

γH2A binds Brc1 to maintain genome integrity during S-phase

Jessica S Williams^{1,5,6}, R Scott Williams^{1,2,5,6}, Claire L Dovey¹, Grant Guenther^{1,2}, John A Tainer^{1,2,4,*} and Paul Russell^{1,3,*}

¹Department of Molecular Biology, The Scripps Research Institute, La Jolla, CA, USA, ²Skaggs Institute for Chemical Biology, The Scripps Research Institute, La Jolla, CA, USA, ³Department of Cell Biology, The Scripps Research Institute, La Jolla, CA, USA and ⁴Life Sciences Division, Department of Molecular Biology, Lawrence Berkeley National Laboratory, Berkeley, CA, USA

ATM^{Tel1} and ATR^{Rad3} checkpoint kinases phosphorylate the C-terminus of histone H2AX (H2A in yeasts) in chromatin flanking DNA damage, establishing a recruitment platform for checkpoint and repair proteins. Phospho-H2A/X (yH2A/X)-binding proteins at double-strand breaks (DSBs) have been characterized, but those required for replication stress responses are unknown. Here, we present genetic, biochemical, small angle X-ray scattering (SAXS), and X-ray structural studies of the Schizosaccharomyces pombe Brc1, a 6-BRCT-domain protein that is structurally related to Saccharomyces cerevisiae Rtt107 and mammalian PTIP. Brc1 binds yH2A to form spontaneous and DNA damage-induced nuclear foci. Spontaneous Brc1 foci colocalize with ribosomal DNA repeats, a region prone to fork pausing and genomic instability, whereas DNA damageinduced Brc1 foci colocalize with DSB response factors. yH2A binding is critical for Brc1 function. The 1.45 Å resolution crystal structure of Brc1-yH2A complex shows how variable BRCT insertion loops sculpt tandem-BRCT phosphoprotein-binding pockets to facilitate unique phosphoprotein-interaction specificities, and unveils an acidic DNA-mimicking Brc1 surface. From these results, Brc1 docking to yH2A emerges as a critical chromatin-specific response to replication-associated DNA damage.

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⁵Present address: Department of Health and Human Services, Laboratory of Structural Biology, National Institute of Environmental Health Sciences, National Institutes of Health, NC 27709, USA ⁶These authors contributed equally to this work

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Introduction

DNA damage responses for maintaining genome integrity require that DNA repair and checkpoint proteins work in concert with factors that bind or modify chromatin at DNA lesions (Stucki and Jackson, 2006; Harper and Elledge, 2007). Central to this integration of chromatin metabolism and DNA repair, the checkpoint kinases ATM^{Tel1} and ATR^{Rad3/Mec1} phosphorylate the carboxyl terminus of histone H2AX (or H2A in yeasts) in multi-kilobase regions of chromatin surrounding DNA double-strand breaks (DSBs), creating a phospho-H2A/X (yH2A/X) recruitment platform for checkpoint and DSB repair factors (Rogakou et al, 1998; Celeste et al, 2003; Stucki and Jackson, 2006; Williams et al, 2007). Although functions of γ H2A/X at DSBs have been extensively investigated, its functions in cellular responses to spontaneous and DNA replication-associated DNA damage have received comparatively little attention and remain a mystery. As cells are constantly exposed to endogenous (i.e. metabolic) sources of DNA damage that stall or collapse replication forks, and the most widely used anti-cancer drugs cause S-phase-specific DNA damage, such unknown effectors of ATM/ATR-dependent yH2A/X signalling during replication are arguably the most crucial.

Evidence for such unknown S-phase yH2A effectors comes from genetic and microscopic studies of the Schizosaccharomyces pombe checkpoint mediator protein Crb2. In response to DSB producing ionizing radiation (IR), the Crb2 tandem-BRCT (BRCA1 carboxy terminus) domains directly bind γ H2A in a phosphorylation-dependent manner (Nakamura et al, 2004; Du et al, 2006; Kilkenny et al, 2008). γH2A is required for Crb2 IR-induced focus formation and for amplification and maintenance of DNA damage checkpoint signals. However, H2A phosphorylation is also critical for fission yeast responses to hydroxyurea (HU), which stalls replication forks without creating DNA double breaks, and for survival of exposure to camptothecin (CPT) (Redon et al, 2003; Nakamura et al, 2004), a prototypical anti-cancer drug that collapses replication forks by poisoning topoisomerase I (Pommier, 2006). Yet, ablation of Crb2 and yH2A uncovers non-epistatic genetic interactions (Du et al, 2006), suggesting the existence of additional important yH2A-interacting factors.

Candidate S-phase γ H2A effectors include *S. pombe* Brc1 and budding yeast Rtt107 (Esc4), which are members of a poorly understood family of 6-BRCT-domain proteins that appear to be conserved in fungi. Brc1 and Rtt107 are required for survival of CPT treatment, but are dispensable for survival of IR (Verkade *et al*, 1999; Rouse, 2004; Sheedy *et al*, 2005; Roberts *et al*, 2006; Zappulla *et al*, 2006). Rtt107 is required for stabilization of stalled replication forks and reestablishment of collapsed forks (Rouse, 2004; Chin *et al*, 2006; Roberts *et al*, 2008), a function, which may involve localization to sites of DNA replication-associated damage through uncharacterized protein–protein interactions. However, the

^{*}Corresponding authors. JA Tainer or P Russell, Department of Molecular Biology, The Scripps Research Institute, 10550 North Torrey Pines Road, La Jolla, CA 92037, USA. Tel.: +1 858 784 8119; Fax: +1 858 784 2277; E-mail: jat@scripps.edu or

Tel.: +1 858 784 8273; Fax: +1 858 784 8273;

E-mail: prussell@scripps.edu

mechanism of DNA damage recruitment of these 6-BRCT genome maintenance factors and how they enable specific repair of replication-associated damage remains unknown. Brc1 and Rtt107 are structurally related to mammalian PTIP proteins that have 6-BRCT domains and participate in DNA damage responses (Muñoz and Rouse, 2009). Although it is unclear as to what extent these proteins are functionally related, a detailed structure defining the functional interactions for any member of this group would provide a critical framework for improved understanding of their structural and functional relationships.

Here, we provide integrated microscopic, genetic, biochemical and structural results showing that the C-terminal pair of Brc1 BRCT domains (BRCT₅–BRCT₆) dock to γ H2A and act in S-phase damage responses. Abrogating Brc1– γ H2A interactions confers sensitivity to replication fork-damaging agents. A high-resolution X-ray structure of the BRCT₅– BRCT₆: γ H2A phosphoprotein complex reveals stereo-specific recognition of pSer phosphates by BRCT repeats, and a probable general mechanism for generating alternate phosphoprotein interaction specificities in the large BRCT-containing superfamily of replication and DNA repair proteins. Together, these results define Brc1 as a major effector of Tel1/Rad3 and γ H2A signalling in mechanisms that protect genome integrity from replication-associated DNA damage.

Results

Brc1 protects the genome from endogenous DNA damage

Brc1 is required for survival of genotoxins that interfere with DNA replication (Verkade *et al*, 1999; Sheedy *et al*, 2005; Dovey and Russell, 2007), but we suspected it might also protect genome integrity during an unperturbed cell cycle. We tested this by monitoring nuclear foci of Rad22^{Rad52}, a critical HR protein that mediates assembly of Rad51 onto ssDNA (New *et al*, 1998). Live-cell analysis revealed that spontaneous Rad22-YFP foci were increased ~2-fold in *brc1*\Delta cells (Figure 1A). Interestingly, a similar effect is seen in *hta-AQ* cells that lack the C-terminal SQ phosphorylation motif required to form γ H2A (Nakamura *et al*, 2004).

