Mechanistic insight into cadmium-induced inactivation of the Bloom protein

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Cadmium is a toxic metal that inactivates DNA-repair proteins via multiple mechanisms, including zinc substitution. In this study, we investigated the effect of Cd2+ on the Bloom protein (BLM), a DNA-repair helicase carrying a zinc-binding domain (ZBD) and playing a critical role to ensure genomic stability. One characteristics of BLM-deficient cells is the elevated rate of sister chromatid exchanges, a phenomenon that is also induced by Cd2+. Here, we show that Cd2+ strongly inhibits both ATPase and helicase activities of BLM. Cd2+ primarily prevents BLM-DNA interaction via its binding to sulfhydryl groups of solvent-exposed cysteine residues and, concomitantly, promotes the formation of large BLM multimers/aggregates. In contrast to previously described Cd2+ effects on other zinc-containing DNA-repair proteins, the ZBD appears to play a minor role in the Cd2+-mediated inhibition. While the Cd2+-dependent formation of inactive multimers and the defect of DNA-binding were fully reversible upon addition of EDTA, the inhibition of the DNA unwinding activity was not counteracted by EDTA, indicating another mechanism of inhibition by Cd2+ relative to the targeting of a catalytic residue. Altogether, our results provide new clues for understanding the mechanism behind the ZBD-independent inactivation of BLM by Cd2+ leading to accumulation of DNA double-strand breaks.

Bloom’s syndrome (BS) is a rare, autosomal and recessive disease resulting from the mutational inactivation of a human RecQ family helicase encoded by the blm gene1. BS is characterized by proportional dwarfism, erythema on sun-exposed skin, hyper- or hypo-pigmented skin areas, immunodeficiency and subfertility2. Persons with BS have a high predisposition to cancer and increased risk for early-onset type-II diabetes3. The blm gene encodes BLM, a 1417-amino acids protein containing several conserved motifs including a zinc-binding domain (ZBD). Previous works have shown that mutation of any of the four conserved Cys residues of the ZBD leads to the BS4,5. Moreover, we have previously shown that the ZBD of RecQ helicases plays a key role in protein folding and is involved in DNA-binding6. Thus, alteration of the zinc coordination state and potentially metal-catalyzed oxidation could impair BLM-mediated DNA-repair processing events. In addition to numerous cytological characteristics including high rates of loss of heterozygosity7–9, chromosome abnormalities (telomere fusions, ring chromosomes and quadriradial chromosomes10), the most striking feature of BLM-deficient cells or cells bearing an impaired BLM mutant is characterized by elevated rates of sister chromatid exchanges (SCEs)11. Interestingly, it was shown that Cadmium (Cd) also provoked elevated rates of SCEs in human cell cultures12. Thus, the effect of Cd2+ on human cell lines shares cytological characters with BLM-deficient cells, establishing a connection between BLM and Cd2+.

Cd2+ is considered as an important health hazard due to its long retention time and bioaccumulation in human body13. Epidemiological and animal experiments have revealed multifactorial carcinogenic properties of cadmium14. Exposure to Cd2+ is associated with cancers of lung, prostate, pancreas and kidney15. Among the various carcinogenic effects of Cd2+, DNA damage accumulation due to inhibition of DNA-repair enzymes is considered as one of the major underlying process16,17. Unlike numerous toxic metal compounds, Cd2+ is considered as weakly mutagenic. Nevertheless, Cd2+ is known to severely increase the genotoxic effects of various mutagens in mammalian cells, including ionizing radiations and DNA alkylating agents used at low non-cytotoxic concentrations18,19. Many studies using yeast or human cells suggest that DNA-repair systems represent highly sensitive targets for Cd2+. However, the precise mechanism behind carcinogenicity remains to be determined. Although

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many in vitro studies on Cd\textsuperscript{2+}-mediated toxic effects have been performed with proteins involved in the Base and Nucleotide Excision Repair (BER/NER)\textsuperscript{20,21}, Mismatch Repair (MMR)\textsuperscript{22,23} and Non-Homologous End-Joining (NHEJ)\textsuperscript{24}, it is a difficult task to highlight a general/common mechanism underlying Cd\textsuperscript{2+}-mediated inhibition of DNA-repair systems. Nevertheless, it appears that detrimental effects of Cd\textsuperscript{2+} on DNA-repair proteins occur through the binding of Cd\textsuperscript{2+} to functional sulfhydryl groups\textsuperscript{23,25}, and the replacement of Zn\textsuperscript{2+} by Cd\textsuperscript{2+} in ZBDs represents one cause for protein dysfunctions.

