicing events that require SF3B2 happens in damaged samples, indicating that this factor is especially relevant in stress conditions.

Indeed, a different analysis considering all genes that show an alternative splicing upon irradiation indicates that only 14% did so both in control and in SF3B2 depleted cells, whereas 46% of the genes suffer DNA damage-induced alternative splicing only when SF3B2 was present, suggesting they require this factor for such event. Strikingly, an additional 40% of the genes suffer damage induced alternative splicing specifically in SF3B2 depleted cells, indicating that when the SF3B complex was absent the splicing landscape is severely affected and new events appear.

Interestingly, the pattern of gain (+) or loss (-) of specific events of alternative splicing was similar in all situations (Table 3) with the exception of SF3B2-depleted cells upon irradiation, that was more pronounced in agreement with a strong role of the SF3B complex in DNA damage-induced alternative splicing (Savage et al., 2014).

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Despite the strong quantitative difference in splicing, qualitatively the types of events were similarly distributed in all cases (Table 3).

In order to validate the array, we studied the mRNA level of several splicing variants of genes that were identified as positives in the analysis. Due to our interest, we focused mainly on those that are related to DNA resection, recombination or the DNA damage response. To do so, we depleted SF3B2 using siRNA and induced or not DNA damage (10 Gy irradiation; Figure 2A). Cells where incubated during 6 hours to allow accumulation of DNA damage-dependent isoforms. We used qPCR and sets of isoformspecific primers (Table 4) to study the level of different variants. In all cases, we included an analysis of a "common isoform" that is present ubiquitously in all conditions. The ratio between the alternative variant and the "common isoform" was normalized to control cells. As shown in Figure 2B, the levels of a specific BRCA1 isoform increased upon DNA damage, but SF3B2 depletion blocks the accumulation of such BRCA1 mRNA specie even in untreated cells. Thus, we described a SF3B2- and DNA-damage induced splicing variant of this mRNA. Differently, we confirmed that RAD51 and EXO1 have a damaged-dependent isoform that is independent of SF3B2 (Figures 2C and 2D). Also, in agreement with the array data, the levels of a DNA2 mRNA variant increased specifically with DNA damage in the absence of SF3B2 (Figure 2E). PIF1 mRNA alternative isoform expression increased both upon irradiation and upon SF3B2 depletion in an epistatic manner (Figure 2F). Finally, we studied ATR, whose alternative splicing is dependent on SF3B2 regardless the presence or absence of DNA damage (Figure 2G). In summary, in agreement with the array, SF3B complex and/or DNA damage presence controls the alternative splicing of DNA repair factors.

CtIP controls mRNA expression and splicing of several genes

As mentioned previously, SF3B directly interacts and regulates the resection factor

CtIP. Interestingly, CtIP is a multifunctional protein that works in DNA repair, but also in other processes, including transcription (Wu and Lee, 2006). Moreover, other proteins related to DNA end resection, such as BRCA1, also have a role in RNA metabolism (Kleiman et al., 2005; Veras et al., 2009). Thus, we wondered whether CtIP could also play role in RNA splicing due to its connection with SF3B. To test this idea, and as for SF3B2, we used the splicing microarray to analyze RNA abundance and splicing genome-wide in cells depleted or not for CtIP, both in damaged and untreated conditions (Figure 1A for depletion of CtIP). When studying genome wide expression level of human genes, we observed that upon depletion of CtIP, and despite the assigned function in transcription, only 74 mRNAs showed altered abundance: 36 were upregulated and 38 downregulated (Figure 3A and B and Tables 5 and 6; Note that CtIP itself is among the downregulated ones, as expected due to the effect of the shRNA). Only 12 genes were exclusively upregulated in cells exposed to IR in cells depleted for CtIP compared with control cells (Figure 3A and Table 5). However, in undamaged conditions solely 22 genes were upregulated and the levels of only two genes increased in both conditions, with and without damage, in cells downregulated for CtIP (Figure 3A and Table 5). On the other hand, CtIP depletion reduced the expression of 16 genes exclusively in unperturbed cells whereas the expression of 18 genes were decreased in irradiated cells (Figure 3B and Table 6). The expression of only 4 genes was downregulated upon CtIP depletion in both damaged and undamaged cells (Figure 3B and Table 6).

Additionally, we studied mRNA splicing in the same conditions mentioned above and, interestingly, we realized that the depletion of CtIP showed a strong effect on RNA processing of hundreds of genes. As shown in Table 7, columns 4 and 5, CtIP downregulation on its own changed the pattern of mRNA splicing compared to control

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conditions, even though this phenotype was more pronounced in unperturbed cells.

As CtIP and SF3B2 physically interact, we wondered how many of the events that show a CtIP-dependent splicing did also rely on SF3B2 for that process, regardless of the exposure or not to DNA damage (Figure 3C). As expected, the number of splicing events that were dependent on SF3B2, a bona fide splicing factor, was higher than those that require CtIP. Indeed, only less than 30% of those SF3B2-dependent events were diminished upon CtIP depletion. But interestingly, around 70% of the CtIP-dependent splicing events were also affected by SF3B2. Thus, our results suggest that CtIP has a role in splicing, although less prominent than SF3B2. Moreover, most CtIP-dependent splicing events require also the SF3B complex, reinforcing the idea that they might act together. On the contrary, SF3B can readily act on the splicing of many genes independently of CtIP.

Considering the role of CtIP in the response to DNA damage, we decided to simultaneously analyze the effect of CtIP depletion and irradiation on genome-wide alternative splicing. Thus, we carried out another analysis in which we compared all conditions in pairs: control cells without DNA damage (siNT) or exposed to IR (siNT6h), or depleted for CtIP in unperturbed (siCtIP) or damaged cells (siCtIP6h) (Figure 3D). Only 107 genes were altered due to CtIP absence in irradiated cells, whereas 205 did so in undamaged cells (Figure 3D; in green and blue respectively). The splicing of 494 genes was changed specifically in response to DNA damage exclusively in CtIP depleted cells (Figure 3D, in red). In a similar number of genes (423 genes; Figure 3D, yellow), mRNA splicing was modified in response to DNA damage only in cells that retained CtIP. We were particularly interested in those latter mRNA, as they represented DNA damage-dependent mRNA variants that are CtIP dependent. Hence, we reasoned that CtIP might mediate, directly or indirectly, such DNA damage-

dependent alternative mRNA processing. Among them, we found that CtIP controls the DNA damage-dependent splicing of the helicase PIF1. Interestingly, SF3B2 also affects PIF1 splicing (Figure 2F). Strikingly, PIF1 and CtIP physically interact and together contribute to DNA end resection over specific DNA structures such as G-quadruplexes (Jimeno et al., 2018). Thus, in order to study deeply the crosstalk between CtIP, DNA end resection and RNA splicing, we decided to focus on the altered splicing of PIF1.