To explore whether $brc1\Delta$ cells require HR repair for survival, we tested for genetic interactions between $brc1\Delta$ and $rad22\Delta$. Tetrad analysis uncovered a severe synergistic growth defect in $brc1\Delta$ $rad22\Delta$ cells (Figure 1B). Double mutants were either inviable or formed microcolonies containing highly elongated cells indicative of a prolonged DNA damage checkpoint arrest (Figure 1B). As even one broken replication fork is lethal in $rad22\Delta$ cells (Roseaulin *et al*, 2008), $brc1\Delta$ cells are likely suffering increased rates of replication fork collapse or rearrangement that are repaired by Rad22-dependent HR.

Brc1 forms γH2A-dependent nuclear foci

Brc1 has 4 BRCT domains (BRCT₁-BRCT₂-BRCT₃-BRCT₄) near its N-terminus followed by an acidic linker region and a tandem pair of BRCTs (BRCT₅-BRCT₆) at its extreme C-terminus (Figure 1C). We postulated that as a BRCT domain-containing protein that is required for genome stability, Brc1 might localize into nuclear foci. Expression of functional, C-terminally tagged GFP-Brc1 from its genomic locus was insufficient for microscopic detection. However, when GFP-Brc1 was expressed from the moderate strength *nmt41* promoter, it formed distinct nuclear foci in \sim 15–25% of cells in an asynchronous population (Figure 1D).

As $brc1\Delta$ and hta-AQ mutations confer sensitivity to an overlapping spectrum of S-phase DNA-damaging agents (Verkade *et al*, 1999; Nakamura *et al*, 2004; Sheedy *et al*, 2005), we hypothesized that Brc1 foci formation might require γ H2A. Indeed, spontaneous nuclear GFP-Brc1 foci were absent in *hta*-AQ cells (Figure 1E).

Rad3 and Tell create γ H2A (Nakamura *et al*, 2004). Rad3-Rad26^{ATRIP/Ddc2} complex associates with DNA lesions by binding the single-stranded (ss) DNA-binding protein replication protein A (Zou and Elledge, 2003), whereas Tell associates with DSBs by binding Nbs1 (Falck *et al*, 2005; You *et al*, 2005). To investigate which types of interactions create spontaneous Brc1 foci, we expressed GFP-Brc1 in *rad3* Δ or *tel1* Δ strains. GFP-Brc1 foci were detected in both mutant backgrounds (Figure 1F), although they were reduced in number (Figure 1G), indicating that Rad3 and Tell both regulate Brc1 localization. As predicted, GFP-Brc1 foci were abolished in a *rad3* Δ *tel1* Δ strain (Figure 1F).

Brc1 BRCT₅-BRCT₆ domains bind γH2A

As tandem-BRCT domains in MDC1 and Crb2 bind yH2A/X (Stewart et al, 2003; Stucki et al, 2005; Lou et al, 2006; Kilkenny et al, 2008), and Brc1 foci formation is independent of Crb2 (Supplementary Figure S1), we hypothesized that a pair of BRCT domains in Brc1 may interact directly with yH2A. Sequence analysis of the six Brc1 BRCT domains revealed that only BRCT₅ has phosphoserine-binding consensus motifs (Williams et al, 2004), referred to here as C1: (S/T)G, and C2: (T/S)XK (X is any residue). To test whether a Brc1:yH2A interaction is mediated via direct association, we analysed binding of recombinant Brc1 BRCT₅-BRCT₆ to N-terminal fluorescein-conjugated yH2A peptides by analytical gel filtration and fluorescence polarization (FP) (Figure 2A and B). In isolation, an FITC-yH2A peptide migrates as an ~4 kDa peak in gel filtration, but coelutes at \sim 25 kDa in the presence of purified BRCT₅-BRCT₆, consistent with a direct Brc1-γH2A interaction. FP-binding analysis shows Brc1 BRCT₅-BRCT₆ binds the γ H2A tail with a 2.6- μ M affinity, similar to other BRCT-phosphoprotein interactions (Figure 2A and B; Manke et al, 2003; Stucki et al, 2005). A Brc1 C1 motif mutant (T672A) does not bind, and interactions with an unphosphorylated H2A peptide are not detected by either method, indicating the interaction is phosphorylation dependent (Figure 2A and B). The hMDC1 BRCT domains bind human yH2AX C-terminal tail (Stucki et al, 2005), but unlike the Brc1- γ H2A interaction, hMDC1 specifically recognizes the human 'SQEY-coo(-)' γ H2AX C-terminus, and will not bind the yeast 'SQEL-coo(-)' carboxy terminus (Stucki *et al*, 2005). Thus, critical differences in these γ H2A interactions are dictated by molecular recognition of the C-terminal H2A residue at the pSer +3 position.

Architecture of the Brc1–γH2A complex

To examine the structural basis for specificity differences between mammalian and yeast γ H2A–BRCT interactions, and provide general insights into the structural determinants of phosphoprotein interaction specificities in the BRCT-containing superfamily of proteins, we solved high-resolution X-ray structures for unliganded Brc1 BRCT₅–BRCT₆ and the

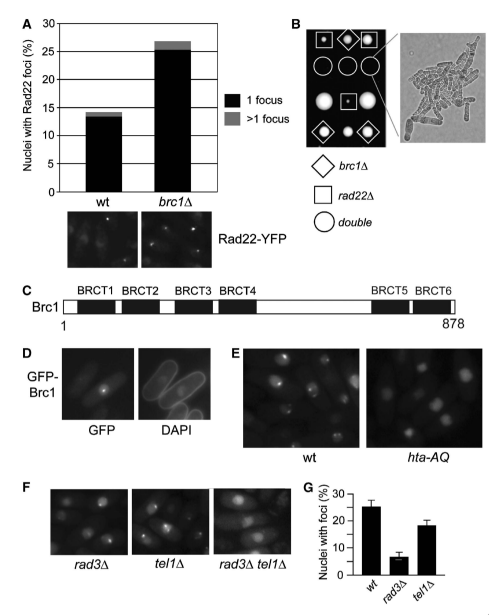


Figure 1 Brc1 is required for genome stability and forms γ H2A-dependent spontaneous nuclear foci. (A) Deletion of $brc1^+$ causes an increase in Rad22-YFP foci. Cells were grown to mid-log phase in YES medium and imaged. Foci numbers from >1000 cells for each strain were quantified. (B) Tetrad analysis of genetic crosses between $brc1\Delta$ and $rad22\Delta$ reveals a severe synergistic growth in the double mutants. Representative spores from three asci are shown. (C) Schematic of the Brc1 BRCT domain organization. BRCT₅ and BRCT₆ form a tandem pair containing a consensus phosphoprotein-binding motif. (D) Cells were grown to mid-log phase in selective medium and fixed for 10 min in 70% ethanol at room temperature. DAPI staining was performed to visualize the nuclei and cell morphology. (E) Brc1 foci formation is dependent on phosphorylation of γ H2A. Live cell microscopy of ectopically expressed GFP-Brc1 driven by the *nmt41* promoter in wild-type (wt) or *hta-AQ* cells. (F) Brc1 foci formation is abolished in the *rad3Atel1*\Delta mutant. (G) Brc1 foci formation is reduced in the *rad3*\Delta and *tel1*\Delta mutants. Live cell microscopy of GFP-Brc1 was examined in the indicated strain backgrounds.

