Regulated Degradation of the Inner Nuclear Membrane Protein SUN2 Maintains Nuclear Envelope Architecture and Function

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Abstract

Nuclear architecture and functions depend on dynamic interactions between nuclear components (such as chromatin) and inner nuclear membrane (INM) proteins. Mutations in INM proteins interfering with these interactions result in disease. However, mechanisms controlling the levels and turnover of INM proteins remain unknown. Here, we describe a mechanism of regulated degradation of the INM SUN domain-containing protein 2 (SUN2). We show that Casein Kinase II and the C-terminal domain Nuclear Envelope Phosphatase 1 (CTDNEP1) have opposing effects on SUN2 levels by regulating SUN2 binding to the ubiquitin ligase Skp/Cullin1/F-BoxβTrCP (SCFβTrCP). Upon binding to phosphorylated SUN2, SCFβTrCP promotes its ubiquitination. Ubiquitinated SUN2 is membrane extracted by the AAA ATPase p97 and delivered to the proteasome for degradation. Importantly, accumulation of non-degradable SUN2 results in aberrant nuclear architecture, vulnerability to DNA damage and increased lagging chromosomes in mitosis. These findings uncover a central role of proteolysis in INM protein homeostasis.
**Introduction**

The organization of eukaryotic chromosomes within the cell nucleus depends on regulated and dynamic interactions with the nuclear envelope (NE). These interactions influence a myriad of cellular processes, from gene regulation and repair to cell motility and fate (Mekhail and Moazed, 2010). Chromosome interactions involve primarily proteins at the inner nuclear membrane, which together with the outer nuclear membrane (ONM), forms the NE. While continuous with the remaining ER membrane, the INM has a distinct set of proteins (Pawar and Kutay, 2021). Mutations in INM proteins are associated with a wide range of diseases, such as muscular dystrophies and premature aging syndromes, underscoring their importance for nuclear architecture and function.

Among the diverse INM proteome, Sad1p/UNC84 (SUN) domain-containing proteins SUN1 and SUN2 are unique as they couple nuclear and cytoplasmic events via LINC (Linker of nucleoskeleton and cytoskeleton) complexes (Crisp et al., 2006; Hodzic et al., 2004; Padmakumar et al., 2005). These molecular bridges across the NE are mediated by interactions of the luminal domains of SUN1 or SUN2 homotrimers in the INM and Nesprin proteins at the ONM (Crisp et al., 2006; Padmakumar et al., 2005; Sosa et al., 2012). Other domains of SUN and Nesprin proteins engage with various components in the nucleus and the cytoskeleton respectively, conferring the LINC complex with diverse functions. Indeed, by transducing forces from the cytoskeleton into nucleus, SUN1 and SUN2 have been implicated in controlling nuclear positioning within the cell (Lei et al., 2009; Zhang et al., 2009), NE anchoring of specific chromatin loci, facilitating NE disassembly during mitosis (Turgay et al., 2014) and DNA repair (Lei et al., 2012; Lottersberger et al., 2015). SUN proteins have also
been implicated in HIV replication and trafficking within host cells (Donahue et al., 2016). In these various functions, SUN1 and SUN2 appear to play both redundant and unique roles. However, our understanding of SUN proteins and the mechanisms by which they contribute to nuclear organization are far from complete.

Recent work in budding yeast showed that protein homeostasis at the INM relies on ubiquitin–dependent protein degradation (Foresti et al., 2014; Khmelinskii et al., 2014). This process is mainly mediated by a INM localized ubiquitin ligase complex, the Asi complex, which promotes ubiquitination of damaged and mislocalized membrane proteins (Foresti et al., 2014; Khmelinskii et al., 2014; Natarajan et al., 2020). Proteins ubiquitinated by the Asi complex are subsequently extracted from the INM into the nucleoplasm by a conserved ATPase complex called Cdc48 in yeast and p97 in mammals and delivered to the proteasome for degradation. Mechanistically, Asi-mediated protein degradation is similar to the ER-associated protein degradation (ERAD), a vital quality control process in ER membranes exposed to the cytosol (Christianson and Carvalho, 2022; Wu and Rapoport, 2018). In fact, the yeast ERAD ubiquitin ligase Doa10 also localizes to the INM and degrades both nucleoplasmic and INM proteins (Boban et al., 2014; Deng and Hochstrasser, 2006; Swanson et al., 2001). Despite the conservation of the ERAD process across all eukaryotes, sequence-based homologues of the Asi complex appear to be circumscribed to certain fungi.

While a functional counterpart of the yeast Asi complex has not been identified in higher eukaryotes, there are hints that the ubiquitin system influences protein homeostasis at the INM in mammals. Disease-associated
mutations in the Lamin-B receptor (LBR) appear to be removed from the INM through a process involving ubiquitination (Tsai et al., 2016). Perturbations of NE homeostasis, such as mutations in Torsin ATPase, result in INM herniations enriched in polyubiquitin conjugates (Laudermilch et al., 2016). Proteomics studies suggest links between the Skp/Cullin1/F-Boxβ-TrCP1/2 (SCF β-TrCP1/2) ubiquitin ligase complex and the INM protein SUN2 (Coyaud et al., 2015; Huttlin et al., 2021, 2017; Loveless et al., 2015; Low et al., 2014; Sowa et al., 2009). The INM protein Emerin has also been shown to interact with ubiquitin ligases (Khanna et al., 2018). However, the significance of these observations and the overall role of protein degradation to INM protein homeostasis remain largely unexplored.

Here we show that SUN2 levels at the INM are controlled by an ERAD-like mechanism. This process involves SUN2 ubiquitination by the SCFβ-TrCP ubiquitin ligase, membrane extraction by the p97 ATPase complex and degradation by the proteasome. We further show that the opposing activities of Casein Kinase II and the C-terminal domain Nuclear Envelope Phosphatase 1 (CTDNEP1) regulate SUN2 degradation. This kinase/phosphatase balance regulates SCFβ-TrCP binding to a non-canonical recognition motif in the SUN2 nucleoplasmic domain that is required for subsequent SUN2 ubiquitination. Finally, we show that accumulation of non-degradable SUN2 results in defects in nuclear envelope architecture that impact nuclear functions.
Results

**SUN2 stability depends on a non-canonical recognition motif for the SCFβTrCP ubiquitin ligase**

The SCFβTrCP ubiquitin ligase promotes ubiquitination and degradation of substrates containing defined recognition motifs (Frescas and Pagano, 2008). To study the relation between SUN2 and SCFβTrCP, we searched the SUN2 nucleoplasmic region for potential SCFβTrCP binding motifs. While the canonical DSGXXS recognition motif is absent, SUN2 contains two regions with related amino acid sequences, hereafter called Site 1 and Site 2, respectively. Importantly, Sites 1 and 2 are conserved across vertebrates and are absent in SUN1, a SUN2 homologue (Figure 1- figure supplement 1A-C).

Binding of SCFβTrCP to canonical and non-canonical sites on substrates requires their prior phosphorylation at two consensus serine residues (Frescas and Pagano, 2008). To test the potential involvement of Site 1 and Site 2 in SUN2 degradation, the critical consensus serine residues were mutated to alanine to abolish phosphorylation, and to aspartate to mimic constitutive phosphorylation (Figure 1A). Wild-type SUN2 (hereafter called SUN2 WT) as well as Site 1 and Site 2 mutants were stably integrated at the single FRT site of HEK293 TReX Flp-In (HEK TF) cell lines under a doxycycline/tetracycline-inducible promoter. The various SUN2 constructs were expressed as C-terminal fusions to superfolder green fluorescent protein (GFP) and hemagglutinin (HA) tags for easy detection (Figure 1B). Upon doxycycline (Dox) induction, there was robust detection of SUN2 WT by flow cytometry. Alanine mutations in Site 1 or Site 2 (Site 1A or Site 2A, respectively) resulted in even higher steady state SUN2 levels (Figure 1C). This increase was particularly noticeable for Site 2A mutant and was
further increased by combining alanine mutations in both sites. Conversely, phosphomimetic aspartate mutations at Site 1 and 2 (Site 1\textsuperscript{D} and Site 2\textsuperscript{D}, respectively) resulted in much lower SUN2 steady state levels, with the Site 2\textsuperscript{D} mutant displaying again a stronger effect (Figure 1D). Combination of aspartate mutations in both sites (Site 1\textsuperscript{D},2\textsuperscript{D}) resulted in further decrease in SUN2 steady state levels, suggesting that phosphorylation at Sites 1 and 2 have an additive effect. Since all SUN2 derivatives are expressed from the same promoter, these data suggest that SUN2 stability is controlled by non-canonical SCF\textsuperscript{βTrCP} recognition motifs, with Site 2 having a more prominent contribution.