CtIP controls mRNA splicing of PIF1

PIF1 is a helicase with a 5'-3' polarity. In humans there are only one PIF1 gene, but it was known to produce two well studied different transcripts. A short transcript (2295nt) produces the longer protein isoform (707aa) called PIF1B, which is located in the mitochondria. On the other hand, the longer transcript (2688nt) is translated into a smaller protein variant named PIF1 α (641aa) that is localized in the nucleus (Futami et al., 2007). The difference between both proteins is the presence of a mitochondrial localization domain in PIF1B, which also lacks the signal to translocate into the nucleus. Our array data showed additional splicing changes on the PIF1 α backbone that were CtIP- and DNA damage-dependent (Figure 4A). We studied the inclusion of exon 3 (Figure 4A (I)), exon 4 (Figure 4A (II)), exon 9 (Figure 4A (III)) and exon 10 (Figure 4A (IV)). All these optional events are combinatorial and not mutually exclusive, so a mix of all the possible different species of mRNA coexist in all condition, regardless CtIP or DNA damage, but the array predicts changes on different conditions. We studied the abundance of the different alternative events by qPCR in cells depleted or not for CtIp with an siRNA and exposed or not to 10Gy of ionizing radiation (Figures 4B-G). The presence of an exon 2-3 junction was only slightly increased upon irradiation or CtIP depletion, but the changes were not significant (Figure 4C). A similar, but in this case significant, change was observed for exon 4 inclusion event

(Figure 4D). Exon 8-9 junction presence was strongly increased in response to irradiation in control cells, and both its inclusion and DNA damage-accrue was completely dependent on CtIP presence (Figure 4E). Exon 10 inclusion event in the mRNA analysis rendered no statistically significant changes in any conditions (Figure 4F). These results suggested that the clearest CtIP-dependent alternative events in response to exogenous damage occur in the exon 4 inclusion and, more specially, in exon 8-9 junction of *PIF1* gene.

In order to determine which activity of CtIP is involved in the splicing of *PIF1*, we carried out several qPCR to analyze the presence of the 8-9 junction on PIF1 mRNA, but in cells bearing different mutated versions of CtIP (Figure 4G). We used GFP-CtIP as control, the resection defective T847A CDK phosphorylation site mutant (Huertas and Jackson, 2009) and E894A sumoylation mutants (Soria-Bretones et al., 2017). Lastly, and considering that BRCA1 interacts with CtIP (Yu et al., 2006) and has been involved in damage-dependent alternative splicing (Savage et al., 2014), the S327A CDK phosphorylation mutant, that does not interact with BRCA1 and still resects albeit at a slower pace (Cruz-García et al., 2014). The data were normalized to the control GFP-CtIP, set as 1. Strikingly, both CDK defective phosphorylation mutants GFP-CtIP-T847A and GFP-CtIP-S327A caused a consistent increase in this splicing event, albeit only statistically significant in the T847A CtIP version (Figure 4F). This suggested CDK phosphorylation of CtIP inhibits the splicing of, at least, PIF1 exon 8-9 junction. This is likely resection independent, as the E894A mutant, a sumoylation defective CtIP that is equally impaired in resection as the T847A mutant, did not share such phenotype (Figure 4G). Thus, specific posttranscriptional modifications seem also required for CtIP role in splicing. Specifically, CDK phosphorylation of CtIP blocks such role, suggesting that the splicing activity of CtIP happens mainly in G1.

PIF1 splicing variants modulate DNA repair

Taking together these results, we decided to study the relevance of those CtIP- and DNA damage-dependent *PIF1* mRNA splicing events in DNA repair process in human cells. Hence, we created two different *PIF1* splicing variants cDNA constructs. First, a *PIF1* containing all exons ("total *PIF1*"; *tPIF1*), which correspond to the canonical PIF1 α . In contrast, we also created a *PIF1* variant artificial cDNA that lacks both exon 4 and exon 9, the two exons which inclusion was more dependent on CtIP and DNA damage (Figures 4C and 4D). We named it "variant *PIF1*" (*vPIF1*). Both genes were expressed from pCDNA. In order to detect the expression of either variant in cells, we tagged both isoforms with GFP. An empty pCDNA plasmid was also used as control in our experiments. We transfected each plasmid (pCDNA, GFP-vPIF1 or GFP-tPIF1) into U2OS in order to study the effect of either isoform in human cells in response to DNA damage. Expression of the proteins coded by those variants is shown in Figure 5A.

Considering that in all the conditions tested for the splicing analysis we could always observe a mixture of different splicing variant, including the canonical PIF1 α we decided to leave the expression of the endogenous *PIF1* gene unperturbed, and combine it with the expression of the different PIF1 variants. First, we analyzed the ability of cells to survive to the DNA damaging agent camptothecin (CPT) when expressing the already spliced vPIF1 and tPIF1 constitutively. Strikingly, constant expression of the tPIF1 rendered cells sensitive to DNA damage when compared with cells expressing the empty plasmid (Figure 5B). This was not observed when the CtIP- and DNA damage-dependent spliced form vPIF1 was constitutively expressed (Figure 5B), suggesting that continuous expression of tPIF1 hampers DNA repair and, therefore, agreeing with the idea that a switch from tPIF1 to vPIF1 by DNA damage- and CtIP-induced alternative splicing ensures an adequate response.

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Considering the fact that PIF1 and CtIP physically interact and cooperate in DNA end processing (Jimeno et al., 2018), we decided to study DNA end resection in response to irradiation (10 Gy) in cells expressing either PIF1 construct. However, the overexpression of either isoform caused no effect in DNA resection or the recruitment of the resection and recombination factor BRCA1 (Figures 5C and D). As a control, we also analyzed the cell cycle in those cells to test whether such overexpression caused any change in the progress of cell cycle (Figure 5E). The lack of effect on RPA foci formation, an early event, and the timing of the observed splicing changes (6 h after irradiation) suggested that the transition from tPIF1 to vPIF expression might be more relevant for cell survival at later events of the DNA damage response and, therefore, it is separated from the resection role of PIF1.

Early expression of vPIF1 delays DNA repair

In order to understand why changes in *PIF1* splicing to produce vPIF1 was only induced in response to DNA damage and that isoform was not constitutively expressed, we set to analyze the repair of DSBs at early time points in the presence of PIF1 isoforms. Interestingly, we observed that constitutive expression of vPIF1, albeit enhancing the long-term survival in response to DNA damage (Figure 5B) hampers or delays the recruitment of both NHEJ and HR proteins. Indeed, early after irradiation, the recruitment of the NHEJ factors 53BP1 and RIF1 was mildly impaired (Figure 6A and 6B). More strikingly, the recruitment of the essential HR factor RAD51 was severely impaired by constitutive expression of vPIF1 (Figure 6C). This was not observed when tPIF1 was overexpressed, confirming that this isoform does not block repair. Thus, our data suggest that a timely expression of different isoforms of PIF1 fine-tunes the response to DNA damage. Indeed, it seems that tPIF1 presence is permissive for early events, such the recruitment of 53BP1, RIF1 or RAD51, but in the

long term is deleterious for cell survival in response to DSBs. vPIF1, on the contrary, increases the resistance to DNA damaging agents, despite the fact that expressed too early slows down DSB repair.