BRCT₅-BRCT₆: γ H2A complex (Figure 2C, see Materials and methods). Three key observations from the Brc1: γ H2A structure extend our understanding of BRCT domain-phosphoprotein recognition and specificity. First, the 1.45 Å structure of the BRCT₅-BRCT₆: γ H2A complex provides an unprecedented high-resolution visualization of the BRCT-phosphoprotein interaction, revealing critical conserved determinants of phosphoserine recognition and binding specificity. Second, comparative analysis of the second primary BRCT peptide-binding determinant, the pSer +3 binding pocket, shows how variable sequences and structures at the BRCT-BRCT interface mold pSer +3 pockets to accommodate different

peptide target specificities. Third, structural conservation of an exposed acidic Brc1 and MDC1 surface outside of the γ H2A/X tail binding cleft suggests additional histone core particle interactions augment phosphorylated histone tail binding in histone-interacting BRCT proteins.

Each Brc1 C-terminal BRCT domain folds as an α - β sandwich with a central four-stranded parallel β -sheet that is flanked on one side by two helices (α 1 and α 3), and on the another by helix α 2. BRCT₅ and BRCT₆ also contain unique helical elements (α I1 and α II') inserted between strands β 2 and β 3. Packing of an α 2- α 1'- α 3' three-helix bundle forms a $40 \times 30 \times 70$ Å elongated dual repeat BRCT-BRCT interface

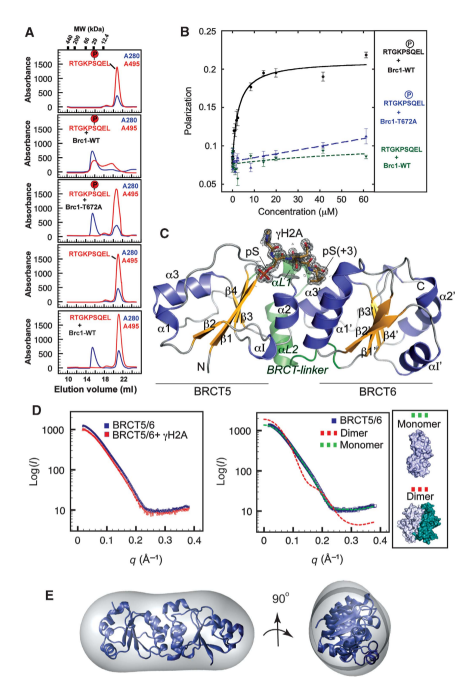


Figure 2 Brc1 BRCT₅–BRCT₆ domains interact directly with the phosphorylated γH2A C-terminal tail. (**A**) Analytical gel filtration binding analysis of Brc1–γH2A protein–peptide interactions. BRCT₅–BRCT₆–γH2A binding was monitored by following coelution of FITC-labelled γH2A peptides that absorb at 495 nm (red trace) with recombinant BRCT₅–BRCT₆ (blue trace). All peptides are FITC conjugated on the N-terminal amide and bear carboxylated C-termini. γH2A peptide coelution with wild-type BRCT₅–BRCT₆ is observed only for the phosphorylated γH2A peptide and is disrupted by pSer binding site mutation T672A. (**B**) FP-binding analysis of the Brc1–γH2A.1 interaction. Brc1 binds the γH2A tail with an apparent affinity of 2.6 µM, whereas the Brc1–T672A mutant or unphosphorylated peptide display no detectable binding. (**C**) X-ray crystal structure of the Brc1–γH2A complex. The γH2A peptide (gold with grey electron density) binds at the interface between Brc1 BRCT₅ and BRCT₆ domains (yellow β-sheets and blue α-helices). Sigma-A weighted 1.45 Å Fo–Fc density contoured at 3 σ was calculated before building the peptide chain. (**D**) Brc1 BRCT5/6 and the BRCT5/6–γH2A complex were analysed by SAXS (left). No significant change in Brc1 oligomeric state occurs on peptide binding. SAXS scattering for apo BRCT5/6 fits well to a monomeric Brc1 model (χ^2 = 4.9), but not to the dimeric NCS-related dimer model (χ^2 = 54.1) found in the apo-Brc1 crystal structure (see inset for NCS dimer structure). (**E**) Low resolution averaged *ab initio* GASBOR shape reconstruction for apo BRCT5/6 closely matches the monomeric structure.

resembling other family members (Williams *et al*, 2001; Glover *et al*, 2004). γ H2A binds in the groove between the BRCT domains. Conformational change of helix α 1, and repositioning of the BRCT₅ and BRCT₆ repeats relative to one another accompany peptide binding. α 1 conformational change seems appropriate to relay BRCT peptide-binding state to neighbouring regions in BRCT-containing proteins, or to other binding partners.

As Brc1 BRCT₅–BRCT₆ associates as a non-crystallographic dimer with an extensive interface (1590 \AA^2 buried solvent-

accessible surface) in the unliganded crystal form, but is monomeric in the peptide-bound structure, we hypothesized monomer-dimer transitions might occur with yH2A binding. To test this, we evaluated the solution oligomeric state of BRCT₅-BRCT₆ and the BRCT₅-BRCT₆/ γ H2A complex in solution with small angle X-ray scattering (SAXS), which can provide accurate shape and assembly information in solution (Putnam et al, 2007; Hura et al, 2009). X-ray solution scattering obtained from purified BRCT₅-BRCT₆ in either the presence or absence of peptide reveals a monomeric assembly state in solution and is not consistent with dimeric models (Figure 2D). Also, low-resolution ab initio solution structures for apo- and peptide-liganded states closely match the elongated $40 \times 40 \times 70$ Å architectures of monomeric BRCT₅-BRCT₆ (Figure 2E). Gel filtration also shows both unbound and peptide-bound BRCT₅-BRCT₆ migrate with an apparent molecular weight of ~ 25 kDa, consistent with the monomeric form and a 1:1 protein:peptide-binding stoichiometry (Figure 2A). Together, SAXS and gel filtration analyses show that Brc1 BRCT5-BRCT6 is monomeric in solution and interacts with the yH2A.1 tail with 1:1 heterodimer stoichiometry.

Brc1–γH2A tail recognition

Three surface pockets embrace the pSer, pSer +2 Glu and pSer +3 Leu residues of the γ H2A tail (Figure 3A–E) and a 604 Å^2 solvent-accessible surface is buried at the Brc1- γ H2A interface. Residues N-terminal to yH2A pSer129 (Lys127 and Pro128) do not make direct protein contacts. γH2A binding is likely a conserved function as residues lining the walls of the interaction pockets are conserved in fungal Brc1 orthologs (Figure 3F). Interestingly, these residues are also conserved in mammalian PTIP, which requires yH2AX for the formation of IR-induced foci, although this is thought to involve an indirect mechanism requiring MDC1 and RNF8 (Stucki et al, 2005; Gong et al, 2009). The binding pockets exhibit precise shape and electrostatic complementarity with the bound peptide, and seem well suited for yH2A binding (Figure 3A). Comparisons of our high-resolution X-ray structures of the yH2A-free and yH2A-bound Brc1 show how flexure and rearrangements at the BRCT5-BRCT6 interface accommodate yH2A binding (Figure 3E). Strikingly, the partially formed pSer +3 (Leu) interaction pocket in the apo structure changes to mold around yH2A Leu132, revealing induced-fit yH2A binding to a structurally plastic Brc1 pocket (Figure 3E).