To directly assess the effect of these mutations on SUN2 degradation, we performed chase experiments upon inhibition of protein translation with cycloheximide (CHX). We observed that SUN2 WT was degraded with a half-life of ~240 min (Figure 1E). Consistent with the analysis at steady state, SUN2 Site 2\textsuperscript{A} mutant was stable while the phosphomimetic aspartate mutation dramatically accelerated SUN2 degradation. SUN2 degradation was further accelerated by combining aspartate mutations at Sites 1 and 2 (Figure 1E). Interestingly, turnover of endogenous SUN2 was influenced and mirrored the turnover of the various transgenes (Figure 1E). Given that SUN2 functions as a trimer (Sosa et al., 2012), this observation likely reflects assembly of endogenous and transgenic molecules into trimeric SUN2 complexes. In contrast, the SUN2 homologue SUN1 was stable under all the conditions tested. A small but reproducible increase in SUN1 steady state levels was observed in cells expressing unstable SUN2 variants. These data confirmed our initial observations and showed that SUN2 degradation is controlled by non-canonical SCF\textsuperscript{βTrCP} recognition motifs.
INM degradation of SUN2 by an ERAD-like process

To test the role of the ubiquitin ligase SCF^{βTrCP} in SUN2 degradation, its activity was acutely blocked with MLN4924. This well-characterized small molecule inhibits Cullin RING ligases like SCF^{βTrCP} by preventing their activation by neddylation (31). Acute SCF^{βTrCP} inhibition with MLN4924 resulted in the stabilization of both WT (Figure 2A, 2B) and Site 2^\text{D} (Figure 2A, 2C) SUN2, whereas the steady state levels of the stable SUN2 Site 2^\text{A} mutant were unaffected (Figure 2A). Given that MLN4924 is a general inhibitor of Cullin RING ligases, the contribution of SCF^{βTrCP} in SUN2 degradation was directly tested upon depletion of the F-Box proteins βTrCP1 and βTrCP2. Depletion of βTrCP1 did not affect the levels of SUN2 Site 2^\text{D}, while βTrCP2 depletion resulted in increased steady state levels of this short-lived SUN2 mutant (Figure 2- figure supplement 1A-B). Further increase in SUN2 Site 2^\text{D} levels was observed when βTrCP1 and βTrCP2 were depleted simultaneously showing that they are redundant in promoting SUN2 degradation. Depletion of other F-Box proteins, such as FBXO2 and FBXO6, had no effect on SUN2 Site 2^\text{D} levels indicating that the effect observed for βTrCP1 and 2 is specific (Figure 2- figure supplement 1A-B). These data indicate that SCF^{βTrCP} ubiquitin ligase promotes SUN2 degradation and highlight the importance of Site 2 in the process.

Next, we tested the role of the p97 ATPase complex, which extracts ubiquitinated proteins from the membrane during ERAD (Christianson and Carvalho, 2022; Wu and Rapoport, 2018). Inhibition of p97 with CB-5803 also resulted in stabilization of both SUN2 WT (Figure 2A) and phosphomimetic mutant Site 2^\text{D} (Figure 2A-C). Moreover, SUN2 was stabilized upon inhibition of the proteasome with Bortezomib (Figure 2A-C), while inhibition of lysosomal protein degradation with Bafilomycin A
had no effect on SUN2 levels (Figure 2C). Similar results were obtained when the levels of phosphomimetic mutant Site 2^p were analysed by immunofluorescence (Figure 2D). Together, our data suggest SUN2 is degraded via an ERAD-like process involving the ubiquitin ligase SCF\(\beta\text{TrCP}\), the p97 ATPase complex and the proteasome.

**Site 2 is critical for SUN2 recognition and ubiquitination by SCF\(\beta\text{TrCP}\)**

Analysis of protein degradation suggests a critical role of Site 2 in SUN2 recognition and ubiquitination by SCF\(\beta\text{TrCP}\). To directly analyse the contribution of Sites 1 and 2 in SUN2 binding to SCF\(\beta\text{TrCP}\) we used immunoprecipitation. We found that endogenous \(\beta\)-TrCP coprecipitated with SUN2 WT. SUN2 binding to \(\beta\)-TrCP was largely unaffected by Site 1^A and 1^D mutations. In contrast, \(\beta\)-TrCP binding to SUN2 was lost in cells expressing the Site 2^A mutant, which cannot be phosphorylated (Figure 3A). Conversely, the amount of \(\beta\)-TrCP coprecipitated by the phosphomimetic SUN2 Site 2^p was greatly increased, even if Site 2^p levels were much lower. The amount of \(\beta\)-TrCP coprecipitated with SUN2 WT and Site 2^p was further increased in cells treated with the p97 inhibitor CB-5083, suggesting that inhibition of SUN2 degradation stabilized its interaction with the SCF\(\beta\text{TrCP}\) ubiquitin ligase. Under these conditions, neddyalted Cullin-1 also coprecipitated, indicating that SUN2 interacts primarily with the active pool of SCF\(\beta\text{TrCP}\) (Figure 3A).

Next, we used immunoprecipitation under denaturing conditions to analyze SUN2 ubiquitination. Ubiquitinated SUN2 WT was readily detected in unperturbed cells and the amount of ubiquitin conjugates increased in cells treated with inhibitors of p97 ATPase or the proteasome, which hinder protein retrotranslocation or degradation, respectively (Figure 3B).
Mutations in Site 1 resulted in levels of ubiquitin conjugates comparable to SUN2 WT. Consistent with its fast turnover, Site 2D mutant displayed increased ubiquitin conjugates, which became even more prominent upon p97 or proteasome inhibition. Importantly, ubiquitin conjugates to SUN2 WT and its derivatives were virtually undetectable upon inhibition of SCFβTrCP with MLN4924 (Figure 3B). Irrespective of the conditions used, ubiquitin conjugates were absent on SUN2 Site 2A (Figure 3B). Thus, SUN2 ubiquitination correlates with its ability to interact with SCFβTrCP. Altogether, these data indicate SUN2 Site 2 is a bona fide SCFβTrCP binding motif, recognized by βTrCP in a phosphorylation-dependent manner.

**Genome-wide screening identifies Casein Kinase 2 as a positive regulator of SUN2 degradation**

To gain further insight on the mechanism of SUN2 degradation, we performed a genome-wide CRISPR-Cas9-based genetic screen using the short lived SUN2 Site 2D mutant as a reporter. HEK TF cells with the SUN2 Site 2D-GFP-HA transgene were transduced with the Toronto KnockOut (TKO) lentivirus library consisting of an average of 4 sgRNAs to each of the 18,053 human genes and additional control sgRNAs (Hart et al., 2017). Mutant cells with high levels of GFP (top 1%), suggestive of impaired degradation of SUN2 Site 2D-GFP-HA, were isolated by flow cytometry at days 9, 13 and 17 after transduction as described (van de Weijer et al., 2020). Analysis of multiple timepoints allows identification of mutations in essential genes, which manifest early but drop out from later timepoints (Hart et al., 2015). Next-generation sequencing was used to sequence and quantify sgRNAs in reference and sorted cell populations (Figure 4A and Figure 4- figure supplement 1A). Gene rankings were generated using the MAGeCK algorithm (Li et al., 2014). Consistent with our earlier results, the
F-BOX protein βTrCP2 (encoded by FBXW11) and other core and regulatory subunits of the SCFβTrCP ubiquitin ligase, the p97 ATPase complex and the proteasome were among the highest scoring hits in our screen (Figure 4B). Independent sgRNAs were used to further validate some of these hits (Figure 4- figure supplement 1A-B) and the results demonstrated the suitability of our screen in identifying modulators of SUN2 degradation.

Another top hit from the screen was the Casein Kinase 2 β subunit (CSNK2B). As observed for other essential genes, the CSNK2B score was particularly high at day 9 and dropped at later timepoints (Figure 4B and Figure 4- figure supplement 1A). One of the genes encoding for Casein Kinase 2 α subunit (CSNK2A2) was also a hit in the screen although with a much lower score (Figure 4- figure supplement 1A). This is consistent with Casein Kinase 2 (CK2) functioning as a dimer of α-β subunit dimers, with CSNK2B being the sole gene encoding for the β subunit, while the α subunit is encoded by three largely redundant isoforms (Venerando et al., 2014).

Importantly, these data raise the possibility that CK2 promotes SUN2 phosphorylation thereby facilitating its recognition and ubiquitination by SCFβTrCP ubiquitin ligase. Consistent with this model, both CK2 and SCFβTrCP ubiquitin ligase were show to have nuclear localization thereby being able to access SUN2 at the INM.

The role of CSNK2B in SUN2 degradation was validated using additional sgRNAs (Figure 4- figure supplement 1C). Given that the depletion of essential genes, like CSNK2B, is complicated and can give confounding effects, we used a well-characterized CK2 inhibitor, tetrabromocinnamic acid (TBCA), to study the role of CK2 in SUN2 degradation. Consistent with
the gene depletion experiments, acute CK2 inhibition with TBCA resulted in an increase of the steady levels of both SUN2 WT and Site 2D mutant (Figure 4C, D). SUN2 WT steady levels were comparable between cells subjected to CK2 or p97 inhibition (Figure 4C, D). Importantly, phosphomimetic mutation in SUN2 Site 2D showed an intermediate response, while the double mutant SUN2 Sites 1D,2D was largely refractory to CK2 inhibition (Figure 4C, D). Thus, CK2 inhibition slows down SUN2 degradation.

Next, we used immunoprecipitation to test whether CK2 regulates SUN2 degradation by controlling its interaction with the ubiquitin ligase SCFβTrCP. Indeed, acute CK2 inhibition with TBCA decreased the interaction between SUN2 and βTrCP (Figure 4E). Together, these data indicate that CK2 activity promotes SCFβTrCP binding to SUN2 thereby facilitating its ubiquitination and degradation.