Differential localization of PIF1 variants

As mentioned, there are two well-characterized PIF1 isoforms, the nuclear PIF1 α and the mitochondrial PIF1 β . Although both our constructs, tPIF1 and vPIF1, are based on the nuclear form, PIF1 α , we wondered whether vPIF1, which lacks part of the helicase domain, might show a different localization. To test it, we performed a cell fractionation assay depleting endogenous PIF1 for clarity. As shown Figure 7A, the protein produced by the tPIF1 construct was mostly chromatin associated as expected (green arrow), whereas the vPIF1, was mainly cytoplasmic (red arrow). Those localizations did not change if cells were exposed to exogenous DNA damage (Figure 7B).

DISCUSSION

In this work, we have discovered that not only the splicing complex SF3B but also the DNA repair factor CtIP controls the mRNA splicing of several genes, including proteins involved in DDR. Additionally, we have characterized the relevance of some splicing events on *PIF1* gene dependent on CtIP and DNA damage for the DDR and DNA repair.

Our data suggest that the SF3B complex controls the splicing and abundance of different mRNAs, both under unchallenged conditions and especially as a response to DNA damage. In fact, several members of this protein complex have been identified by different genome-wide analysis as targets of the DDR (Beli et al., 2012; Bennetzen et al., 2010; Elia et al., 2015; Matsuoka et al., 2007). This is especially relevant in

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response to DNA damage, as a large set of genes have a differential splicing in SF3B2 depleted cells compared with control cells specifically upon exposure to ionizing irradiation. Interestingly, many cellular pathways are represented in this set of genes, from stress signalling and apoptosis to organelle organization and vesicle transport. Especially striking is the large effect in the mitotic cell cycle, RNA metabolism and protein modifications, known targets of the DNA damage response (Supplementary Figure 2). These DNA damage-dependent changes in mRNA metabolism agree with previous studies that reported large changes in mRNA expression (Gasch et al., 2001; Rieger et al., 2004). In SF3B case, we hypothesize that both aspects, gene expression and splicing changes, are indeed related to the splicing activity of the complex. Although regulation of transcription has been primarily associated for such alterations in gene expression, it has been increasingly clear that post-transcriptional modifications could indeed affect mRNA levels in response to DNA damage (Boucas et al., 2012; Fan et al., 2002). Indeed, up to 50% of the changes on mRNA level in response to genotoxic agents could be attributed to mRNA turnover and not transcription (Boucas et al., 2012; Fan et al., 2002). Alternative splicing is known to affect mRNA stability in response to DNA damage by creating non-productive transcripts that are subjected to degradation by the non-sense mediated decay pathway (Barbier et al., 2007; Ip et al., 2011). In other cases, direct splicing-dependent gene expression repression or activation in response to DNA damage has been observed (Ip et al., 2011; Pleiss et al., 2007). Along those lines, we propose that alternative splicing controlled by the SF3B complex affects generally expression levels and the accumulation of alternative spliced mRNA in response to DNA damage. This global response will help the cells to fine-tune the response to broken DNA. Additionally, our splicing array data shows that SF3B2 affects also the splicing of key DDR factors, such as BRCA1, RAD51, RIF1, DNA2 and EXO1. This

might reflect a critical role of the SF3B complex in preparing the cells to the exposure to genotoxic agents.

Similarly, we show that CtIP affects the mRNA accumulation of hundreds of transcripts. This agrees, in principle, with the established role of CtIP in transcription (Koipally and Georgopoulos, 2002; Li et al., 1999; Liao et al., 2010; Liu and Lee, 2006; Liu et al., 2014; Sum et al., 2002; Xu et al., 2007). However, we also observe a prominent effect in splicing genome wide. Such role seems to require the functional interaction with the SF3B complex, as the majority of those events were altered also when SF3B2 was depleted. Splicing occurs cotranscriptionally and there is an intense crosstalk between transcription efficiency and splicing (Aslanzadeh et al., 2018; Braunschweig et al., 2014; Fong et al., 2014; Howe et al., 2003; Ip et al., 2011). Thus, transcription impairment could modify splicing efficiency and, on the contrary and as discussed for SF3B above, defective splicing might affect accumulation of mRNA that can be, erroneously, interpreted as transcriptional defects. Thus, in the case of CtIP is not so clear if those two roles, in transcription and splicing, are really two independent functions or simply both sides of the same coin. In any case, the regulation of the accumulation of different species of mRNA of many genes might explain why CtIP seems to play so many different roles in many processes. Indeed, there are evidences of other cases in which a RNA metabolism protein modulates its role in DDR through regulating the splicing of other factors involved in that process (Pederiva et al., 2016; Savage et al., 2014; Shkreta and Chabot, 2015). For example, PRMT5 regulates its effects on DNA repair by controlling the RNA splicing of several epigenetic regulators, especially the histone H4 acetyl-transferase TIP60 (KAT5) and the histone H4 methyltransferase SUV4-20H2 (KMT5C) (Hamard et al., 2018).

We propose that, for many phenotypes, and specially including DNA end resection,

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recombination and the DNA damage response, CtIP and SF3B probably participates at two different levels, directly, working as repair factors, but also indirectly as general transcription/splicing regulators. In that regard, they closely resemble its interactor BRCA1, which also seems to be involved at many different levels. Such double involvement of some RNA factors in DNA repair, which has simultaneous RNA-mediated and RNA-indirect roles in homologous recombination, was also observed by other authors (Anantha et al., 2013). This creates complex regulatory networks that are able to integrate multiple cellular signals and elicit sophisticated response that fine-tune the response.