Thr672 (motif C1) and Lys710 (motif C2) (Figure 3B and D) from BRCT₅ and an associated network of protein-solventphosphate interactions mediate the yH2A pSer129 interaction. An analysis of phosphoserine oxygen-coordinating atoms indicates the phosphate moiety is likely bound as a polarized anion with two negatively charged oxygen atoms tetrahedrally coordinated by protein and solvent, and a third neutral, doubly bonded oxygen with linear coordination. Consistent with oxygen coordination, refinement of the complex to 1.45 Å (Figure 3C) with relaxed geometric restraints on the phosphoserine yields bond lengths appropriate for the two equidistant P–O bonds (1.55 Å), and a third shorter P = O(1.48 Å) bond (Figure 3D). We therefore propose that stereospecificity of the pSer interaction is a critical conserved determinant of BRCT-peptide binding, which maximizes coulombic attraction between the charged phosphate oxygen and the conserved C2 motif lysine (Brc1 K710). Selection for a polarized phosphoserine (versus resonance binding) probably contributes to moderate (high nM to low mM) binding affinities for four-residue BRCT molecular recognition sequences, despite the modest size of the BRCT proteinphosphoprotein interfaces. This phosphoserine interaction geometry may also ensure specific BRCT-phosphoprotein recognition of phosphoserine-phosphorylated targets over phospho-mimicking sequences containing Asp or Glu.

γ H2A-pSer and pSer + 3 contacts are critical for Brc1 foci formation and survival of S-phase damage

On the basis of our Brc1– γ H2A structure, we used mutational analysis to directly assess the functions for γ H2A-pSer, pSer +2 and pSer +3 contact residues in mediating spontaneous Brc1 nuclear focus formation. A salt bridge between Brc1 Arg704 and γ H2A Glu131 mediates the pSer +2 interaction. An R704E mutant localized to nuclear foci, albeit at a reduced frequency compared with WT, suggesting that the Brc1– γ H2A pSer +2 interaction is important, but not essential for γ H2A binding in cells (Figure 3G). In contrast, despite their nuclear localization, Brc1 foci formation was ablated for the T672A (pSer binding), K710M (pSer binding) and R707E (pSer +3 carboxylate binding) mutants. Thus, structural and *in vivo* observations suggest that the critical determinants of Brc1– γ H2A binding are phosphoserine binding and recognition of the pSer +3 position.

To assess the significance of the Brc1 BRCT₅–BRCT₆– γ H2A interaction and Brc1 foci formation, we introduced the T672A and K710M mutations into the genomic *brc1*⁺ locus and challenged these mutants with chronic exposure to S-phase genotoxins. Both mutants were sensitive to CPT and the DNA alkylating agent methyl methanesulphonate (MMS) (Figure 3H), which also interferes with DNA replication, indicating that γ H2A binding and foci formation are critical for Brc1 function. Interestingly, comparison to a *brc1* Δ mutant revealed that T672A and K710M are hypomorphic mutations (Figure 3H), indicating that Brc1 may retain submicroscopic interactions with DNA lesions when it cannot bind γ H2A.

To evaluate the genetic epistasis relationships between Brc1 and γ H2A, we introduced *brc1* Δ , *brc1-T672A* or *brc1*-K710M mutations into the hta-AQ background and tested genotoxin sensitivity relative to the parent strains. All of the brc1 mutants were sensitive to 4 mM HU, as was the hta-AQ parent, and this sensitivity was not further enhanced by combining the mutations (Supplementary Figure S2). These data suggest that in response to replication fork arrest caused by HU, the function of γ H2A can be largely if not entirely explained by its recruitment of Brc1. The results were largely the same in cells treated with 2 µM CPT (Supplementary Figure S2). However, when tested in media containing 5 µM CPT, the brc1-T672A hta-AQ or brc1-K710M hta-AQ strains were more sensitive than the parent strains. There are several possible explanations for these data, with our favoured model being that yH2A has both Brc1 dependent and independent functions in recovery from collapsed replication forks.

Sculpting of BRCT pSer + 3 binding pockets

To understand Brc1 pSer +3 binding and specificity determinants, we compared the Brc1 pSer +3 (Leu) recognition

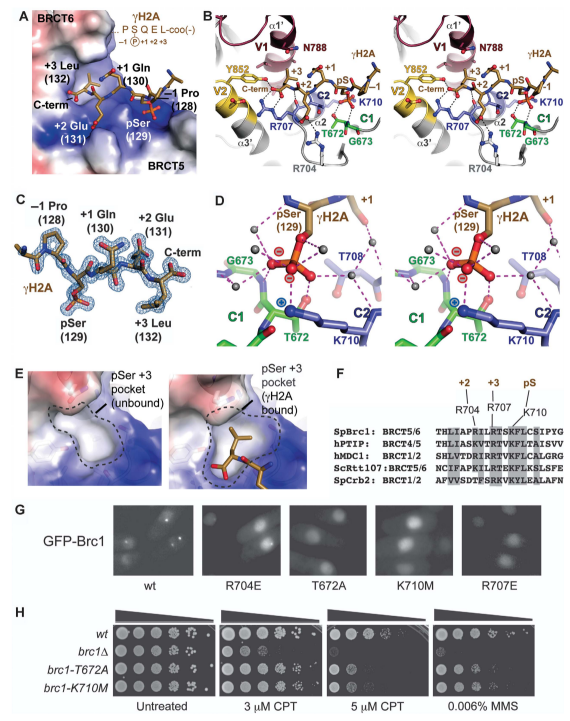


Figure 3 Brc1– γ H2A interactions. (**A**) The γ H2A-binding pocket of Brc1 BRCT₅–BRCT₆ harbours two positively charged grooves (blue electrostatic potential) for the pSer and pSer + 2 (Glu) residues, and a hydrophobic pocket for the pSer + 3 (Leu). The γ H2A peptide (gold) is displayed overlaid on an electrostatic surface of the Brc1 phosphoprotein-binding surface. (**B**) γ H2A is bound by constant BRCT pSer binding motifs C1 (green) and C2 (blue) and variable regions V1 (maroon) and V2 (yellow). A stereoview is displayed. (**C**) Final Sigma-A weighted 2Fo-Fc electron density (blue) at 1.45 Å is contoured at 1.5 σ and displayed for the peptide (gold). (**D**) Molecular interactions and water hydration shell for the γ H2A phosphoserine. Stereospecific coordination of the pSer phosphate moiety maximizes charge–charge interactions with C2 motif K710 epsilon amino group. (**E**) Structural plasticity of the Brc1 pSer + 3 Leu binding pocket. A partially formed + 3 pocket in the apo-Brc1 structure rearranges to interact with γ H2A Leu132 on phosphoprotein binding (right). (**F**) Sequence conservation of the Brc1 peptide-binding pocket of Brc1 ablates spontaneous foci formation for mutants T672A, K710M and R707E. Expression of the GFP-Brc1 proteins was induced in a wt strain for 18 h at 30°C and live fluorescence microscopy was performed. (**H**) Genetic analysis of the Brc1 γ H2A-binding mutants. Ten-fold serial dilutions of cells were exposed to the indicated DNA-damaging agent and incubated at 30°C for 3 days.

pocket with structures of the MDC1– γ H2AX complex (Stucki *et al*, 2005) and the BRCA1–BRCtide complex (Williams *et al*, 2004) (Figure 4). In all three structures, two variable surface

regions (V1 and V2), and one constant motif (C2) converge to form contiguous pSer +3 binding pockets (Figure 4A–C). Whereas V2 and C2 are structurally similar for all interac-