The phosphatase CTDNEP1 is a negative regulator of SUN2 degradation

Under normal conditions, SUN2 is a relatively stable protein (Figure 5C) (Buchwalter et al., 2019). Therefore, we hypothesized that CK2-dependent SUN2 degradation is counteracted by the activity of a protein phosphatase. Given its localization to the INM and phosphatase activity towards Serine/Threonine residues, the C-terminal Domain Nuclear Envelope Phosphatase 1 (CTDNEP1) was an attractive candidate to regulate SUN2 turnover (Bahmanyar et al., 2014; Kim et al., 2007). To test this possibility, we analysed the levels of endogenous SUN2 upon CTDNEP1 depletion. Consistent with our hypothesis, the steady state levels of endogenous SUN2 and of SUN2 WT transgene were strongly reduced in CTDNEP1 depleted
cells (Figure 5A, B and D). This effect was specific to SUN2 as the levels of other INM proteins such as SUN1 (Figure 5A, B) and LBR (Figure 5- figure supplement 1A) remained unchanged. In CTDNEP1 KO cells, the lower levels of endogenous SUN2 (Figure 5C) and SUN2 WT transgene (Figure 5- figure supplement 1B) were due to their accelerated degradation, as monitored by cycloheximide chase experiments. Importantly, increased turnover of endogenous SUN2 (Figure 5- figure supplement 1C and D) and transgenic SUN2 WT (Figure 5D) in CTDNEP1 depleted cells was reversed by inhibition of the SCFβTrCP ubiquitin ligase, the p97 ATPase and the proteasome. Moreover, regulation of SUN2 turnover required CTDNEP1 phosphatase activity. As shown in Figures 5E and Figure 5- figure supplement 1E, re-expression of wild-type CTDNEP1 (CTDNEP1-Flag) but not of a catalytically inactive CTDNEP1 (CTDNEP1(PD)-Flag) mutant in CTDNEP1 KO cells restored the levels of both endogenous and transgenic SUN2. Thus, the nuclear envelope phosphatase CTDNEP1 is a negative regulator of SUN2 degradation.

Together with our earlier results, these data also suggest a model in which SUN2 levels are controlled by a phospho-switch, with the kinase CK2 promoting SUN2 degradation and the phosphatase CTDNEP1 counteracting it. In support of this model, the reduction of SUN2 levels in CTDNEP1 KO cells is counteracted by acute CK2 inhibition with TBCA (Figure 5F and Figure 5- figure supplement 1F). A second prediction of the model is that the SUN2 Site 2A, which cannot be phosphorylated in the main site controlling SUN2 degradation, is insensitive to the loss of CTDNEP1. Indeed, depletion of CTDNEP1 while resulting in low levels of SUN2 WT transgene, did not affect SUN2 Site 2A mutant level (Figure 5G and Figure 5- figure supplement 1G). Importantly, the levels of endogenous SUN2 were...
still reduced in these cells, confirming the depletion of CTDNEP1 (Figure 5G). Therefore, CK2 and CTDNEP1 have opposing effects on SUN2 turnover.

**Aberrant nuclear architecture and function by expression of non-degradable SUN2**

Next, we investigated the importance of SUN2 regulated degradation for nuclear morphology and function. To this end, we analysed nuclear morphology in HeLa cells expressing either SUN2 WT or the non-degradable SUN2 Site2A mutant. In parental HeLa cells, most nuclei display a smooth or slightly wrinkled surface. Expression of SUN2 WT from a Dox-inducible promoter for 24 hours increased the fraction of wrinkled nuclei as expected (Donahue et al., 2016). A small fraction of SUN2 WT expressing cells showed aberrant nuclei, which were highly convoluted and multi-lobed. Nearly half of the cells expressing non-degradable SUN2 Site2A displayed these highly aberrant nuclei, which were never observed in parental HeLa cells (Figure 6B). Importantly, nuclear morphology greatly improved upon shutting off expression of SUN2 WT for 24 hours. In contrast, aberrant nuclear morphologies persisted even 24 hours after shutting-off SUN2 Site 2A expression, consistent with the long half-life of this protein (Figure 6A, B and Figure 6- figure supplement 1A). In the aberrant nuclei, SUN2 Site 2A was distributed heterogeneously, accumulating in patches or lines. Besides SUN2 Site 2A, these structures were enriched in certain INM proteins, such as LBR and EMD. However, other INM proteins like LAP 2β and SUN1 were unaffected (Figure 6C). Thus, accumulation of non-degradable SUN2 drives aberrant nuclear envelope architecture and redistribution of specific INM proteins.
SUN2 was implicated in facilitating nuclear envelope disassembly during mitosis (Turgay et al., 2014). We wondered if the changes in nuclear architecture due to accumulation of non-degradable SUN2 Site 2A would impact mitotic progression. Timelapse fluorescence microscopy showed that cells expressing SUN2 WT and Site 2A progress through the various stages of mitosis with normal kinetics (Figure 6- figure supplement 1B). In particular, nuclear envelope breakdown, chromosome alignment at the metaphase plate and anaphase onset occurred with similar timings. In addition, the frequency of multipolar spindles was low in both cases (Figure 6- figure supplement 1B). However, cells expressing SUN2 Site 2A showed higher frequency of lagging chromosomes during anaphase (Figure 6- figure supplement 1C-D). Curiously, these cells also showed an increased association of chromosomes with the nuclear membrane (Figure 6- figure supplement 1D). Thus, accumulation of non-degradable SUN2 may impact chromosome dynamics, potentially by interfering with interactions between chromosomes and the INM.

SUN2 has also been implicated in DNA damage repair, particularly in the repair of double strand breaks (Lei et al., 2012; Lottersberger et al., 2015). While the precise role of SUN2 in DNA repair is ill defined, it may control the mobility of damaged loci thereby affecting the efficiency of the repair process (Lottersberger et al., 2015). Therefore, we asked whether accumulation of non-degradable SUN2 Site 2A impacts the cell’s ability to repair DNA double strand breaks. To assess the efficiency of DNA repair we monitored the DNA damage marker γH2AX (Ser139-phosphorylated histone H2AX) (Rogakou et al., 1998) in U2OS cells subjected to ionising radiation (IR) to trigger double strand breaks. Parental and SUN2 Site 2A expressing cells have comparable levels of γH2AX both prior and soon after
IR treatment (Figure 6D-E). This indicates that expression of non-degradable SUN2 is (1) insufficient to trigger DNA damage; and (2) does not prevent cells from mounting a DNA damage response upon induction of double strand breaks with IR. At 24 hours post IR, parental cells show nearly basal levels of γH2AX indicating that DNA damage has been repaired. In contrast, the levels of γH2AX in SUN2 Site2A are higher 24 hours post IR, likely due to unresolved DNA damage. Thus, accumulation of non-degradable SUN2 Site 2A appears to compromise efficient DNA repair.

Altogether, these data indicate that accumulation of non-degradable SUN2 affects nuclear architecture and impacts nuclear functions.
Discussion

We uncovered that the INM protein SUN2, a subunit of the LINC complex, undergoes regulated turnover (Figure 7). The kinase CK2 and the phosphatase CTDNEP1 act respectively, as positive and negative regulators of SUN2 degradation by influencing the phosphorylation state of non-canonical binding sites for the SCF$^{βTrCP}$ ubiquitin ligase, primarily Site 2. Binding of SCF$^{βTrCP}$ to phosphorylated Site 2 results in SUN2 ubiquitination and the recruitment of the p97 ATPase complex, which facilitates SUN2 membrane extraction and delivery to the proteasome for degradation. Cells expressing non-degradable SUN2 display defects in nuclear architecture highlighting the importance of this ERAD-like process in maintaining INM protein homeostasis in mammalian cells.

While SUN2 and the LINC complex play important functions in cell homeostasis, conditions exist in which the forces they transduce into the nucleus are deleterious. It has been shown that aberrant DNA repair resulting from increased chromatin mobility at sites of double strand breaks depends on the LINC complex (Lottersberger et al., 2015). Consistent with these data, we observe that expression of non-degradable SUN2 results in impaired repair of double strand breaks. Downregulation of the LINC complex resulting in the decoupling of the nucleus from the cytoskeleton has been observed in cells under extreme cycles of mechanical load (Gilbert et al., 2019). This nuclear decoupling protects DNA from mechanical-induced damage and appears to involve a decrease in SUN2 levels by some unknown mechanism. Thus, it is appealing to consider that downregulation of LINC complexes by localized SUN2 degradation at sites of DNA damage or extreme mechanical load may facilitate DNA repair or
prevent mechanical-induced damage. Future studies should test these possibilities directly.

Mechanistically, the involvement of a soluble ubiquitin ligase, SCFβTrCP, in SUN2 degradation presents an important difference to conventional ERAD, which is normally mediated by transmembrane ubiquitin ligases (Christianson and Carvalho, 2022). However, the participation of SCFβTrCP in an ERAD-like process is not unprecedented. In HIV infected cells, SCFβTrCP promotes the degradation of the integral membrane protein CD4 (Florence Margottin et al., 1998). In this case, virally-encoded Vpu protein functions as an adaptor, binding to the substrate CD4 in the ER membrane and recruiting the ubiquitin ligase SCFβTrCP via the cytosolic tail (Magadán et al., 2010; F Margottin et al., 1998; Zhang et al., 2013). Based on these observations, it will be interesting to test whether SCFβTrCP has a more widespread role in the degradation of membrane proteins. Interestingly, a soluble ubiquitin ligase is also implicated in downregulating the yeast SUN-domain containing protein Mps3 by an ERAD-like process (Koch et al., 2019). In this case, the degradation of Mps3, which is important for the distribution of nuclear pore complexes and spindle pole bodies (the yeast equivalent to centrosomes) within the INM (Chen et al., 2014; Jaspersen et al., 2006), facilitates cell cycle progression. Like in the case of SUN2, accumulation of non-degradable Mps3 affects nuclear morphology (Koch et al., 2019).