Other proteins are likely involved in such complex networks, including the helicase PIF1. Interestingly, PIF1 and CtIP directly interacts and participates in resection of DSBs at atypical DNA structures such as G-quadruplexes (Jimeno et al., 2018). But additionally, CtIP controls a DNA damage alternative splicing of PIF1 that modulates the response to DNA damaging agents. A proficient DDR seems to require a timely switch between the two forms described here, the tPIF1 and vPIF1. Whereas vPIF1 slightly impairs early events in DNA repair, its presence ensures a better survival to DNA damage. On the contrary, the presence of tPIF1 does not affect those early events, but if maintained in time compromise viability of cells exposed to camptothecin. Interestingly, the main difference between both forms is the change in cellular localization and the presence of an active helicase domain. Whereas tPIF1 maintains such activity, vPIF1 lacks exon 9 and, therefore, misses part of the active site. Thus, it is likely that tPIF1 participates on DNA end resection and DNA repair early on, working on a DNA substrate as a helicase. However, our data suggest that cells prefer to reduce this active PIF1 pool to ensure survival. One possibility is that the switch between both isoforms reduces the active pool of the helicase, both by sending the protein to the

cytoplasm and by destroying the active site, to avoid interference of the helicase with very late steps of DNA repair or the DDR. Alternatively, it is formally possible that the vPIF1 plays some active role in the cytoplasm that facilitates survival. Considering that another isoform, PIF1ß is essential for mitochondrial metabolism, we could not exclude this hypothetical function, although it will imply a role that does not require a functional helicase domain. In any case, PIF1 alternative splicing illustrates how CtIP might have additional effects in the response to DNA damage that has been, so far, overlooked. Strikingly, our data suggest that the regulation of PIF1 is modulated by the threonine 847 of CtIP, although not because resection is impaired of this mutant. This residue is a well stablished CDK site (Huertas, 2010; Polato et al., 2014). Interestingly, and albeit less clearly, CDK phosphorylation of CtIP S327 seems also involved. Hence, it is possible that upon DNA damage but only in cells in the G1 phase of the cell cycle, CtIP activates the expression of the isoform called vPIF1. This will separate the different roles of CtIP in a cell cycle dependent manner, with the DNA damage function mainly on G1 and the DNA end resection and homologous recombination exclusively in S and G2.

In agreement with a tight relationship between BRCA1, CtIP and the SF3B complex, all three are intimately related to cancer appearance and, more specifically, with breast cancer incidence (Gokmen-Polar et al., 2019; Maguire et al., 2015; Paul and Paul, 2014; Soria-Bretones et al., 2013). In the case of CtIP and BRCA1, this connection with cancer has been mostly explained as a defective DNA repair. However, considering the involvement of CtIP, described here, and BRCA1 (Savage et al., 2014) in RNA splicing, maybe the cancer connection should be revisited on the light of those novel roles. Conversely, recent studies have detected recurrent mutations in components of the spliceosome in myelodysplastic syndromes (Pollyea et al., 2019; Shiozawa et al., 2018),

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renal cell carcinoma (Verma and Das, 2018; Yang et al., 2017), chronic lymphocytic leukemia (Agrawal et al., 2017; Maleki et al., 2019), lung adenocarcinoma (Kim et al., 2018; Mao et al., 2019), breast cancer (Gokmen-Polar et al., 2019; Zhao et al., 2019) or pancreatic cancer (Tian et al., 2015; Zhou et al., 2017). Moreover, alterations in expression of splicing factors, including SF3B, can derive in various types of cancers (Alsafadi et al., 2016; Goswami et al., 2014; Maguire et al., 2015; Zheng et al., 2018). This probably reflects the importance of mRNA splice variants of several important genes in apoptosis, metabolism, and angiogenesis (Grosso et al., 2008). However, another tantalizing possibility, not yet analysed in detail, is that some of those connections of SF3B with cancer might be a consequence of its more direct role in DNA repair. Furthermore, it might be possible to exploit the defective DNA repair and DDR in SF3B deficient cancer for therapeutic interventions. Thus, this crosstalk between DNA repair and the DDR and splicing might become in the future an important target for cancer treatment.

MATERIALS AND METHODS

Cell lines and growth conditions

U2OS human cell lines were grown in DMEM (Sigma-Aldrich) supplemented with 10% fetal bovine serum (Sigma-Aldrich), 2 mM L-glutamine (Sigma-Aldrich) and 100 units/mlpenicillin and 100 μ g/ml streptomycin (Sigma-Aldrich). For cells expressing GFP-tPIF1 and GFP-vPIF1, medium was supplemented with 0.5 mg/ml G418 (Sigma).

siRNAs, plasmids and transfections

siRNA duplexes were obtained from Sigma-Aldrich, Dharmacon or Qiagen (Table 8) and were transfected using RNAiMax Lipofectamine Reagent Mix (Life Technologies), according to the manufacturer's instructions. Plasmid transfection of U2OS cells with

PIF1 variants was carried out using FuGENE 6 Transfection Reagent (Promega) according to the manufacturer's protocol.

Cell fractionation

U2OS cells stably expressing the different PIF1 isoforms were subjected to nuclear/cytoplasm fractionation to analyze the distribution of both PIF1 variants in the different cellular compartments. Cells were transfected with siRNA against PIF1 (Table 8) to remove endogenous PIF1 and 48 hours later they were irradiated or not with 10 Gy ionizing radiation. One-hour post-irradiation, cells were harvested in Resuspension Buffer (10 mM HEPES pH 7.9, 10 mM KCl, 1.5 mM MgCl2, 0.34 M sucrose, 10% glycerol, 1 mM DTT and protease inhibitors). To disrupt cellular membranes, Triton X-100 was added to cell suspensions to a final concentration of 0.1%, and samples were incubated on ice for 8 minutes. Cells were then centrifuged at 4°C at 1300 g for 5 minutes to separate the cytoplasmic fraction from the nuclei pellet. Nuclei were then washed once with Resuspension Buffer and incubated in 100 μ l of this buffer containing 90 U/ml Benzonase (VWR) and 0.1 mg/ml BSA for 1 hour on ice. Cytoplasmic and nuclear fractions were analyzed by western blotting.

SDS-PAGE and Western blot analysis

Protein extracts were prepared in 2× Laemmli buffer (4% SDS, 20% glycerol, 125 mM Tris-HCl, pH 6.8) and passed 10 times through a 0.5 mm needle–mounted syringe to reduce viscosity. Proteins were resolved by SDS-PAGE and transferred to low fluorescence PVDF membranes (Immobilon-FL, Millipore). Membranes were blocked with Odyssey Blocking Buffer (LI-COR) and blotted with the appropriate primary antibody (Table 9) and infra-red dyed secondary antibodies (LI-COR) (Table 10). Antibodies were prepared in Blocking Buffer supplemented with 0.1% Tween-20.

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Membranes were air-dried in the dark and scanned in an Odyssey Infrared Imaging System (LI-COR), and images were analyzed with ImageStudio software (LI-COR). \$%

Immunofluorescence and microscopy

For RPA foci visualization, U2OS cells knocked-down for different proteins were seeded on coverslips. At 1 h after either irradiation (10 Gy), coverslips were washed once with PBS followed by treatment with Pre-extraction Buffer (25 mM Tris-HCl, pH 7.5, 50 mM NaCl, 1 mM EDTA, 3 mM MgCl2, 300 mM sucrose and 0.2% Triton X-100) for 5 min on ice. To visualize RAD51 foci, cells were cultured for 3 h after irradiation. For visualize BRCA1 foci, Pre-extraction Buffer 2 (10mM PIPES pH 6.8, 300 mM sucrose, 50mM Nacl, 3mM EDTA, 25X proteinase inhibitor and 0.5% Triton X-100) was used. Cells were fixed with 4% paraformaldehyde (w/v) in PBS for 15 min. Following two washes with PBS, cells were blocked for 1 h with 5% FBS in PBS, costained with the appropriate primary antibodies (Table 9) in blocking solution overnight at 4°C or for 2 h at room temperature, washed again with PBS and then coimmunostained with the appropriate secondary antibodies for 1 h (Table 10) in Blocking Buffer. After washing with PBS and drying with ethanol 70% and 100% washes, coverslips were mounted into glass slides using Vectashield mounting medium with DAPI (Vector Laboratories). RPA foci immunofluorescences were analyzed using a Leica Fluorescence microscope.