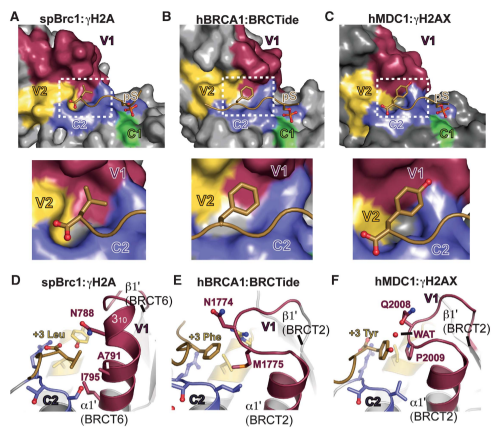


Figure 4 BRCT V-loop structural variability molds the pSer +3 binding pocket and defines peptide-binding specificity. (**A**) V-loop pSer +3 binding pocket molding in the Brc1– γ H2A complex. A surface representation of Brc1 is displayed for BRCT phosphoprotein interactions with constant regions (C1—green and C2—blue), and variable BRCT peptide-binding loops (V1—maroon and V2—yellow). Inset: The Brc1 pSer +3 binding pocket tightly contours around the γ H2A Leu132 carboxyl terminus. (**B**) hBRCA1–BRCTide complex constant and variable loops. Inset: BRCA1 phosphoprotein-binding pocket (RCSB 1T2V) is larger than Brc1, and binds Phe at γ H2AX pSer +3. (**C**) hMDC1– γ H2AX complex constant and variable loops. Inset: the mammalian γ H2AX-binding pocket is larger than the Brc1 pocket, and binds Tyr, but not Leu or Phe at the γ H2AX pSer +3 position. (**D**) Side view of the Brc1 pSer +3 binding pocket. V1 of Brc1 forms an extended loop and 3_{10} helix that surrounds γ H2AA pSer +3 Leu on three sides. (**E**, **F**) The BRCA1 and hMDC1 V1 loops do not form 3_{10} helical structures like Brc1, but adopt extended loops contributing two residues to γ H2AX pSer +3 binding pocket.

tions, the V1 regions (comprising the $\beta 1' - \alpha 1'$ connecting loop and $\alpha 1'$ of the second BRCT of the BRCT pair) show marked structural heterogeneity, and are a primary pSer +3 pocket sculpting and specificity determinant.

In Brc1, the V1 $\alpha l'$ is capped by a 3_{10} helix. Three $\alpha l'$ residues (Asn788, Ala791 and Ile795) form the base of the hydrophobic γ H2A pSer +3 (Leu132) side-chain-binding pocket. Two residues (Tyr852 from V2 and Arg707 from C2) directly bind the yH2A C-terminal carboxylate. Carboxyl terminus positioning by Tyr852 and Arg707, and precise pocket contouring around yH2A Leu132 Co1 and Co2, suggests that Brc1 binding to larger aromatic side chains (Phe or Tyr) would require major distortions of the binding pocket. Thus, Brc1 seems evolved to specifically bind Leu at pSer +3. In contrast to Brc1, two residues in extended $\beta 1' - \alpha 1'$ loops of BRCA1 and MDC1 make the pSer + 3 sidechain contacts. These fundamentally different V1 pocket configurations form larger pockets adapted for binding Phe/Tyr (BRCA1) (Williams et al, 2001; Manke et al, 2003) or Tyr (MDC1) (Stucki et al, 2005), but not Leu at the pSer +3 position as seen here for Brc1 (Figure 4). Thus, our current structural data show how sequence and structural variability in the BRCT V1 loops gives rise to markedly different pSer +3 binding pocket topologies and binding specificities.

Spontaneous Brc1 foci do not colocalize with Crb2 or Rad22

Crb2 and Brc1 both bind yH2A through C-terminal BRCT repeats; yet they have different functions: Crb2 is required for Chk1 activation and efficient DSB repair, whereas Brc1 is required for recovery from replication fork collapse. To investigate whether Crb2 and Brc1 localize into distinct structures, we first compared the foci patterns of GFP-Brc1 or YFP-Crb2 in unperturbed cells. This analysis revealed a large difference in the percentage of cells with GFP-Brc1 foci $(\sim 15\%)$ or YFP-Crb2 foci $(\sim 2\%)$ (Figure 5A). Analysis of a strain that expressed both fluorescently tagged proteins confirmed that most GFP-Brc1 foci do not overlap with CFP-Crb2 foci in untreated cells (Figure 5B). This analysis was extended by examining a strain that coexpressed GFP-Brc1 and Rad22-RFP, in which only $\sim 5\%$ of the GFP-Brc1 foci overlapped with Rad22-RFP foci (Figure 6A). This pattern contrasts with the nearly total overlap of YFP-Crb2 and Rad22-CFP foci in IR-treated cells (Du et al, 2003).

From these data, we hypothesized that most of the spontaneous Brc1 foci occur at sites of replication fork arrest, which do not activate the Crb2-dependent DNA damage checkpoint and are not substrates for Rad22-dependent HR repair. To test this hypothesis, we compared GFP-Brc1 and YFP-Crb2 foci in cells treated with HU, a compound that

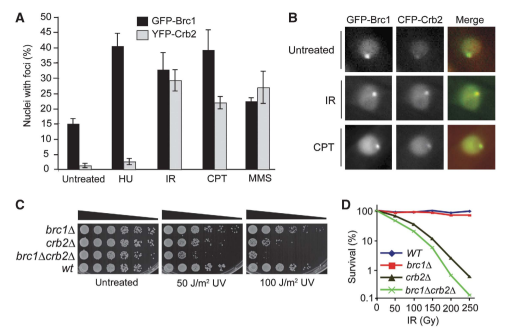


Figure 5 Specificity of DNA damage-induced foci formation by Brc1 and Crb2. (**A**) Foci formed by Crb2 and Brc1 increase in response to IR, CPT and MMS, whereas spontaneous and HU-induced Brc1 foci occur more frequently than Crb2. Cells expressing either GFP-Brc1 or 2YFP-Crb2 were grown for 17 h at 25°C in minimal medium before being split. Treatment was with 12 mM HU for 4 h, 30 μ M CPT for 4 h, 0.03% MMS for 5 h or 120 Gy IR plus 2 h recovery. (**B**) Brc1 and Crb2 foci colocalize following exogenous DNA damage. Cells expressing GFP-Brc1 (green) and CFP-Crb2 (red) were grown for 17 h at 25°C in minimal medium and left untreated or subjected to treatment with 30 μ M CPT for 5 h or 120 Gy IR plus 2 h recovery before imaging. The weak focus in the untreated CFP-Crb2 image is GFP bleed through into the red channel. (**C**) Synthetic additivity of *brc1* and *crb2* mutations. Five-fold serial dilutions of cells were exposed to the indicated doses of UV and incubated at 30°C for 2 days. (**D**) The *brc1* $\Delta crb2\Delta$ double mutant shows enhanced sensitivity to IR.

causes replication fork arrest by depleting dNTPs. Consistent with earlier results (Du *et al*, 2006), HU treatment did not significantly increase YFP-Crb2 foci (Figure 5A). In contrast, HU treatment caused a large increase in cells with GFP-Brc1 foci (Figure 5A). As observed for spontaneous GFP-Brc1 foci, HU-induced GFP-Brc1 foci were abolished in *hta-AQ* cells (Supplementary Figure S3).