Extraction of ERAD substrates from the membrane into the cytosol, a step known as retrotranslocation, is driven by the pulling force of p97 ATPase and facilitated by transmembrane segments of ERAD ubiquitin ligases or accessory factors (Christianson and Carvalho, 2022; Wu and Rapoport, 2018). In the case of a soluble ubiquitin ligase like SCFβTrCP, how membrane
substrates are retrotranslocated is unclear. Our genome-wide CRISPR screen did not identify any membrane proteins that could potentially mediate SUN2 retrotranslocation suggesting redundancy between multiple components. Alternatively, the pulling force imposed by p97 may be sufficient to extract SUN2 from the INM membrane.

Regulated protein degradation by ERAD is well documented, particularly in the control of sterol biosynthesis (Christianson and Carvalho, 2022; Johnson and DeBose-Boyd, 2018). In these cases, the degradation of rate limiting enzymes depends on the concentration of specific sterol metabolites in the ER membrane and functions as a feedback mechanism controlling sterol biosynthesis. To our knowledge, SUN2 is the first substrate shown to be controlled by a kinase/phosphatase balance. Given the requirement of phosphorylation for SCFβTrCP binding to substrates, the involvement of a kinase was expected (Frescas and Pagano, 2008). In fact, CK2 has been implicated in the degradation of other SCFβTrCP substrates such as Vpu-mediated degradation of CD4 (Florence Margottin et al., 1998) and Cyclin F (Mavrommati et al., 2018). In contrast, the involvement of CTDNEP1 is unexpected as this phosphatase has been mainly implicated in dephosphorylation of Lipin, a key regulator of lipid metabolism (Barger et al., 2022; Zhang and Reue, 2017). In particular, CTDNEP1 was recently implicated in coordinating membrane biogenesis with cell division and loss of CTDNEP1 results in excess membrane production which affect chromosome dynamics (Merta et al., 2021). A role for SUN2 in lipid metabolism was also recently proposed (Lee et al., 2022), suggesting that the convoluted nuclei observed in cells expressing non-degradable SUN2 may result from excessive production of NE membranes. Thus, future studies should explore further how the function of CTDNEP1 in controlling
SUN2 levels relates to its role in lipid regulation. Another important open question is whether CTDNEP1 and CK2 act directly on SUN2.

While studies in rats and cultured cells suggest that INM proteins are generally long lived (Buchwalter et al., 2019; Toyama et al., 2013), our results show that the INM proteome can be remodelled in a highly regulated fashion. Other recent examples show that certain conditions, like mutations (Tsai et al., 2016) or ER stress (Buchwalter et al., 2019), can trigger the turnover of specific INM proteins. Thus, it appears that the turnover of INM proteins is likely to be temporal and spatially restricted, allowing fine-tuning of INM properties without jeopardizing its overall structure and critical functions in nuclear organization.
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Declaration of Interests

The authors declare no competing interests.
Figure Legends

Figure 1: SUN2 stability depends on a non-canonical recognition motif for the SCF$_{\betaTrCP}$ ubiquitin ligase

(A) SUN2 nucleoplasmic regions containing non-canonical recognition motifs for the SCF$_{\betaTrCP}$ hereafter called Sites 1 and 2. In each case, the residues mutated to Alanine (purple) or to Aspartate (red) are indicated. The canonical SCF$_{\betaTrCP}$ recognition motif (DSGXXS) is indicated. SCF$_{\betaTrCP}$ binding occurs only to serine residues (Cyan) that are phosphorylated.

(B) Schematic representation of SUN2-GFP-3HA construct used in most experiments. Sites 1 and 2 in the SUN2 nucleoplasmic region are indicated in Cyan.

(C, D) Flow cytometry analysis of doxycycline-induced expression of SUN2 WT or derivatives in HEK TF cells. Analysis was performed 24 h after SUN2 expression.

(E) Analysis of the stability of SUN2 WT and the indicated mutants after inhibition of protein synthesis by cycloheximide (CHX). Cell extracts were analyzed by SDS–PAGE and immunoblotting. Transgenic WT and mutant SUN2 were detected with anti-HA antibody. Endogenous SUN2 was detected with anti-SUN2 antibody. This antibody recognizes the C-terminal peptide of SUN2 and is deficient in recognizing SUN2 if tagged C-terminally. Endogenous SUN1 was detected with anti-SUN1 antibody. Tubulin was used as a loading control and detected with an anti-Tubulin antibody.

Figure 2: INM degradation of SUN2 by an ERAD-like process

(A) Flow cytometry analysis of tetracycline-induced expression of SUN2 WT, Site 2$^A$ and Site 2$^B$ in HEK TF cells. Analysis was performed 24 h post-
induction in cells left untreated, incubated 4 h with inhibitors to SCFβTrCP
neddylation (MLN-4924; 1 µM), p97 (CB-5083; 2.5 µM, CB) or the
proteasome (Bortezomib; 1 µM, Btz).

(B) Analysis of SUN2 WT steady state levels in HEK TF cells. Analysis was
performed 24 h post-induction in cells left untreated, incubated 4 h with
inhibitors to SCFβTrCP neddylation (MLN-4924; 1 µM), p97 (CB-5083; 2.5
µM, CB) or the proteasome (Bortezomib; 1 µM, Btz). Cell extracts were
analyzed by SDS–PAGE and immunoblotting. SUN2 WT was detected with
anti-HA antibodies. GAPDH was used as a loading control and detected with
an anti-GAPDH antibody.

(C) Analysis of SUN2 Site 2D steady state levels in HEK TF cells. Analysis
was performed 24 h post-induction in cells incubated 4 h with DMSO
(vehicle) or the inhibitors to SCFβTrCP neddylation (MLN-4924; 1 µM), p97
(CB-5083; 2.5 µM, CB), the proteasome (Bortezomib; 1 µM, Btz and 10 µM
MG132) or incubated 6 h with bafilomycin A (1 µM, Baf) that inhibits
lysosomal delivery. Cell extracts were analyzed by SDS–PAGE and
immunoblotting. SUN2 Site 2D was detected with anti-HA antibodies. LC3B
was analyzed to confirm effectiveness of bafilomycin A treatment and was
detected with an anti-LC3B antibody. GAPDH was used as a loading control
and detected with an anti-GAPDH antibody.

(D) Immunofluorescence in HeLa cells expressing SUN2 Site 2D. Expression
of SUN2 Site 2D was induced with Dox for 24h and incubated for 4 h with
DMSO (vehicle) or the inhibitors to SCFβTrCP neddylation (MLN-4924; 1 µM),
p97 (CB-5083; 2.5 µM, CB), the proteasome (Bortezomib; 1 µM, Btz). SUN2
Site 2D was detected with anti-HA antibodies. The ER marker Calnexin was
detected with an anti-Calnexin antibody and DNA was labelled with DAPI.
Scale bar; 10 µm.
Figure 3: Site 2 is critical for SUN2 recognition and ubiquitination by SCFβ-TrCP

(A) Binding of WT and the indicated SUN2 mutants to subunits of the ubiquitin ligase SCFβ-TrCP and the ATPase p97 analyzed by immunoprecipitation. Expression of SUN2 derivatives was induced with Dox for 24h and treated with DMSO (-) or CB-5083 (+) for 4 h before cell lysis and immunoprecipitation with anti-HA coated beads. Eluted proteins were separated by SDS-PAGE and analyzed by western blotting with the indicated antibodies.

(B) Ubiquitination of WT and the indicated SUN2 mutants in cells incubated 4 h with DMSO (vehicle) or the inhibitors to SCFβ-TrCP neddylation (MLN-4924; 1 µM), p97 (CB-5083; 2.5 µM, CB), the proteasome (Bortezomib; 1 µM, Btz). Upon immunoprecipitation with anti-HA coated beads, eluted proteins were separated by SDS-PAGE and analyzed by western blotting with anti-HA and anti-ubiquitin antibodies.

Figure 4: Genome-wide screening identifies Casein kinase 2 as a positive regulator of SUN2 degradation

(A) Workflow of the CRISPR-Cas9 genome-wide screen.

(B) Significance score of the genes analyzed in the screen calculated by the MAGeCK algorithm. The x-axis represents the genes in alphabetical order. The y-axis shows the −log(αRRA) significance value. The −log(αRRA) cut-off was arbitrarily set at 5 (dashed line). Significantly enriched genes are annotated and colour-coded: genes related to SCF ubiquitin ligase core subunits (blue), NEDD8 conjugation cascade (red), ubiquitin conjugation...
cascade (orange), p97 and co-factors (green) and kinase (pink).

Proteasome subunits are shown in a grey dotted box.

(C-D) Flow cytometry (C) and western blotting (D) analysis of Dox-induced expression of WT, Site 2A, Site 2D and Site 1D, 2D SUN2 in HEK TF cells. Analysis was performed 24 h post-induction in cells incubated 6 h with inhibitors to Casein Kinase 2 (TBCA; 100 µM) and p97 (CB-5083; 2.5 µM, CB). SUN2 derivatives were detected with anti-HA antibody. CK2 inhibition was confirmed by blotting with anti-phospho-CK2 substrates antibody. GAPDH was used as a loading control and detected with an anti-GAPDH antibody.

(E) Immunoprecipitation of SUN2 WT from cells incubated 6h with DMSO (vehicle), Casein Kinase 2 inhibitor (TBCA; 100 µM) and p97 inhibitor (CB-5083; 2.5 µM, CB). Cell lysates were immunoprecipitated with anti-HA coated beads. Eluted proteins were separated by SDS-PAGE and analyzed by western blotting with the indicated antibodies.