For 53BP1 visualization, U2OS cells were seeded and transfected as previous described. Once collected, cells were fixed with methanol (VWR) for 10 min on ice, followed by treatment with acetone (Sigma) for 30 sec on ice. For RIF1 foci visualization, cells were fixed with 4% PFA for 15 min, washed twice with 1× PBS and then permeabilized for 15 min with 0.25% Triton diluted in 1× PBS. Samples were immunostained as described above with the appropriate primary and secondary antibodies (Tables 9 and

10). Images obtained with a Leica Fluorescence microscope were then analyzed using Metamorph to count the number, intensity and size of the foci.

Cell cycle analysis

Cells were fixed with cold 70% ethanol overnight, incubated with 250 μ g/ml RNase A (Sigma) and 10 μ g/ml propidium iodide (Fluka) at 37°C for 30 min. For each replicate, 10.000 cells were analyzed with a FACSCalibur (BD). Cell cycle distribution data were further analyzed using ModFit LT 3.0 software (Verity Software House Inc).

RNA extraction, reverse transcription and quantitative PCR

RNA extracts were obtained from cells using NZY Total RNA Isolation kit (Nzytech) according to manufacturer's instructions. To obtain complementary DNA (cDNA), 1 μ g RNA was subjected to RQ1 DNase treatment (Promega) prior to reverse transcription reaction using Maxima H Minus First Strand cDNA Synthesis kit (Thermo Scientific) according to manufacturer's instructions. Quantitative PCR from cDNA was performed to check siRNA-mediated knock-down of several proteins. For this, iTaq Universal SYBR Green Supermix (Bio-Rad) was used following manufacturer's instructions. DNA primers used for qPCR are listed in Table 4. Q-PCR was performed in an Applied Biosystem 7500 FAST Real-Time PCR system. The comparative threshold cycle (Ct) method was used to determine relative transcripts levels (Bulletin 5279, Real-Time PCR Applications Guide, Bio-Rad), using β -actin expression as internal control. Expression levels relative to β -actin were determined with the formula $2^{\Delta \Delta_{\alpha}}$ (Livak and Schmittgen, 2001).

Microarray

To analyse RNA splicing genome wide from cells depleted or not for SF3B2 or CtIP in damaged (10 Gy) and untreated conditions the splicing microarray Transcriptome

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Arrays HTA & MTA (Affymetrix) was used as previously described (Jimeno-González et al., 2015). RNA was obtained using an RNA isolation kit, as explaineed in section above, in triplicates. The purity and quality of isolated RNA were assessed by RNA 6000 Nano assay on a 2100 Bioanalyzer (Agilent Technologies). Total RNA from each sample (250 ng) was used to generate amplified and biotinylated sense-strand cDNA from the entire expressed genome according to the GeneChip WT PLUS Reagent Kit User Manual (P/N 703174 Rev 1; Affymetrix Inc.). GeneChip HTA Arrays were hybridized for 16 h in a 45 °C oven, rotated at 60 rpm. According to the GeneChip Expression Wash, Stain and Scan Manual (PN 702731 Rev 3; Affymetrix Inc.), the arrays were washed and stained using the Fluidics Station 450 and finally were scanned using the GeneChip Scanner 3000 7G. Raw data were extracted from the scanned images and analyzed with the Affymetrix Command Console 2.0 Software. The raw array data were preprocessed and normalized using the Robust Multichip Average method. Data were processed further using Transcriptome Analysis Console (TAC) Software from Affymetrix, which performs a gene-level analysis or an alternative splicing analysis. For gene-level analysis, gene expression changes with P < 0.05(ANOVA) and a llinear fold changel ≥ 2 were considered significant. For alternative splicing analysis, splicing (SI) calculated the index was as SI=Log2[(Probe-set1Doxintensity/Gene1Doxintensity)/(Probe-set1ControlIntensity/G ene1ControlIntensity)]. Splicing changes were considered significant when the lsplicing index | was ≥ 2 and p < 0.05 (ANOVA).

Array data also were analyzed using AltAnalyze software (Emig et al., 2010) version 2.0 with core probe set filtering using DABG (detected above background; P value cutoff of 0.05) and microarray analysis of differential splicing (MiDAS exon analysis parameters; P value cutoff of 0.05). The signs of the splicing index values obtained

from AltAnalyze were changed to use the same splicing index definition throughout the manuscript. AltAnalyze incorporates a library of splicing annotations from UCSC KnownAlt database. GeneChip HTA 2.0 arrays include probes to detect 245.349 different transcript variants, supported by a variable number of ESTs in the databases. Intronic regions often are represented in the ESTs databases because of the retrotranscription of unspliced pre-mRNAs, aberrant splicing forms, and spurious firing of cryptic promoters inside introns, with functional or nonfunctional implications. For that reason, GeneChip HTA 2.0 arrays allow the identification of an intron-retention event

(media.affymetrix.com/support/technical/datasheets/hta_array_2_0_datasheet.pdf).

Statistical analysis

Statistical significance was determined with a Student's *t*-test or ANOVA as indicated using PRISM software (Graphpad Software Inc.). Statistically significant differences were labelled with one, two or three asterisks for P < 0.05, P < 0.01 or P < 0.001, respectively.