Spontaneous Brc1 foci colocalize with rDNA

As HU-induced replication fork arrest triggers Brc1 foci formation, we inquired whether spontaneous Brc1 foci occur in genomic regions prone to fork arrest and genomic instability. The most obvious candidate was the ribosomal DNA (rDNA) repeats where polar replication fork barriers (RFBs) are required to maintain the integrity of the rDNA locus (Krings and Bastia, 2004; Sanchez-Gorostiaga *et al*, 2004). These repetitive rDNA arrays are located on the outer arms of chromosome III and protrude into the nucleolus (Uzawa and Yanagida, 1992). We therefore coexpressed GFP-Brc1 with RFP-tagged Gar1, a protein that associates the small nucleolar RNA. Approximately 60% of GFP-Brc1 foci were perinucleolar in untreated cells (Figure 6B), indicating that the majority of spontaneous Brc1 foci occur in the rDNA loci.

DNA damage-induced Brc1 foci colocalize with Rad22 and Crb2

As $brc1\Delta$ cells are sensitive to genotoxins such as CPT and MMS that cause DNA damage during S-phase, we hypothesized that exposure to these clastogens might induce Brc1 foci. Indeed, these agents caused a large increase in GFP-Brc1 foci (Figure 5A). IR had a similar effect, which was unexpected because $brc1\Delta$ cells are not IR sensitive (Figure 5D; Dovey and Russell, 2007). In all these treatments, GFP-Brc1 foci formation required γ H2A (Supplementary Figure S3). These observations suggested that besides forming foci in response to replication arrest, Brc1 also forms foci at DSBs. Indeed, we found that IR caused a large increase in the percentage of GFP-Brc1 foci that overlap with Rad22-YFP (Figure 6A). Moreover, the percentage of GFP-Brc1 foci that were perinucleolar decreased in response to IR (Figure 6B), suggesting that Brc1 is recruited to IR-induced DSBs occurring throughout the genome.

CPT, MMS and IR induction of GFP-Brc1 foci were paralleled by YFP-Crb2 foci formation (Figure 5A). To test whether Brc1 and Crb2 colocalize at DSBs, cells coexpressing GFP-Brc1 and CFP-Crb2 were exposed to IR or CPT. Both treatments caused extensive colocalization of CFP-Crb2 and GFP-Brc1 foci (Figure 5B).

The IR induction of GFP-Brc1 foci suggested that Brc1 might have an unappreciated function in DSB repair. We examined this possibility by carrying out genetic epistasis analysis with *brc1* Δ and *crb2* Δ . As observed earlier, *brc1* Δ cells were not sensitive to IR or UV. However, the *brc1* Δ *crb2* Δ double mutant displayed additive sensitivity to these genotoxins (Figure 5C and D). This result shows that Brc1 and Crb2 share a DSB repair function, or a unique function of Brc1 can be bypassed through a Crb2-dependent mechanism.

Discussion

Brc1 is a critical S-phase yH2A effector

Although MDC1– γ H2AX and Crb2– γ H2A interactions are critical for responding to IR-induced DSBs (Stewart *et al*, 2003; Nakamura *et al*, 2004; Stucki *et al*, 2005; Du *et al*,

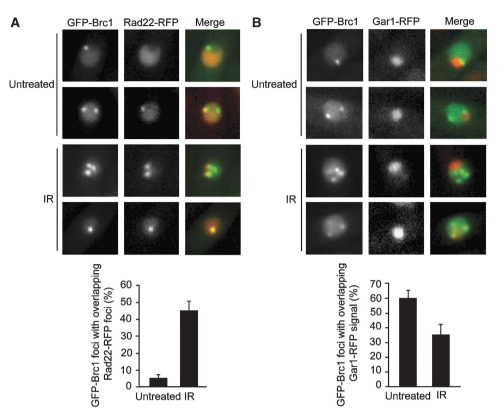


Figure 6 Brc1 foci localize to sites of genome instability. (A) IR-induced Brc1 foci frequently colocalize with the Rad22 DNA repair protein. Live cells expressing GFP-Brc1 (green) and Rad22-RFP (red) were photographed. Images were merged in the right panel. Samples were left untreated or exposed to 120 Gy IR followed by a 1 h recovery. Only $\sim 5\%$ of spontaneous GFP-Brc1 and Rad22-RFP foci colocalize under normal conditions, whereas an increased overlap in signal is observed following induction of DSBs. For each sample, foci were scored in at least 300 cells and mean values of colocalized foci per field were plotted with error bars representing the standard deviation of the mean. (B) Live cells expressing GFP-Brc1 (green) and nucleolar Gar1-RFP (red) were photographed. Images were merged in the right panel. Samples were left untreated or exposed to 120 Gy IR followed by a 1 h recovery. Quantification is shown on the right. For each sample, foci were scored in at least 300 cells and the mean values of colocalized foci per field were plotted with error bars representing the standard deviation of the mean.

2006; Lou et al, 2006; Kilkenny et al, 2008), the effectors of yH2A signalling in response to replication stress or S-phase DNA damage have been elusive. Here, we identify Brc1 as a critical yH2A-docking factor required for recovery from replication-associated DNA damage in fission yeast. With a high-resolution visualization of the Brc1-yH2A binding interface, our results reveal how key conserved determinants of phosphate recognition combined with evolutionary sculpting of BRCT pSer +3 binding pockets achieve maximum binding affinity with minimalist four-residue phosphorylated target recognition sequences. Disruption of this interface ablates Brc1 foci formation and confers sensitivity to genotoxins that disturb DNA replication. Taken together, these data show that Tel1/Rad3-regulated binding of Brc1 to yH2A is a critical chromatin-specific response to DNA damage in S-phase.

Distinct types of *γH2A*-dependent Brc1 foci

ATM and ATR checkpoint kinases regulate cellular responses to many forms of genotoxic stress, which vary with respect to the damage type, chromosomal context and the replicative state of the genome. These kinases also have many substrates, few of which are understood in detail. Chromatinspecific responses largely flow through γ H2A formation, raising the question of how these responses are tailored to specific circumstances of DNA damage. From the results presented here, it emerges that there are at least two γ H2A-docking proteins in *S. pombe*, Crb2 and Brc1, which both use C-terminal BRCT domains to bind γ H2A.

Notably, there are at least two types of Brc1 foci important for tailoring chromatin-specific responses to varying types of genotoxic stress: those that colocalize with Crb2 and Rad22. and those that do not. Both types require γ H2A. The former class is induced by clastogens that cause DSBs such as CPT or IR, whereas the latter class arises spontaneously and largely in the rDNA locus, or in response to HU-induced fork arrest. As Rad22 foci increase and Rad22 is required for viability in *brc1* Δ cells, Brc1 foci likely mark stalled replication forks. Failure to stabilize these forks leads to their collapse and creates a requirement for Rad22-mediated HR repair. Once a replication fork collapses, as when CPT poisons topoisomerase I, both Brc1 and Crb2 are required for efficient recovery along with Rad22-mediated HR repair. These Brc1 and Crb2 functions are substantially enhanced through yH2A interactions.

What accounts for the differential localization of Brc1 and Crb2 when both proteins bind γ H2A? Part of the explanation must involve Crb2's requirement for two types of histone modifications to form IR-induced foci (Du *et al*, 2004, 2006; Sanders *et al*, 2004). Proximal to its γ H2A-binding BRCT domains, Crb2 has Tudor domains that bind Set9-methylated histone H4 (H4K20Me) (Sanders *et al*, 2004; Botuyan *et al*,

2006; Du *et al*, 2006). Mutations abolishing γ H2A or H4K20Me binding are epistatic. As bulk H4K20Me is constitutive in fission yeast, DNA damage may alter chromatin structure and expose H4K20Me to control Crb2 localization (Du *et al*, 2006). So H4K20Me may be demethylated at stalled replication forks, or chromatin structure around stalled forks may not expose H4K20Me for productive engagement by the Crb2 Tudor domains. By relying on a histone code, checkpoint kinases can differentially regulate Crb2 and Brc1 localization by a shared histone modification.