Figure 5: The phosphatase CTDNEP1 is a negative regulator of SUN2 degradation

(A) Steady state levels of endogenous SUN2 in HEK TF transfected with an empty vector (EV) or sgRNAs targeting CTDNEP1. Cell extracts were analyzed by SDS–PAGE and immunoblotting with anti- SUN2 and anti-SUN1 antibodies. Calnexin was used as a loading control and detected with an anti-Calnexin antibody.

(B) Immunofluorescence in parental and CTDNEP1 KO HeLa cells. Expression of SUN2 Site 2D was induced with Dox for 24h. Endogenous
SUN2 and SUN1 were detected with anti SUN2 and anti-SUN1 antibodies respectively. DNA was labelled with DAPI. Scale bar; 10µm.

(C) Analysis of endogenous SUN2 stability after inhibition of protein synthesis by cycloheximide (CHX) in parental and CTDNEP1 KO HEK TF cells. Cell extracts were analyzed by SDS–PAGE and immunoblotting. Endogenous SUN2 and SUN1 were detected with anti SUN2 and anti-SUN1 antibodies, respectively. Tubulin was used as a loading control and detected with an anti-Tubulin antibody. Three independent experiments were quantified on the right.

(D) Flow cytometry analysis of SUN2 WT in HEK TF cells transfected with an empty vector (EV) or with a sgRNA targeting CTDNEP1. Analysis was performed 24 h post- Dox induction in cells left untreated or incubated 4 h with inhibitors to SCFβTrCP neddylation (MLN-4924; 1 µM), p97 (CB-5083; 2.5 µM, CB) or the proteasome (Bortezomib; 1 µM, Btz).

(E) Analysis of transgenic (+ Dox; left) and endogenous SUN2 (No Dox; right) steady state levels in HEK TF SUN2 WT-expressing parental and CTDNEP1 KO cells transfected with an empty vector (EV), a vector encoding FLAG-tagged WT CTDNEP1 or phosphatase dead CTDNEP1 (PD). Exogenous SUN2 WT and endogenous SUN2 were detected with anti-HA and anti-SUN2 antibodies, respectively. CTDNEP1-FLAG was detected with anti-FLAG antibodies. GAPDH was used as a loading control and detected with an anti-GAPDH antibody.

(F) Flow cytometry analysis of SUN2 WT in CTDNEP1 KO HEK TF cells. Analysis was performed 24 h post-Dox induction in cells incubated 6h with DMSO (vehicle), Casein Kinase 2 inhibitor (TBCA; 100 µM) and p97 inhibitor (CB-5083; 2.5 µM, CB).
Analysis of transgenic (+Dox; left) and endogenous SUN2 (No Dox; right) steady state levels in HEK TF SUN2 WT or Site 2A cells transfected with an empty vector (EV) or with a sgRNA targeting CTDNEP1. Transgenic and endogenous SUN2 were detected with anti-HA and anti-SUN2 antibodies, respectively. GAPDH was used as a loading control and detected with an anti-GAPDH antibody. The abundance of SUN2 in relation to GAPDH was quantified and is shown below each lane.

Figure 6: Accumulation of non-degradable SUN2 results in aberrant nuclear architecture and function

(A) Micrographs of parental HeLa or cells expressing SUN2 WT or Site2A. Cells were analyzed 24h after inducing SUN2 expression (left) or 24h after turning off SUN2 expression (right). SUN2 was detected with GFP and DNA was labelled with DAPI.

(B) Quantification of nuclear morphology in HeLa cells as in (A). Three independent experiments, with 50-150 cells for each condition in each repeat were quantified. Representative examples of nuclear morphologies are shown on the right. Error bars indicate standard deviation.

(C) Immunofluorescence in HeLa cells upon induction of SUN2 Site 2A expression for 24h. SUN2 Site 2A was detected with GFP. Anti-LAP2β, -LBR, -EMD and -SUN1 antibodies were used to detect these INM proteins. DNA was labelled with DAPI.

(D) Immunofluorescence in U2OS parental and SUN2 Site 2A-expressing cells irradiated with 5Gy ionising radiation (IR). Time after irradiation is indicated. Non-irradiated cells were used as control (shown on the top).
SUN2 Site 2A expression was induced 24h prior irradiation. \( \gamma \)H2AX foci were detected with an anti-\( \gamma \)H2AX antibody. Scale bar; 10\( \mu \)m.

(E) Quantification of \( \gamma \)H2AX foci per nucleus detected in (D). Three independent experiments were analyzed and 30-40 nuclei were quantified for each condition in each replicate. Error bar represents standard error of the means (SEM). Ordinary one-way ANOVA and Tukey’s multiple comparisons were performed (**** p<0.0001).

**Figure 7: A kinase-phosphatase balance regulates ER-associated degradation of SUN2 from the INM**

The scheme illustrates the opposing effects of the kinase CK2 and the phosphatase CTDNEP1 on SUN2 degradation. Phosphorylation of SUN2 Site 2 promotes the binding of SCF\( ^{\beta TrCP} \) ubiquitin ligase and subsequent SUN2 ubiquitination, membrane extraction by the p97 ATPase complex and delivery to the proteasome for degradation. SUN2 is depicted as a monomer for simplicity however it functions as a homotrimer.
METHOD DETAILS

Plasmids
For stable integration of SUN2 constructs into the FRT sites of Flp-In T-Rex HEK cells, SUN2 cDNAs were cloned into the pcDNA5-FRT-TO plasmid (Invitrogen). For CTDNEP1-rescue experiments a lentiviral vector was used with the expression of CTDNEP1 being driven by the EF-1α promoter. For experiments in HeLa and U2OS cells, we used a lentiviral vector with an inducible doxycycline-responsive promoter driving the expression of SUN2 WT and variants. For gene knockouts, a lentiviral pSicoR plasmid with U6 promoter driving sgRNA expression and EF-1a promoter driving the expression of Puromycin-T2A-Flag-Cas9 was used.

Cell culture
All cells were grown at 37°C, 5% CO2 in DMEM medium (Sigma-Aldrich) supplemented with L-Glutamine (2 mM; GIBCO), Penicillin-Streptomycin (10 Units/mL; GIBCO) and 10% FCS (GIBCO). Cells were regularly tested for mycoplasma contamination.

Generation of Flp-In T-Rex HEK (HEK TF) cell lines
Cell lines were made as per manufacturer’s protocol.

Lentiviral production and transduction
Lenti-X cells were co-transfected with lentiviral and packaging plasmids using the Mirus LT1 transfection reagent in 24-well tissue culture plate format. 72 hours later, the media was harvested. For transduction, cells were seeded in a 24-well tissue culture plate (1.5x10⁵ cells for HEK and 7.5x10⁴ cells for HeLa). The next day, 200ul of lentivirus was added onto the cells. 24 hours later, the media was replaced and cells were grown for
further 24 hours. Cells were expanded to 10cm tissue culture-treated petri dish in the presence of antibiotic selection (zeocin or blastocidin for either the constitutive or the inducible lentiviral plasmids, respectively).

**Co-Immunoprecipitation**

Expression of SUN2 constructs in HEK293 TF cells was induced with 1ug/ml Doxycycline 24 hours prior to lysis. Cells were grown to 80% confluency in 6-well format, washed in Tris-buffered Saline (TBS; 50 mM Tris–HCl pH 7.5, 150 mM NaCl) once, and lysed in 1% DMNG (Anatrace) lysis buffer (50 mM Tris–HCl pH 7.5, 150 mM NaCl) containing cOmplete protease inhibitor cocktail (Roche) and 5mM N-Ethylmaleimide (NEM). Cell suspension was solubilised on a rotating wheel for 2 hours at 4°C. Cell debris and nuclei were pelleted at 20,000 g for 20 min at 4°C. Postnuclear supernatants were incubated for 2 hours with pre-equilibrated 20ul anti-HA antibody magnetic beads slurry (Sigma-Aldrich). After three 10min-washes in 0.2% DMNG washing buffer (50 mM Tris–HCl pH 7.5, 150 mM NaCl) on rotating wheel at 4°C, proteins were eluted from the beads in 1x sample buffer for 15 min at 65°C and collected into fresh tubes using magnetic racks. Eluates were subsequently supplemented with 100mM DTT.

**Substrate ubiquitination assay**

Expression of SUN2 constructs in HEK293 TF cells was induced with 1ug/ml Doxycycline 24 hours prior to lysis. Cells at around 80% confluence in a 10 cm dish were lysed in RIPA buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl, 1% Triton X-100, 0.5% Sodium Deoxycholate, 0.1% SDS) containing NEM (5mM) and cOmplete protease inhibitor cocktail (Roche). Cell suspension was solubilised on a rotating wheel for 2 hours at 4°C. Cell
debris and nuclei were pelleted at 20,000 g for 20 min at 4°C. Postnuclear supernatants were incubated for 2 hours with pre-equilibrated 30ul anti-HA antibody magnetic beads slurry (Sigma-Aldrich). After three 15min-washes in RIPA buffer on a rotating wheel at 4°C, proteins were eluted from the beads in 1x sample buffer for 15 min at 65°C and collected into fresh tubes using magnetic racks. Eluates were subsequently supplemented with 100mM DTT.

Translation shut-off experiments
Expression of SUN2 constructs in Flp-In TRex HEK cells was induced with 1ug/ml Doxycycline 24 hours prior to translation shut-off experiments. Cells were incubated with cycloheximide (50 μg/mL) for the indicated time points, after which cells were lysed in 1x sample buffer containing Benzonase (Sigma-Aldrich), cOmplete protease inhibitor cocktail (Roche), and DTT. Lysates were incubated for 30 min at 37°C, after which proteins were separated by SDS-PAGE. Immunoblotting was performed as described below. Representative images of at least three independent experiments are shown. Error bars represent the standard deviation.