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TABLES:

Table 1: Genes upregulated upon SF3B2 depletion

Genes upregulated in	Genes upregulated	Genes upregulated in
undamaged cells	in damaged cells	undamaged and damaged cells
ABCA5	ANKRD18DP	BET1

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AC087073.1	ANKRD45	C12orf39
ANTXR2	CCDC30	CBWD1
ATG12	СНКА	CBWD2
C14orf37	CTD-2651B20.3	CBWD3
Clorf168	EIF5	CBWD6
C1orf63	GABRR3	CBWD7
C21orf91	HAVCR2	CDO1
C9orf156	JAKMIP1	CGRRF1
CCDC66	KLHL41	CTD-2023N9.3
CD274	NME8	FAM133B
CDKN1B	PDE4D	FLVCR1
CREBRE	RP11-386I8 6	GMFB
DNAIC27	RP11-473L15 3	KDM3A
DTD2	RP11-545I10 2	MITD1
ENOX1	RP4-553F4 6	MMP13
FBXW7	RRH	MTHFD2L
GBP1	SI FN12	NPM3
GDAP1	SPATA9	NOO1
GTF2E2	SYNE1	RP11-111F5 3
HFBP1	THEMIS	RSAD2
HSD17B13	TRAM1	SPH AR
IK7F3	TSPV5P	SRD543-4S1
IL 18P1	WDR87	STARD4
II 5	7NF442	
ITPR2	ZNF571	TXNDC2
KANSI 2	2111371	7NF844
KIA A 1467		
MAGOH		
METTI 13		
NCAPG2		
PARP16		
PCNXI 2		
PDIK11		
PPARGCIA		
RP11-159G9 5		
RP11-213G2 2		
RP11-261C10 3		
RP11-46C20 1		
RP13-996F3 4		
S100A8		
SCO1		
SEST SESN1		
SI C25A27		
SRPRB		
SUN3		
THOC7		
TP53INP1		
TRPM7		
IISP54		
ZMAT3		
ZNF669		

Table 2: Genes downregulated upon SF3B2 depletion

Genes downregulated in undamaged cells	Genes downregulated in damaged cells	Genes downregulated in undamaged and damaged cells
AHCYL1	GRAP	CAV1
AL603965.1	KRT27	EXOSC3
ALDH1L1	LINC00482	KDELC2
ALPL	OR6T1	NID1

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ANO1	PRSS22	ST3GAL5
ΔΝΧΔ8	RP1_300G12 2	
	RI 1-500012.2	
ANXA8L1	RP11-429E11.2	
ARHGAP31	SNCAIP	
ATG5	SPINK 14	
BCL2A1	VPS37C	
BECN1		
C12 or f50		
C120I130		
C14orf182		
CCL2		
CCL/		
CCNDBP1		
CDC45		
CHORDON		
CHORDCI		
CLMP		
CSF2		
CTC 42(D10.2		
CIC-436P18.3		
CYB5R2		
F2F1		
FAM49A		
FANCD2		
GALNT1		
GALINII		
GINS1		
HSPB11		
IADS		
IAKS		
ICAM1		
INPP5B		
TIGB5		
KCNH7		
KCNMA1		
KIAA1279		
KLHDC3		
KRT20		
KR16A		
KRT6C		
KRT75		
LGI2		
LRFN5		
LPPC15		
MBNL2		
MMP9		
MOCOS		
MOCOS		
MYH16		
NCEH1		
NDUES3		
ND0133		
NEK/		
NMD3		
DAL M3		
PAPOLA		
PDE9A		
PEPD		
PIEZO2		
PIR		
$DI \Delta 2G7$		
PLOD2		
PPP4R4		
QRFPR		
RCAN2		
DD11 1150115 1		
кг11-11 <i>32</i> П13.1		

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RP11-18H21.1			
RP11-283G6.5			
RP11-295K2.3			
RP11-47I22.3			
RP3-522D1.1			
RP5-968D22.1			
SF3B2			
SGK196			
SLC22A4			
SLCO2A1			
SSTR2			
ST6GALNAC3			
THRS1			
TNEPSEQ			
TOMM34			
TDDD2			
WAC AS1			
AUCVI 1			
AICILI AI 602065 1			
AL003903.1			
ALFL ANO1			
ANVAS			
AINAAO ANVAQL1			
ANAAOLI ADUCAD21			
AKHUAPSI			
BCL2AI DECN1			
G12arf50			
C120f150			
C140f1182			
CCL2			
CCNDDD1			
CCNDBP1			
CUODDC1			
CHORDET			
CLMP CCE2			
CSF2			
CVD5D2			
CIDJK2 E2E1			
FANCD2			
GALNT1			
GINS1			
HSPR11			
IARS			
ICAM1			
INPP5B			
ITGB3			
KCNH7			
KCNMA1			
$KI\Delta \Delta 1279$			
KI HDC3			
KRT20			
KRT64			
KRTGC			
KRT75			
I REN5			

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LRRC15	
MBNL2	
MMP9	
MOCOS	
MYH16	
NCEH1	
NDUFS3	
NEK7	
NMD3	
PALM3	
PAPOLA	
PDE9A	
PEPD	
PIEZO2	
PIR	
PLA2G7	
PLOD2	
PPP4R4	
PRLR	
QRFPR	
RCAN2	
RP11-1152H15.1	
RP11-18H21.1	
RP11-283G6.5	
RP11-295K2.3	
RP11-47I22.3	
RP3-522D1.1	
RP5-968D22.1	
SF3B2	
SGK196	
SLC22A4	
SLCO2A1	
SSTR2	
ST6GALNAC3	
THBS1	
TNFRSF9	
TOMM34	
TPK1	
TPPP3	
WAC-AS1	

Table 3: Specific splicing changes in response to IR and SF3B2 depletion. (+) represents gain and

(-) loss of each specific event

Compa	rison	Irradiated versus non irradiated samples in control cells	Irradiated versus non irradiated samples in SF3B2 depleted cells	SF3B2 depleted versus control cells in untreated conditions	SF3B2 depleted versus control cells in irradiated conditions
(+) terminı	alt-C- 15	274	220	201	375
(+) terminu	alt-N- 15	265	215	185	335
(+) alt-o	oding	39	44	35	53
(+)	nonsense	72	53	59	117

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mediated decay				
(+) retained intron	75	67	62	122
(+) truncated	43	52	54	115
(-) alt-C-terminus	228	234	231	503
(-) alt-N-terminus	245	249	222	479
(-) alt-coding	42	53	93	164
(-) nonsense mediated decay	44	44	77	152
(-) retained intron	61	68	76	175
(-) truncated	63	56	28	66
intron (-) truncated	61 63	68 56	76 28	175 66

Table 4. DNA primers.