BLM facilitates homologous recombination (HR) between diverged homologous sequences\textsuperscript{26}. Among the different DNA-repair systems, HR is remarkable by its ability to accurately repair DNA double-strand breaks. Defects in the HR machinery are often associated with cell cycle deregulations, apoptosis or genomic instability. Until now, HR remains the only DNA-repair pathway for which there is no clear evidence of Cd\textsuperscript{2+}-dependent inhibition, although previous studies have shown characteristic features of HR dysfunction following Cd\textsuperscript{2+} uptake such as elevation of SCEs\textsuperscript{12,15} and deregulation of the MRE11-dependent pathway that interacts with the HR machinery\textsuperscript{24}.

Based on previous observations showing inhibitory effects of Cd\textsuperscript{2+} on zinc-containing DNA-repair proteins and taking into account characteristic phenotypes of Cd\textsuperscript{2+}-exposed human and yeast cells\textsuperscript{2,2}, we addressed in the present study the molecular mechanisms of Cd\textsuperscript{2+}-dependent BLM inactivation. We demonstrated that Cd\textsuperscript{2+} strongly inhibits both the ATPase and helicase activities of recombinant BLM and that Cd\textsuperscript{2+} primarily prevents BLM-DNA interaction due to the formation of large BLM multimers/aggregates. This formation of large multimers/aggregates is dependent on the protein redox state and is fully reversible by EDTA. Consequently, EDTA rescues the DNA-binding defect in the presence of Cd\textsuperscript{2+}. Interestingly, EDTA did not fully counteract the Cd\textsuperscript{2+}-dependent inhibition of the proper DNA unwinding activity. Altogether, our results indicate that Cys in the reduced state are primarily targeted by Cd\textsuperscript{2+}. Surface and solvent-exposed Cys mediate the Cd\textsuperscript{2+}-dependent formation of aggregates (reversible by EDTA) whereas at least one additional residue, playing a key role in the catalytic process, is targeted by Cd\textsuperscript{2+} in an irreversible manner.

**Results**

The full-length BLM helicase (BLM\textsuperscript{full-length}) is composed of 1417 amino acid and contains the helicase core (BLM\textsuperscript{642–1290}) harboring two RecA domains and forming ATP-binding sites, the ZBD (pos. 994–1068) and the winged-helix domain (pos. 1069–1189), whereas the N- and C-terminal domains are more related to protein-protein interactions and nuclear localization\textsuperscript{27,28}. BLM\textsuperscript{642–1290} used throughout this study together with BLM\textsuperscript{full-length} and *E. coli* RecQ (RecQ\textsuperscript{E.coli}), displays full activities comparable to BLM\textsuperscript{full-length}\textsuperscript{27}.

**Cadmium severely impairs both helicase and ATPase activities of BLM.** To determine the effect of Cd\textsuperscript{2+} on BLM activities, we first measured the helicase activities of BLM\textsuperscript{full-length} and BLM\textsuperscript{642–1290} in the presence of increasing concentrations of CdCl\textsubscript{2}. DNA unwinding activities of both proteins were strongly inhibited by Cd\textsuperscript{2+} as measured by a radioactive assay: 95% and 100% of inhibition at 5 and 10 \(\mu\)M Cd\textsuperscript{2+}, respectively (Fig. 1a). Interestingly, the RecQ\textsuperscript{E.coli} unwinding activity was actually inhibited by Cd\textsuperscript{2+} but to a much lesser extent compared to BLM proteins. Indeed, the same inhibition level (>95%) of RecQ\textsuperscript{E.coli} was obtained at 100 \(\mu\) M CdCl\textsubscript{2}. These results, suggesting that BLM represents a more sensitive target for Cd\textsuperscript{2+}, were confirmed by stopped-flow FRET experiments allowing measurements of the unwinding kinetic rate constant of BLM by using partial duplex DNA labeled with fluorescein and hexachlorofluorescein as a donor and acceptor, respectively\textsuperscript{20,29} (Supplementary Fig. S1; see Supplementary Table S1 for details about the structure of the DNA substrate): the levels of Cd\textsuperscript{2+}-dependent inhibition of BLM\textsuperscript{full-length} and BLM\textsuperscript{642–1290} helicase activities were similar and significantly higher than that observed with RecQ\textsuperscript{E.coli} (=10-fold higher). Thus, Cd\textsuperscript{2+} displays a selective profile for BLM and most likely targets its helicase core since no significant difference in the Cd\textsuperscript{2+}-dependent inhibitions of BLM\textsuperscript{full-length} and BLM\textsuperscript{642–1290} was observed.