Immunoblotting
Samples were incubated at 65°C for 10min, separated by SDS–PAGE (Bio-Rad) and proteins were transferred to PVDF membranes (Bio-Rad). Membranes were blocked in 5% Milk or BSA in PBS-Tween20 buffer and then probed with primary antibodies overnight at 4°C on shaker. Secondary antibodies were performed at RT for 1 hour either in 0.5% Milk or BSA in PBS-Tween20 buffer. Membranes were developed by ECL (Western Lightning ECL Pro, Perkin Elmer), and visualized using an Amersham Imager 600 (GE Healthcare Life Sciences).
**Immunofluorescence**

HeLa cells are seeded (7.5x10^4 cells) in DMEM supplemented with Pen/Strep onto round coverslips (13mm, #1.5) in a 12-well tissue culture-treated plate. 24 hours later, media was replaced to 1μg/ml Doxycycline-containing DMEM supplemented with Pen/Strep and incubated for further 24 hours. For Dox wash out experiments, 5.0x10^4 cells were seeded to allow for an additional day for Doxycycline washout protocol. Media was aspirated off and washed once with PBS. Cells were fixed with 4% PFA (methanol-free) for 10min at RT, followed by three PBS washes. Fixed cells were then treated with 10mM Glycine for 5min, followed by three PBS washes. Fixed cells were incubated in blocking buffer (BB; 1% BSA, 0.1% Saponin in PBS) for 30min. Primary antibodies cocktails were prepared in BB. For primary antibody staining, coverslip was carefully placed down onto pre-spotted antibodies cocktail (20ul/cover slip) on clean parafilm and incubated for 1 hour underneath home-made aluminum foil-covered moisturized chamber. Cover slips were washed three times with BB and secondary antibody staining was performed as was done for primary staining. Cover slips, then were incubated in DAPI-containing PBS for 5min and washed three times with PBS. Coverslips were mounted onto glass slide in non-hardening mounting media and sealed with nail polish.

**Confocal fluorescence microscopy**

Fixed cells on slides after immunofluorescence were imaged at 21°C using an inverted Zeiss 880 microscope fitted with an Airyscan detector using ZEN black software. The system was equipped with Plan-Apochromat 63×/1.4-NA oil lens, with an immersion oil (Immersol™ W 2010, Carl Zeiss; refractive index of 1.518). 488 nm argon and 405 nm, 561
nm and 633nm solid state diode lasers were used to excite fluorophores. Z-sections with 0.37μm-thick intervals were collected. The oil objective was covered with an immersion oil (ImmersolT 518 F, Carl Zeiss) with a refractive index of 1.518.

Microscopy images with CZI file format were analyzed using ImageJ (bundled with Java 1.8.0_172) software. Scoring of nuclear morphology was done after maximum intensity projection image processing.

**Live cell imaging**

Time-lapse imaging of paired GFP-SUN2 WT and Site 2^A^ cells was performed on a DeltaVision Elite light microscopy system. Fluorescence images were collected on a 512 × 512-pixel electron-multiplying charge-coupled device camera (QuantEM; Photometrics) using the software package softWoRx (GE Healthcare). Cells were seeded on two-chambered glass-bottom dishes (Lab-Tek) at 30,000 per well. Cell media was replaced with FluoroBrite DMEM media supplemented with 10 % fetal bovine serum, GlutaMAX and SiR-Hoechst at a final concentration of 100 nM 8 hours prior to imaging. For imaging, cells were placed in a 37°C and 5% CO2 environmental chamber (Tokai Hit). Per field of view, seven planes were captured 2 μm apart every 5 min, with light powers at 5% and 15 ms exposures. Maximum-intensity projections were performed using softWoRx, with image cropping and analysis performed using Fiji. Three independent experiments were conducted, with more than 45 cells per cell line being analyzed in each repeat. Statistical analysis and graphing were performed on GraphPad Prism v9.0. Initiation of prophase, nuclear envelope breakdown, metaphase (alignment) and anaphase were analyzed as described (Hayward et al., 2019). Cells with delayed chromosome
release from nuclear membrane were characterised as having one or more chromosomes at the nuclear periphery in the two timepoints following nuclear envelope breakdown.

**DNA damage assay and quantification of γH2AX foci**

U2OS parental and SUN2 Site 2A cells were seeded (7.5x10^4 cells) in DMEM supplemented with Pen/Strep onto round cover slips (16mm, #1) in 35mm tissue culture-treated dishes. SUN2 Site 2A expression was induced 24 hours before ionizing radiation (IR) with the addition of Doxycycline to a final concentration of 1ug/ml. A total dose of 5 Gy IR was performed to induce DNA double strand breaks. Cells were further incubated in Doxycycline-containing media for 1h, 4h, 8h and 24h. Control cells not exposed to IR were harvested after 24 hours of Doxycycline induction. Cells were fixed with 4% PFA (methanol-free) for 10min at RT, followed by three PBS washes. Fixed cells were then permeabilised with 0.2% Triton-X 100 in PBS for 10min, followed by three PBS washes. Fixed cells were blocked with PBS/10% FBS for 2 hours at 4°C. Primary antibody (Anti-γH2AX) were incubated overnight at 4°C in PBS/10% FBS (1:1000) in home-made humidified chambers, followed by 3 times wash with PBS/0.1% Triton-X 100 for 5min. Secondary antibody was prepared in PBS/0.15% FBS, and cells were incubated with secondary antibody at room temperature for 2 hours in the humidified chambers, followed by 3 times wash with PBS/0.1% Triton-X 100 for 5min and one 5min wash with PBS. Cover slips were then incubated in DAPI-containing PBS for 5min, washed three times with PBS, mounted onto glass slide in non-hardening mounting media and sealed with nail polish.
Quantification of γH2AX signal was performed using ImageJ (bundled with Java 1.8.0_172) software. Each nucleus was masked and the number of spots within the mask with size between 0.4-3 microns was counted as foci. More than 90 parental and SUN2 Site 2\textsuperscript{A} cells in each timepoint from three independent experiments were analysed, and statistical analysis was performed using One-way ANOVA method on GraphPad Prism v9.0.

**Generation CRISPR/Cas9-mediated gene knockout cells**

Cells were transfected with single bicistronic gRNA-Cas9 plasmid at 80% confluency with Mirus LT1 transfection reagent in 24-well tissue culture plate format. 48 hours later, cells were trypsinized and expanded into a 10cm tissue culture dish in the presence of 2ug/ml puromycin for 72 hours. Cells were maintained in fresh media (without puromycin) for additional 72 hours before validation by western blotting. Knockout clonal cell lines were generated by single cell sorting using fluorescent-assisted cell sorting (FACS).

**Genome-wide CRISPR/Cas9 screen**

The TKOv3 CRISPR/Cas9 library was a gift from Jason Moffat (Addgene #90294). The sgRNA library and 2nd generation lentiviral packaging plasmids were transfected into HEK293T cells. Virus titre was determined for optimising transduction at MOI of 0.3 in HEK TF SUN2 Site 2\textsuperscript{D} cells in T175 tissue culture flask format. For the screen, at least 125x10\textsuperscript{6} HEK TF SUN2 Site 2\textsuperscript{D} cells were infected in T175 flasks to achieve a 250-fold coverage of the library after selection at day 1. On day 2, fresh media was replaced. On day 3, cells were selected with 2ug/mL puromycin (GIBCO) for 72 h. Cells were seeded into two technical replicates with at least
25x10^6 total cells per replicate in T175 flasks. 24 hours prior to Day 7-sorting, 100 ng/mL Doxycycline (Sigma-Aldrich) was added to the cell medium to induce the expression of the transgene SUN2 Site 2D transgene. On the day of sorting, cells were trypsinized and collected into a single flask. From this, 25x10^6 reference cells were collected, pelleted and frozen. In addition, two replicates of 25x10^6 cells were further seeded for Day 13 sorting. The rest of the cells were subjected to FACS and 1% of the brightest GFP cells was collected using a BD FACSaria3 sorter. Transgene induction, reference population collection, sorting and seeding for Day 17 were repeated as described above. Genomic DNA was extracted from each cell population using a QIAGEN BloodMaxi kit (for reference samples) or BloodMini kit (for sorted samples) according to the manufacturer's protocol. sgRNAs were PCR amplified from the entire isolated genomic DNA using NEBNext Ultra II Q5 Master Mix (NEB) and the primers v2.1-F1 and v2.1-R1, according to the TKOv3 protocol. PCR reactions were pooled again, after which a second PCR was performed to attach indices and sequencing adapters using the primers i5 and i7. The PCR reaction was loaded onto a 2% agarose gel, the 200bp band excised and purified using a GeneJet PCR Purification kit (Thermo Fischer Scientific). Concentration and quality of the purified bands were assessed by using TapeStation and pooled library at 10nM in 30ul were prepared. The library was analyzed by deep sequencing on an Illumina NextSeq500 at the Oxford Genomics Centre. Gene rankings were generated using the MAGeCK algorithm.