 γ H2A's dual function in recruiting both Brc1 and Crb2 to chromatin at CPT-induced DSBs can explain why *hta-AQ* mutants are preferentially sensitive to genotoxins that collapse replication forks (Nakamura *et al*, 2004). As *hta-AQ* mutants are defective in recruiting both Brc1 and Crb2, while *set9* Δ mutants are defective only in recruiting Crb2, γ H2A recruitment of two factors explains the heretofore perplexing observation that the CPT sensitivity of *hta-AQ* mutants exceeds that of *set9* Δ mutants (Du *et al*, 2006).

Dual modes of recruiting Brc1 to damaged DNA?

The Brc1 yH2A-binding mutants are hypomorphic, which implies that Brc1 has both yH2A-dependent and yH2A-independent mechanisms of interacting with DNA lesions. As this situation resembles that for Crb2, it may be a common feature of proteins that interact with γ H2A (Du *et al*, 2006). The yH2A-independent functions of Brc1 may involve the BRCT₁₋₄ domains binding to other DNA or chromatin-associated ligands, although these are unlikely mediated by direct Brc1-phosphoprotein interactions. Our results suggest three specific possibilities. First, Brc1 may interact with Cut5^{TopBP1}, as this protein acts in DNA replication as well as DNA damage checkpoint signalling (Garcia et al, 2005), and it is required for the yH2A-independent mechanism of Crb2 function (Du et al, 2006). Second, Brc1 may bind ssDNA exposed at DNA lesions. Third, Brc1 may interact with nucleosomes at sites of DNA damage independent of post-translational modifications. However, expression of Brc1 BRCT5-BRCT6 domains is both necessary and sufficient for nuclear foci formation (Supplementary Figure S4), suggesting that binding to yH2A is critical for recruitment to sites of genome instability that can be detected microscopically.

Interestingly, the Brc1 BRCT₅–BRCT₆ interdomain linker is formed by two helices (α L1 and α L2) closely resembling the MDC1 helix-loop-helix linker structure (Stucki *et al*, 2005). This was unexpected because BRCT linker structures are the most structurally variable regions in tandem-BRCT repeat domains (Glover *et al*, 2004). Moreover, a conserved contiguous surface patch comprising 12 acidic (Asp, Glu) and polar residues (Tyr, Thr and Ser) populate the BRCT linker and neighbouring helix α 3 of Brc1–BRCT₅, or the equivalent region of MDC1–BRCT₁ (Figure 7A and B). Direct tethering of this electronegative BRCT surface (Figure 7A) adjacent to the histone octamer suggests that additional interactions with the positively charged histone-fold core particle surfaces may occur.

We hypothesize that the conserved electronegative patch of MDC1 and Brc1 γ H2A-interacting BRCT proteins seems adapted for recognition of histone core particle surfaces exposed on disruption of nucleosome architecture at DNA damage (Figure 7C). Charge-charge Brc1-histone interactions could facilitate γ H2A histone exchange and/or localized

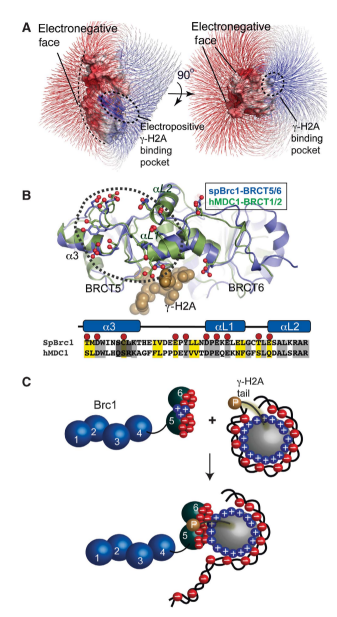


Figure 7 A conserved BRCT electronegative surface on γ H2A and γ H2AX-interacting proteins. (**A**) An electrostatic surface potential (displayed with field lines) for Brc1 shows a highly polarized bar magnet surface with an electropositive γ H2A interaction cleft (blue), and an extensive electronegative face (red). (**B**) A structural overlay of the mammalian γ H2AX-binding MDC1 tandem-BRCT domains with the yeast Brc1– γ H2A complex reveals an extensive conserved electronegative surface. (**C**) Schematic for a proposed two-site nucleosome interaction involving phosphorylation-dependant γ H2A tail binding and nucleosome-Brc1 charge–charge interactions. Docking to the electronegative surface may augment γ H2A histone tail docking through direct interactions with the positively charged DNA-binding nucleosome histone core fold.

remodelling of chromatin structure proximal to damaged DNA through an interaction predicted to compete with histone-DNA binding (Figure 7C). Consistent with this hypothesis, the Brc1 acidic surface exhibits the hallmark characteristics of a DNA mimic (Putnam *et al*, 1999; Putnam and Tainer, 2005).

Spontaneous Brc1 foci at rDNA loci

Brc1 foci are formed more frequently than Crb2 foci in untreated cells, suggesting an important function for Brc1

localization to chromatin during an unperturbed cell cycle. Spontaneous Brc1 foci often colocalize to the nucleolar periphery with the rDNA repeats (Figure 6B), so Brc1 may have specific functions in maintaining rDNA integrity. Eukaryotic cells have genetically programmed sites of replication pausing, the most abundant being polar RFBs in the rDNA loci (Krings and Bastia, 2004; Sanchez-Gorostiaga et al, 2004). RFBs may reduce replisome collisions with the transcription machinery (Takeuchi et al, 2003; Tsang and Carr, 2008), and defects in RFB function lead to genomic instability. Brc1 may therefore facilitate RFB formation or maintenance. An additional $\sim 40\%$ of spontaneous Brc1 foci are not perinucleolar (Figure 6B). Non-rDNA foci may be sites of Brc1 accumulation serving general functions in promoting the stability of genomic regions prone to fork pausing, such as non-specific sites of endogenous DNA damage.

BRCT-phosphoprotein recognition in the DNA damage response

Tandem-BRCT domains are key molecular readout modules for the assembly of phosphorylation-regulated DNA damage response effector and signalling complexes (Manke et al, 2003, Glover et al, 2004). In mammalian cells, ATM and ATR phosphorylate over 700 proteins in response to genotoxic insult (Matsuoka et al, 2007). With over 2400 curated BRCT domain sequences in the PFAM database (http:// pfam.sanger.ac.uk/family?acc = PF00533), it is likely that many other critical phosphoprotein-BRCT interaction combinations have evolved with unique interaction specificities. The molding of BRCT phosphoprotein-binding specificity pockets through evolution of variable BRCT insertion loops, as observed here for the Brc1-yH2A complex, provides a probable general mechanism for generating the variability required for assembly of specific and distinct BRCT-phosphoprotein complexes in vivo.

Materials and methods

Strains, media and genetic analysis

Strain genotypes are listed in Supplementary Table S1. Growth media and methods for *S. pombe* were performed as described (Moreno *et al*, 1991). Ectopic expression of *pRep41-N-GFP-brc1*⁺ for microscopy was under control of the thiamine-repressible *nmt41* promoter. Induction of plasmid expression in selective medium lacking thiamine was for 18–21 h. Spot assays were performed by plating 5- or 10-fold serial dilutions of exponentially growing cells onto YES in the absence or presence of the indicated DNA-damaging agents. Plates were incubated at 30°C and scanned after 2–3 days of growth. The survival assay was performed by counting cells plated in triplicate onto rich medium after exposure to the indicated IR doses. Normalization was to the untreated sample.