ACTB qPCR Fwd	ACGAGGCCCAGAGCAAGA	RT-qPCR of Actin
ACTB qPCR Rvs	GACGATGCCGTGCTCGAT	RT-qPCR of Actin
BRCA1 common isoform Fwd	CCCTTTCACCCATACAC	To validate the microarray of SF3B2 splicing isoforms
BRCA1 common isoform Rvs	AAGTGTTGGAAGCAGG	To validate the microarray of SF3B2 splicing isoforms
RAD51 isoform Fwd	TCCAGAACAGCACCAAAG	To validate the microarray of SF3B2 splicing isoforms
RAD51 isoform Rvs	GTGGTGACTGTTGGAAG	To validate the microarray of SF3B2 splicing isoforms
RAD51 common isoform Fwd	CATVTGGAGGTAGCAGAAG	To validate the microarray of SF3B2 splicing isoforms
RAD51 common isoform Rvs	CTCGTGCTAATCTGGAC	To validate the microarray of SF3B2 splicing isoforms
EXO1 isoform Fwd	CCTCGGAGTGAGAGAAA	To validate the microarray of SF3B2 splicing isoforms
EXO1 isoform Rvs	TGTAGCAATCCCTGTATCCC	To validate the microarray of SF3B2 splicing isoforms
EXO1 common isoform Fwd	CTGAAGTGTTTGTGCCTGAC	To validate the microarray of SF3B2 splicing isoforms
EXO1 common isoform Rvs	CCACAACTGCACCAC	To validate the microarray of SF3B2 splicing
DNA2 isoform Fwd	CAGAGGCAAGCGATGA	To validate the microarray of

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		SF3B2 splicing isoforms
DNA2 isoform Rvs	AACCACAGGCGGTAGAGA	To validate the microarray of SF3B2 splicing isoforms To validate the
DNA2 common isoform Fwd	GGAGAAGAGTGGCAGTT	microarray of SF3B2 splicing isoforms
DNA2 common isoform Rvs	TCTGTCACCTGCCATTAG	To validate the microarray of SF3B2 splicing isoforms
ATR isoform Fwd	GTCAGGAAGGTCTATGTG	To validate the microarray of SF3B2 splicing isoforms
ATR isoform Rvs	GTCCTTGAAAGTACGG	To validate the microarray of SF3B2 splicing isoforms
ATR common isoform Fwd	CACCACAGGCACAATCA	To validate the microarray of SF3B2 splicing isoforms
ATR common isoform Rvs	TCCACTAACACAACTAGCCC	To validate the microarray of SF3B2 splicing isoforms
2 PIF1 Fwd	TGGTGAAGCGGCCTGTGGA	To validate the microarray of CtIP splicing isoforms
3 PIF1 Rvs	GTGAAGAAGATGCTCTGG	microarray of CtIP splicing isoforms
4 PIF1 Rvs	AGTGAGCCCAGGATTCGCTT	microarray of CtIP splicing isoforms To validate the
8 PIF1 Fwd	GTGTTCAGATGAGGTGAC	microarray of CtIP splicing isoforms
9 PIF1 Rvs	TAGTTGAAGGAGCTGG	To validate the microarray of CtIP splicing isoforms
10 PIF1 Rvs	CTGCCTCGAACCCAAC	To validate the microarray of CtIP splicing isoforms

Table 5: Genes upregulated upon CtIP depletion

Genes upregulated in undamaged cells	Genes upregulated in damaged cells	Genes upregulated in undamaged and damaged cells
AC010967.2	AC068535.3	MMP13
AC016582.2	ADAMTS9	TULP3
AC064852.4	CD200R1L	
AC073043.2	DMXL2	
AF064858.7	IL10	
ARL14	IL24	
C9orf153	KLHL6	
C9orf156	MMP3	
CBLN2	RP11-445K13.2	
COL10A1	SPATA9	
EGLN1	SRD5A3-AS1	

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IKZF3	WDR87	
KRT13		
NCEH1		
PARP16		
PPP2R1B		
PRSS44		
RP11-285B24.1		
RP11-290K4.1		
RP11-300J18.2		
RP11-9G1.3		
RP4-553F4.6		
RP4-650F12.2		
SGIP1		
U51244.2		

Table 6: Genes downregulated upon CtIP depletion

Genes downregulated in	Genes downregulated	Genes downregulated in
undamaged cells	in damaged cells	undamaged and damaged cells
DACT3	C8orf4	CtIP
DRAP1	CCL7	GALNT7
ELOVL7	CD69	OLR1
FGF9	CTGF	SMYD2
LINC00482	CXCL10	
MYOCD	GOLT1A	
NDP	INPP4B	
OR6T1	KAT2B	
RBMS3-AS1	MKX	
RCN2	NINJ1	
RP11-1069G10.1	NTRK3	
SEMA3D	PDCD4	
SEZ6	RP5-1172A22.1	
SMIM12-AS1	RP5-968D22.1	
SNORA38B	S1PR3	
VAV2	SLC22A16	
ZNF571	SYVN1	
	TMEM116	
	TMPRSS15	
	TPPP3	

Table 7: Specific splicing changes in response to IR and CtIP depletion. (+) represents gain and (-

) loss of each specific event

	Irradiated versus		Irradiated versus		CtIP depleted		CtIP	depleted
Comparison	non	irradiated	non	irradiated	versus	control	versus	control
	sample	es in	sampl	es in CtIP	cells in	untreated	cells	in

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	control cells	depleted	cells conditions	irradiated		
				conditions		
(+) alt-C-	274	202	87	44		
terminus	274	505	87	44		
(+) alt-N-	265	327	77	46		
terminus						
(+) alt-coding	39	43	28	7		
(+) nonsense	70	(2)	20	12		
mediated decay	12	63	20	12		
(+) retained						
intron	75	96	25	11		
(+) truncated	43	46	26	14		
(-) alt-C-terminus	228	282	142	80		
(-) alt-N-terminus	245	298	129	74		
(-) alt-coding	42	44	29	10		
(-) nonsense	44	57	36	19		
(-) retained	61	83	36	18		
(-) truncated	63	73	15	6		
Table 8: siRNAs used.						
Non Target D	Non Target D-001810-10-20 Dharmacon UGGUUUACAUGUUGUGUGA, UGGUUUACAUGUUUUCUGA and UGGUUUACAUGUUUUCUGA (mix)					

CtIP	Huertas and Jackson 2009	Sigma	GCUAAAACAGGAACGAAUC
SF3B2	HSC.RNAI.N00 6843.12.6	IDT	GGACUUGUCAUUUCAUGUUCUUATT
PIF1	N/A	IDT	GGAUGUUCUCAGGUUGUAUUUAUTT

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Target protein	Source	Supplier/Reference	Application	Dilution
GFP	Rabbit	Santa Cruz (sc-8334)	WB	1:1000
α-tubulin	Rabbit	Sigma (T9026)	WB	1:50000
RPA32	Mouse	Abcam (ab2175)	IF	1:500
γH2AX	Rabbit	Cell Signaling (2577L)	IF	1:500
RAD51	Mouse	Abcam; ab213	IF	1:1000
BRCA1	Mouse	Santa Cruz (sc-6954)	WB, IF	1:1000, 1:200
53BP1	Rabbit	NB100-304, Novus	WB, IF	1:1000,1:500
RIF1	Mouse	Bethyl Laboratories (A300-568A)	WB, IF	1:500, 1:200
PIF1	Mouse	Santa cruz (sc-48377)	WB	1:500

Fable 9): Primary	antibodies used	in this study	y.WB	, western blotting	. IF,	, immunofluorescence.
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Table 10. Secondary antibodies used in this study. WB, western blotting. IF, immunofluorescence

Antibody	Supplier/Reference	Application	Dilution
IRDye 680RD goat anti-mouse IgG (H+L)	LI-COR (926-68070)	WB	1:5000
IRDye 800CW goat anti-rabbit IgG (H+L)	LI-COR (926-32211)	WB	1:5000
Alexa Fluor 594 goat anti-mouse	Invitrogen (A11032)	IF	1:1000
Alexa Fluor 488 goat anti-rabbit	Invitrogen (A11034)	IF	1:1000

FIGURE LEGENDS:

Figure 1: SF3B2 depletion affects gene expression and splicing of many genes.