**Flow cytometry**

Cells were trypsinized and resuspended in FACS buffer (2% FBS, 1 mM EDTA in PBS). Cells were analyzed using BD LSRFortessa X-20 flow cytometer on plate reader standard mode.
Legends to the Supplemental Figures

Figure 1- figure supplement 1: Evolutionary conservation analysis of SUN2 protein sequence.
(A) Schematic of human SUN2 domain organization.
(B) Multiple sequence analysis of SUN2 Site 1 (Top) and Site 2 (Bottom) among primates (chimpanzee, rhesus and green monkeys), other mammals (mouse and dog) and marsupial (Oppossum), amphibian (frog) and bird (chicken). Regions corresponding to human SUN2 Site 1 and Site 2 are indicated by boxes.
(C) Multiple sequence analysis of nucleoplasmic sequences of human SUN1 and SUN2 proteins. Regions corresponding to human SUN2 Site 1 and Site 2 are indicated by boxes.

Figure 2- figure supplement 1: Depletion of β-TrCP1/2 stabilises SUN2 Site 2D
(A) Western blot analysis of SUN2 Site 2D steady state levels in HEK TF cells transfected with an empty vector (EV), sgRNAs against βTrCP1 (three independent gRNAs), βTrCP2 (two independent gRNAs) or both. sgRNAs targeting unrelated F-Box genes FBXO2 and FBXO6 were included as specificity control. Cell extracts were analyzed by SDS–PAGE and immunoblotting. Transgenic SUN2 was detected with anti-HA antibody. Depletion of βTrCP1 is detected with anti-βTrCP1 antibody. Tubulin was used as a loading control and detected with an anti-Tubulin antibody.
(B) Flow cytometry analysis of SUN2 Site 2D steady state levels in HEK TF cells treated as in (A). Cells transfected with EV are shown in gray and specific gRNA for specific genes in blue.
Figure 4- figure supplement 1: Analysis of genome-wide screening

(A) Significance score of the genes analyzed in the genome-wide CRISPR/Cas9 genetic screen calculated by the MAGeCK algorithm as done in Fig 5 for additional timepoints at Day 13 (top) and Day 17 (Bottom) post-TKO lentivirus transduction. Core and regulatory subunits of the SCF ubiquitin ligase are indicated in blue.

(B) Flow cytometry analysis of validation of several hits from the screen with three independent gRNAs in HEK TF SUN2 Site 2^D cells. Cells transfected with EV are shown in gray and with specific gRNA for specific genes in blue (a regulatory component of SCF ubiquitin ligase) and green (p97 and p97 cofactors).

(C) Flow cytometry (Left) and western blot (Right) analysis of SUN2 WT steady state levels in parental HEK TF cells and CTDNEP1 KO cells, which display lower SUN2 levels (see Figure 5), transfected with an empty vector (EV) or sgRNAs targeting CK2b. On the left, cell extracts were analyzed by SDS–PAGE and SUN2 was detected with anti-HA antibody. Depletion of CK2b was confirmed with an anti-CK2b antibody. Tubulin was used as a loading control and detected with an anti-Tubulin antibody.

Figure 5- figure supplement 1: Characterization of the CTDNEP1 role in SUN2 degradation

(A) Immunofluorescence in parental and CTDNEP1 KO Hela cells. Expression of SUN2 Site 2^D was induced with Dox for 24h. Endogenous LBR was detected with anti-LBR antibody. DNA was labelled with DAPI. Scale bar; 10µm.

(B) Analysis of transgenic SUN2 stability after inhibition of protein synthesis by cycloheximide (CHX) in HEK TF SUN2 WT-expressing parental and CTDNEP1 KO cells. Cell extracts were analyzed by SDS–PAGE and
immunoblotting. Transgenic SUN2 is detected with anti-HA. Tubulin was used as a loading control and detected with an anti-Tubulin antibody. Three independent experiments are quantified on the graph below.

(C) The degradation of endogenous SUN2 in CTDNEP1 KO HEK TF cells is curbed by inhibitors to SCFβTrCP neddylation (MLN-4924; 1 µM), p97 ATPase (CB-5083; 2.5 µM, CB) or the proteasome (Bortezomib; 1 µM, Btz). Cell extracts were analyzed by SDS–PAGE and immunoblotting. Endogenous SUN2 was detected with anti SUN2 antibody. Tubulin was used as a loading control and detected with an anti-Tubulin antibody.

(D) Western blot analysis of endogenous SUN2 steady state levels in HEK TF cells transfected with an empty vector (EV), sgRNAs against βTrCP1 (three independent gRNAs), βTrCP2 (two independent gRNAs) or both. sgRNAs targeting unrelated F-Box genes FBXO2 and FBXO6 were included as specificity control. Cell extracts were analyzed by SDS–PAGE and immunoblotting. Endogenous SUN2 was detected with an anti-SUN2 antibody. Depletion of βTrCP1 is detected with anti-βTrCP1 antibody. Tubulin was used as a loading control and detected with an anti-Tubulin antibody.

(E) Flow cytometry analysis of transgenic SUN2 steady state levels in HEK TF SUN2 WT-expressing parental and CTDNEP1 KO cells transfected with an empty vector (EV), a vector encoding FLAG-tagged WT or phosphatase dead (PD) CTDNEP1.

(F) Western blotting analysis of Parental and CTDNEP1KO HEK TF cells incubated 6h with Casein Kinase 2 inhibitor (TBCA; 100 µM) or DMSO as a control. SUN2 was detected with anti-SUN2 antibody. GAPDH was used as a loading control and detected with an anti-GAPDH antibody.
(G) Flow cytometry analysis of transgenic SUN2 steady state levels in HEK TF SUN2 Site 2A-expressing cells transfected with an empty vector (EV) or sgRNAs targeting CTDNEP1.

**Figure 6- figure supplement 1: Analysis of the accumulation of non-degradable SUN2 in mitotic progression**

(A) Flow cytometry analysis of transgenic SUN2 steady state levels in HEK TF expressing SUN2 WT (Left) or Site 2A (Right) either without (0 h) or upon shutoff of the transgenes with washing out Dox for 24, 48 or 72 h after an initial 24 h DOX induction. Negative control indicates cells that were not treated with Dox.

(B) Timelapse microscopy analysis of HeLa cells expressing SUN2 WT or Site 2A, filmed progressing through mitosis. SiR-Hoechst (DNA) is shown in white and transgenic SUN2 in green. All timings are from the final metaphase timepoint. Brightness and contrast is equal in all panels, apart from the bottom SUN2 WT (brightened) panel where the SUN2 WT signal has been increased for better visualisation of its localisation.

(C) Quantification of timing between various events during mitosis in HeLa cells expressing SUN2 WT (Pink) or Site 2A (Light green) from timelapse microscopy experiment in (B). The last panel indicates quantification of percentage of multipolar cells observed in (B). At least 45 cells were quantified per experiment and three independent experiments were performed. Unpaired t-test statistical analysis was performed (*p<0.05, **p<0.01, ***p<0.001).

(D) Quantification of percentage of HeLa cells expressing SUN2 WT (Pink) or Site 2A (Light green) that displayed delayed chromosome release from nuclear membrane (Top left) and cells with lagging chromosomes (Top right). In the bottom panel, quantification of timing between NEBD and loss
of SUN2 from the membrane in HeLa cells expressing SUN2 WT (Pink) or Site 2A (Light green) from timelapse microscopy experiment in (B). At least 45 cells were quantified per experiment and three independent experiments were performed. Unpaired t-test statistical analysis was performed (*p<0.05, **p<0.01, ***p<0.001).
References


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Margottin Florence, Bour SP, Durand H, Selig L, Benichou S, Richard V,


Rogakou EP, Pilch DR, Orr AH, Ivanova VS, Bonner WM. 1998. DNA double-stranded breaks induce histone H2AX phosphorylation on serine 139. *J*


Figure 1: SUN2 stability depends on a non-canonical recognition motif for the SCFβTrCP ubiquitin ligase

A. Site 1

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B. Site 2

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C. Relative cell counts

D. Relative SUN2 protein levels

E. SUN2 WT, SUN2 Site 1, SUN2 Site 2, SUN2 Site 1,2

CHX chase (min) 0 30 60 120 240
Figure 2: INM degradation of SUN2 by an ERAD-like process

A. 

B. 

C. 

D. 

SUN2 WT

SUN2 Site2A

SUN2 Site2D

IB: HA

IB: GAPDH

IB: LC3B

IB: GAPDH
Figure 3: Site 2 is critical for SUN2 recognition and ubiquitination by SCFβTrCP ubiquitin ligase

A.

B.

10% Input

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<td>CB5083:</td>
<td></td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
</tr>
</tbody>
</table>

IP: HA

<table>
<thead>
<tr>
<th></th>
<th>WT</th>
<th>Site1^A</th>
<th>Site1^D</th>
<th>Site2^A</th>
<th>Site2^D</th>
</tr>
</thead>
<tbody>
<tr>
<td>CBS083:</td>
<td></td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>CB5083:</td>
<td></td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
</tr>
</tbody>
</table>

IB: HA

IB: β-TrCP1

Neddylated-Cullin1

IB: Cullin1

IB: p97

IP: HA

IB: Ubiquitin
Figure 4: Genome-wide screening identifies Casein kinase 2 as a positive regulator of SUN2 degradation

A. Lentiviral transduction
TK0v3 genome-wide
CRISPR/Cas9 library
Puromycin selection
Induce SUN2 Site2D expression with Doxycyline

HEK TF
SUN2(Site2D)-sfGFP-3xHA

B. Day 9 Post-transduction

-log10 (positive score)

Genes

C. SUN2 WT
SUN2 Site2A
SUN2 Site2D
SUN2 Site1D,2D

Relative cell counts

D. SUN2 WT
SUN2 Site2A
SUN2 Site2D
SUN2 Site1D,2D

DMSO:
CB-5083:
TBCA:

IB: HA

E. 10% Input
IP:HA

1% Top GFP+

Genomic DNA isolation
Barcoding and mix
Count sgRNA by deep sequencing and calculate gene score

- SCF ubiquitin ligase core subunits
- NEDD8 conjugation cascade
- Ubiquitin conjugation cascade
- p97 and cofactors
- Kinase

- DMSO:
- TBCA:
- CB-5083:
Figure 5: The phosphatase CTDNEP1 is a negative regulator of SUN2 degradation

A.