Microscopy

Cells were photographed using a Nikon Eclipse E800 microscope equipped with a Photometrics Quantix CCD camera. Cells were treated with 12 mM HU (Sigma), $30 \,\mu$ M CPT (Sigma) or 0.03 % MMS (Sigma) and grown for 4–5 h at 30°C or 5–6 h at 25°C. For IR treatment, mid-log phase cells were exposed to the indicated dose and allowed time for recovery at 30°C for 1–4 h. Quantitation was performed by scoring 500 or more nuclei from at least two independent experiments. For DAPI-staining experiments, cells were fixed for 10 min at RT in 70% EtOH and washed with H₂O before treatment with 5 μ l of DAPI (500 μ g/ml).

Protein expression, purification and crystallization

Brc1-BRCT₅-BRCT₆ (residues 659-878) was expressed in *Escherichia coli* Rosetta2 (DE3) (Novagen) at 20°C in Terrific Broth from pET15b as N-terminal 6-histidine tagged protein. After Ni-NTA affinity chromatography and elution with an on-resin thrombin cleavage that removed the 6his-tag, BRCT₅-BRCT₆ was purified with Superdex 75 gel filtration chromatography (GE-Amersham) in BRCT buffer (200 mM NaCl, 20 mM Tris-HCl pH 7.5, 0.1% β -mercaptoethanol). The T672A variant was introduced using the Quikchange (Stratagene).

Crystallization data collection

Crystals of selenomethionine-derivitized BRCT₅-BRCT₆ were grown at 22°C by mixing 2 μ l of 15 mg/ml BRCT₅-BRCT₆ in BRCT buffer with 2 μ l of well solution 1 (20–22% w/v PEG3350 100 mM NaCl, 20 mM ammonium phosphate, 0.1 M sodium citrate pH 5.0). Single crystals were obtained with streak seeding from clustered monoclinic rods that grew within 1–2 days. Native orthorhombic crystals of apo-BRCT₆ used for refinement were grown using well solution 2 (20–22% w/v PEG 3350, 100 mM NaCl, 100 mM K₂HPO₄/ KH₂PO₄ pH 5.0).

All peptides used in this study were synthesized at the Scripps Research Institute core facility. The BRCT₅-BRCT₆- γ H2A complex was preformed before crystallization by mixing BRCT₅-BRCT₆ at 10 mg/ml in BRCT buffer with γ H2A phosphopeptide (Ac-KPpSQEL-COO-) at a 1.5:1 phosphopeptide:protein ratio. In all, 30 × 30 × 500 μ M orthorhombic needles of the complex grew at 22°C within 2–3 days after mixing 1 μ l of complex with 1 μ l well solution 3 (17–20% PEG3350, 200 mM ammonium chloride). For data collection, the crystals were flash frozen in liquid nitrogen after transfer to cryo-protectant solutions comprised of the appropriate mother liquor supplemented with 26% ethylene glycol.

X-ray diffraction data collection and processing

All single crystal X-ray diffraction data were collected at the SIBYLS beamline (BL12.3.1) at the Advanced Light Source, Lawrence Berkeley National Laboratory. X-ray data reduction and scaling was performed with the HKL2000 suite (Otwinowski and Minor, 1997).

X-ray structure determination and refinement

All four selenium positions in the crystallographic asymmetric unit of the monoclinic apo-BRCT₅–BRCT₆ space group (2 selenomethionines per 220 residue BRCT₅–BRCT₆ polypeptide) were located and refined with SOLVE (Terwilliger and Berendzen, 1999) and a 3wavelength Se-MET monoclinic (P21) MAD data set (Supplementary Table S2). Crystallographic phases and experimental electron maps were improved by density modification in RESOLVE (Terwilliger, 2000). Automated model building in RESOLVE and manual fitting in O (v10.0) (Jones *et al*, 1991) produced a model that was preliminarily refined to 1.8 Å using REFMAC (Murshudov *et al*, 1997; Winn *et al*, 2001) and the MAD data set remote wavelength (λ 3). This model was used for molecular replacement search with MOLREP (Vagin and Teplyakov, 2000) to determine the orientation of the structure in the related native P212121 crystal form. Subsequent cycles of positional and anisotropic temperature factor refinement against the 1.55 Å orthorhombic data set.

Phases for the $BRCT_5-BRCT_6-\gamma$ H2A complex were obtained by molecular replacement in MOLREP and the unbound $BRCT_5-BRCT_6$ structure. The $BRCT_5-BRCT_6-\gamma$ H2A complex crystal form contains one complex in the asymmetric unit. The phosphorylated γ H2A peptide chain was traced into unambiguous sigma-A weighted 1.45Å Fo-Fc and 2Fo-Fc difference electron density maps, and the model was refined as for the apo-BRCT₅-BRCT₆ at 1.55Å $(R/R_{free} = 17.9/22.5)$ and the BRCT₅-BRCT₆- γ H2A complex at 1.45Å $(R/R_{free} = 13.1/18.8)$ exhibit excellent geometric parameters (Supplementary Table S2).

Accession codes

Coordinates and structure factors for BRCT₅-BRCT₆ (RCSB accession code, 3L40) and the BRCT₅-BRCT₆- γ H2A complex (RCSB accession code, 3L41) are in the Protein Data Bank.

Peptide binding studies

FP equilibrium binding measurements with amino-terminal fluorescein (FITC)-conjugated peptides were taken on a Fluoromax-3 (Horiba/Yobin-Yvon) with constant wavelength excitation at 490 nm and emission monitored at 520 nM. Titrations used a constant concentration (10 nM) of phosphorylated (FITC-KRTGKPpSQEL) or unphosphorylated (FITC-KRTGKPSQEL) H2A peptide, and increasing amounts ($25 \text{ nM}-80 \mu M$) of WT-BRCT₅–BRCT₆ or T672A–BRCT₅–BRCT₆. A measure of 400 μ l binding reactions were equilibrated for 10 min at 20°C in binding buffer (40 mM NaCl, 20 mM Tris–HCl pH 7.5, 5% glycerol, 0.1 mg/ml bovine serum albumin, 0.1% β-mercaptoethanol) before FP measurement. Equilibrium binding constants were calculated with KaleidaGraph (v4.03).

Small angle X-ray scattering

SAXS scattering data for samples of apo- for apo-BRCT₅–BRCT₆ (at 2.5, 5.0 and 10.0 mg/ml), and for the BRCT₅–BRCT₆– γ H2A complex (at 5.0 and 10.0 mg/ml) in the momentum transfer range 0.015 < S <0.388 Å⁻¹ was collected at beamline 12.3.1 (SIBYLS) at the Advanced Light Source.

SAXS data analysis

No evidence of aggregation for samples was evident as judged by linearity of Guinier plots. Radius of gyration (Rg) and pair distance distribution functions P(r) were evaluated using GNOM (Svergun, 1992). Theoretical scattering was generated with CRYSOL (Svergun *et al*, 1995). *Ab initio* solution envelopes were obtained by averaging 10 independent dummy residue models from reciprocal space minimization in GASBOR (Svergun *et al*, 2001).

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Supplementary data

Supplementary data are available at *The EMBO Journal* Online (http://www.embojournal.org).

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Conflict of interest

The authors declare that they have no conflict of interest.

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