A, Representative western blot showing the expression levels of SF3B2 and CtIP upon depletion with the indicated shRNAs in cells exposed or not to 10 Gy of ionizing

radiation. α -tubulin blot was used as loading control. **B**, Distribution of the genes upregulated upon SF3B2 depletion regarding the exposure or not to DNA damage. Fold change (FC)>2, p-value<0.05. Gene expression was measured using the GeneChip HTA Array as described in the methods section. The number of upregulated genes in undamaged cells (green), 6 h after exposure to irradiation (10 Gy; pink) or both (yellow) is shown in a Venn diagram. **C**, Same as A but for downregulated genes. **D**, the GeneChip HTA Array was used to look for genes that changed their splicing upon SF3B2 depletion, as mentioned in the methods section. Other details as in A.

Figure 2: Splicing changes in DDR factors in cells depleted of SF3B2.

A, Representative western blot showing the expression levels of SF3B2 upon depletion with a siRNA against SF3B2 or a control sequence (siNT) in cells exposed or not to 10 Gy of ionizing radiation. α -tubulin blot was used as loading control. **B-G**, Specific RNA isoforms levels of the indicated genes were calculated as the ratio between the abundance of the specific splicing form normalized with the total amount of each gene RNA by quantitative RT-PCR using specific primers in cells transfected with the indicated siRNAs and 6 hours after irradiation or mock treatment. See Materials and Methods for details. A schematic representation of the splicing events measured is shown in each case on the top. The common splicing event analyzed is shown in green, oligos are represented as arrows. The specific splicing that changes upon SF3B2 depletion is shown in orange. The graphs represent the average and standard deviation of three independent experiments. Statistical significance was calculated using an ANOVA test. * p<0.05,** p<0.01 and *** p<0.005.

Figure 3: CtIP depletion affects gene expression and splicing of many genes.

A, Distribution of the genes upregulated upon CtIP depletion regarding the exposure

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or not to DNA damage. Fold change (FC)>2, p-value<0.05. Gene expression was measured using the GeneChip HTA Array as described in the Materials and Methods section. The number of upregulated genes in undamaged cells (green), 6 h after exposure to irradiation (10 Gy; pink) or both (yellow) is shown in a Venn diagram. **B**, Same as A but for downregulated genes. **C**, the GeneChip HTA Array was used to look for genes that changed their splicing upon SF3B2 and/or CtIP depletion, as mentioned in the Materials and Methods section. The Venn digram represents the one that change when CtIP (pink), SF3B2 (green) or both are downregulated. Other details as in A. **D**, Differential splicing events between different conditions: undamaged cells depleted for CtIP (siCtIP) or transfected with a non-target siRNA (siNT); or RNA collected 6h after irradiation in cells depleted for CtIP (siCtIP_6h) or control cells (siNT_6h).

Figure 4: PIF1 splicing changes upon CtIP depletion.

A, Schematic representation of PIF1 with the exons (boxes) and splicing events analyzed (roman numbers). **B**, Representative western blots showing the expression levels of CtIP upon depletion with a siRNA against CtIP or a control sequence (siNT) in cells exposed or not to 10 Gy of ionizing radiation. α -tubulin blot was used as loading control. **C**, Analysis of the exon 2 and exon 3 junction using quantitative PCR using primers located in exon 2 and 3 (see panel A) in cells depleted (siCtIP) or not (siNT) for CtIP, upon exposure to IR (black bars) or in unchallenged conditions (white bars). The presence of a constitutive exon was used as a control. The abundance of such event was normalized to the control cells in undamaged conditions. The average and standard deviation of three independent experiments is shown. Statistical significance was calculated using an ANOVA test. * p<0.05,** p<0.01 and *** p<0.005. **D**, same as B but for the inclusion of Exon 9. Primers in Exon 8 and

9 were used (see panel A). Other details as panel B. F, same as B but for the inclusion of Exon 10. Primers in Exon 8 and 10 were used (see panel A). Other details as panel B. G, Study of the exon 8-9 junction in cells depleted for endogenous CtIP and expressing the indicated mutants of CtIP. Other details as panel B.

Figure 5: Expression of different PIF1 splicing variants.

A, Western blot showing the abundance of different PIF1 isoforms in cells transfected with the empty pCDNA plasmid or pCDNA bearing the vPIF1 or tPIF1 splicing variants. **B**, Survival to different doses of camptothecin (CPT) in cells overexpressing the tPIF1 or vPIF1 isoforms as indicated. Cells transfected with the empty pCDNA vector were used as a control. The average and standard deviation of three independent experiments is shown. **C**, Percentage of cells positive for RPA foci upon exposure to 10 Gy of ionizing radiation. Cells expressed the indicated PIF1 variants. An empty pCDNA vector was used as a control. The average and standard deviation of three independent experiments is shown. No statistically significant differences were found using an ANOVA test. **D**, Same as A, but for BRCA1 foci. **E**, cell cycle analysis of cells transfected with the indicated plasmids. The percentage of cells in each cell cycle phase was analyzed as described in the Materials and Methods section. The average and standard deviation of three independent experiments is shown.

Figure 6: PIF1 splicing variants affect the recruitment of DDR factors.

A, Average number of 53BP1 foci per cell upon exposure to 10Gy of IR in cells transfected with the indicated vectors. The average and standard deviation of three independent experiments is shown. Statistical significance was calculated using an ANOVA test. * p<0.05. **B**, same as A but for RIF1 foci. **C**, same as A but for RAD51 foci.

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Figure 7: Cellular localization of PIF1 splicing variants.

A, Protein samples from undamaged cells expressing the indicated PIF1 isoforms were fractionated as described in the Materials and Methods section. Cytoplasmic, nuclear and chromatin fractions were resolved in SDS-PAGE and blotted with the indicated antibodies. vPIF1 is marked with a red arrow. tPIF1 is located with a green arrow. A representative experiment is shown. **B**, same as A, but in cells exposed to 10 Gy of ionizing radiation.





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С

A

Genes which splicing depends on SF3B2 CtIP















cytoplasm chromatin cytoplasm nucleus nucleus chromatin GFP-tPIF1 GFP-vPIF1 GFP GFP-tPIF1 GFP-vPIF1 GFP + . + + + + + + + . + -+ anti-GFP anti-GFP anti- α -tubulin anti- α -tubulin