B.

C.

D.

E.

F.

G.
Figure 6: Aberrant nuclear architecture and function by expression of non-degradable SUN2

A. 

24hrs SUN2 ON  

24hrs SUN2 ON  24hrs SUN2 OFF

DAPI  GFP

Parental  

Sun2 WT

Sun2 Site2a

B. 

24hrs SUN2 ON  24hrs SUN2 ON  24hrs SUN2 OFF

Percentages of nuclear phenotype, %

Parental  

SUN2 WT

SUN2 Site2a

C. 

SUN2 Site2a

DAPI  GFP  Lap2b  LBR

DAPI  GFP  EMD

DAPI  GFP  SUN1

D. 

Parental  SUN2 Site2a

No IR

Time post-IR

1 hour

24 hours

E. 

Parental  SUN2 Site2a

Number of γH2AX foci / nucleus

Time post-IR (Hours)

No IR  1  24

**** (p<0.0001)
Figure 7: A kinase-phosphatase balance regulates ER-associated degradation of SUN2 from the INM
Figure 1-figure Supplement 1: Evolutionary conservation analysis of SUN2 protein sequence

A. Schematic of human SUN2 domain organization.

B. Multiple sequence analysis of SUN2 Site 1 and Site 2 (CLUSTAL O (1.2.4))

Site 1

<table>
<thead>
<tr>
<th>Species</th>
<th>Sequence</th>
<th>Length</th>
</tr>
</thead>
<tbody>
<tr>
<td>HUMAN</td>
<td>GDGS-SSSGG-SSVAGSQTSLFK</td>
<td>30</td>
</tr>
<tr>
<td>CHIMPANZEE</td>
<td>GDGS-SSSGG-SSVAGSQTSLFK</td>
<td>33</td>
</tr>
<tr>
<td>RHESUS MONKEY</td>
<td>GDGS-SSSGG-SSVAGSQTSLFK</td>
<td>38</td>
</tr>
<tr>
<td>GREEN MONKEY</td>
<td>GDGS-SSSGG-SSVAGSQTSLFK</td>
<td>37</td>
</tr>
<tr>
<td>MOUSE</td>
<td>DNGSSGSSGA-SSVAGSGGTQVFK</td>
<td>36</td>
</tr>
<tr>
<td>DOG</td>
<td>GDGS-SSSGG-SSVMQGQTFK</td>
<td>36</td>
</tr>
<tr>
<td>PIG</td>
<td>GDGS-SSSGG-SSVAGSQTSLFK</td>
<td>116</td>
</tr>
<tr>
<td>WHALE</td>
<td>GDGS-SSSGG-SSVAGSQTSLFK</td>
<td>36</td>
</tr>
<tr>
<td>OPPOSUM</td>
<td>GDGS-SSSGG-SSVLMIQHVPKF</td>
<td>36</td>
</tr>
<tr>
<td>FROG</td>
<td>EDDKRS------SSMESHSLYK</td>
<td>37</td>
</tr>
<tr>
<td>CHICKEN</td>
<td>EDEDG--------SSSLGGPQLPFK</td>
<td>34</td>
</tr>
</tbody>
</table>

Site 2

<table>
<thead>
<tr>
<th>Species</th>
<th>Sequence</th>
<th>Length</th>
</tr>
</thead>
<tbody>
<tr>
<td>HUMAN</td>
<td>RKATredniGSSSSGYSSEDDYV</td>
<td>131</td>
</tr>
<tr>
<td>CHIMPANZEE</td>
<td>RKATredniGSSSSGYSSEDDYV</td>
<td>136</td>
</tr>
<tr>
<td>RHESUS MONKEY</td>
<td>RKATredniGSSSSGYSSEDDYV</td>
<td>152</td>
</tr>
<tr>
<td>GREEN MONKEY</td>
<td>RKATrediGSSSSGYSSEDDYV</td>
<td>152</td>
</tr>
<tr>
<td>MOUSE</td>
<td>SKASEGFGSSGSSGYSSEDDLA</td>
<td>141</td>
</tr>
<tr>
<td>DOG</td>
<td>DKASEGFGSSGSSGYSSEDDFA</td>
<td>141</td>
</tr>
<tr>
<td>PIG</td>
<td>GKADEGFGSSGSSGYSSEDDYV</td>
<td>225</td>
</tr>
<tr>
<td>WHALE</td>
<td>NKLSEGFGSSGSSGYSSEDDYV</td>
<td>145</td>
</tr>
<tr>
<td>OPPOSUM</td>
<td>DKISYDTYGSSSSGYSSEDDYS</td>
<td>150</td>
</tr>
<tr>
<td>FROG</td>
<td>TQPSYGTSSSSGYSSEEDNT</td>
<td>154</td>
</tr>
<tr>
<td>CHICKEN</td>
<td>-SRTYDYaGSSSSGYSSEDDYa</td>
<td>152</td>
</tr>
</tbody>
</table>

C. Multiple sequence analysis of comparing SUN1 and SUN2 (CLUSTAL O (1.2.4)).

```
Figure 2 - figure Supplement 1: Genetic depletion of B-TrCP1/2 stabilises SUN2 Site 2<sup>0</sup> transgene

A. SUN2 Site<sup>20</sup>

B. gBTRCP1_1, gBTRCP1_2, gBTRCP1_3, gBTRCP2_1, gBTRCP2_2, gBTRCP1+2, gFBXO2_1, gFBXO2_2, gFBXO6_1, gFBXO6_2.

Relative cell counts

SUN2: sfGFP-3xHA
Figure 4 -figure Supplement 1: Analysis of genome-wide screening

A. Timepoint 2 (Day 13)

B. Timepoint 3 (Day 17)

C. Parental CTDNEP1KO

Parental EV 

CTDNEP1 KO EV 

IB: HA

IB: CK2b

IB: Tubulin
Figure 5 - figure Supplement 1: Characterization of the CTDNEP1 role in SUN2 degradation

A. 

![Immunostaining images](image)

B. 

<table>
<thead>
<tr>
<th>CHX chase (min)</th>
<th>0</th>
<th>240</th>
</tr>
</thead>
<tbody>
<tr>
<td>DMSO:</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>MLN-4924:</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>CB5083:</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Bortezomib:</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

IB: HA (short exp)
IB: HA (long exp)
IB: Tubulin

C. 

CTDNEP1 KO

<table>
<thead>
<tr>
<th>CHX chase (min)</th>
<th>0</th>
<th>240</th>
</tr>
</thead>
<tbody>
<tr>
<td>DMSO:</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>MLN-4924:</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>CB5083:</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Bortezomib:</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

IB: SUN2
IB: Tubulin

D. 

CTDNEP1KO

E. 

SUN2 WT

F. 

CTDNEP1KO

G. 

SUN2 Site2^a
**Figure 6**: Analysis of the accumulation of non-degradable SUN2 in mitotic progression

**A.**

![ WT Site2\(^{a}\) 

<table>
<thead>
<tr>
<th>SUN2-sfGFP-3xHA</th>
<th>Negative control</th>
<th>SUN2 transgene shut-off</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1" alt="WT" /></td>
<td><img src="image2" alt="Site2(^{a})" /></td>
<td>0 hour</td>
</tr>
</tbody>
</table>

**B.**

<table>
<thead>
<tr>
<th>Metaphase</th>
<th>Anaphase</th>
<th>Telophase</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 min</td>
<td>5 min</td>
<td>10 min</td>
</tr>
<tr>
<td>SIR-Hoechst SUN2 WT</td>
<td>SIR-Hoechst SUN2 Site2(^{a})</td>
<td>2X brightened SUN2 WT</td>
</tr>
</tbody>
</table>

**C.**

<table>
<thead>
<tr>
<th>Prophase - NEBD (min)</th>
<th>NEBD - Alignment (min)</th>
<th>Alignment - Anaphase (min)</th>
<th>Multipolar Cells (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image3" alt="Prophase" /></td>
<td><img src="image4" alt="NEBD - Alignment" /></td>
<td><img src="image5" alt="Alignment - Anaphase" /></td>
<td><img src="image6" alt="Multipolar Cells" /></td>
</tr>
</tbody>
</table>

| ![Prophase](image7) | ![NEBD - Alignment](image8) | ![Alignment - Anaphase](image9) | ![Multipolar Cells](image10) |

**D.**

- **Cells with delayed chromosome release from nuclear membrane (%):**
  - n.s. (p = 0.8640)
  - 50
  - 40
  - 30
  - 20
  - 10
  - 0

- **Cell with lagging chromosomes (%):**
  - *** (p = 8.64 \(10^{-5}\))
  - 80
  - 60
  - 40
  - 20
  - 0

- **Multipolar Cells (%):**
  - n.s. (p = 1.000)
  - 20
  - 15
  - 10
  - 5
  - 0

- **NEBD - SUN2 Membrane Loss (min):**
  - * (p = 0.0114)

- **SUN2 WT**
- **SUN2 Site2\(^{a}\)**