We next measured the ATPase activity of the three proteins in the presence of increasing CdCl\textsubscript{2} concentrations. Again, Cd\textsuperscript{2+} strongly inhibited the ATPase activities of both BLM\textsuperscript{full-length} and BLM\textsuperscript{642–1290}, while the ATPase activity of RecQ\textsuperscript{E.coli} was much less inhibited (Fig. 1b). The corresponding IC\textsubscript{50} values (inhibition concentration 50%) determined for BLM\textsuperscript{full-length} and BLM\textsuperscript{642–1290} were 6.7 and 7.3 \(\mu\)M, respectively, whereas the IC\textsubscript{50} was much higher (65 \(\mu\)M) for RecQ\textsuperscript{E.coli} (Table 1). This result closely parallels, at least qualitatively, the one obtained for the inhibition of DNA unwinding activity as explained above.

**Stoichiometry of Cadmium binding to BLM.** Before addressing the mechanism of BLM inhibition, we assessed whether BLM is a direct target of Cd\textsuperscript{2+}. We then investigated the Cd\textsuperscript{2+}:BLM\textsuperscript{642–1290} stoichiometry using a fluorescence-based assay (see Methods). The Cd\textsuperscript{2+}:RecQ\textsuperscript{E.coli} stoichiometry was studied in parallel for comparison. We found a stoichiometry of 11–12 Cd\textsuperscript{2+} per BLM\textsuperscript{642–1290} molecule (Fig. 2a, left) and a significantly lower stoichiometry was found for RecQ\textsuperscript{E.coli}, 6–7 Cd\textsuperscript{2+} per protein (Fig. 2a, right). Interestingly, each representation displayed two slopes, reflecting two types of sites characterized by distinct accessibilities and affinities. Previous studies have reported that free sulfhydryl groups of Cys are good candidates for Cd\textsuperscript{2+} binding\textsuperscript{23,25}. BLM\textsuperscript{full-length} and BLM\textsuperscript{642–1290} contain 30 and 19 Cys, respectively (11 in RecQ\textsuperscript{E.coli}). To determine whether Cd\textsuperscript{2+} actually targets Cys of BLM\textsuperscript{642–1290} and RecQ\textsuperscript{E.coli}, stoichiometry experiments were repeated in the presence of N-Ethylmaleimide (NEM), a thiol-alkylating agent that forms stable covalent thioether bonds with sulfhydryls of reduced Cys. NEM treatment significantly altered Cd\textsuperscript{2+} stoichiometry with two features: the Cd\textsuperscript{2+}:protein stoichiometry was decreased from 11–12 to 5 and from 6–7 to 3–4, for BLM\textsuperscript{642–1290} and RecQ\textsuperscript{E.coli}, respectively (Fig. 2b). Second, in contrast to experiments performed without NEM, the stoichiometry curves displayed one slope, corresponding to the low-affinity binding site cluster for both proteins. Altogether, the results show that these helicases display at least two types of Cys clusters. Assuming that (i) Cd\textsuperscript{2+} possesses a higher affinity for Cys localized at the surface compared to residues localized in the protein core, (ii) NEM only interacts with
surface residues for steric hindrance reason, we conclude that the 1st Cys cluster (surface) is characterized by high affinity for Cd$^{2+}$ without NEM but does not anymore interact with Cd$^{2+}$ in the presence of NEM while the 2nd cluster (protein core) is characterized by weaker affinity for Cd$^{2+}$ due to lower accessibility and is not influenced by NEM. We estimated that the 1st and 2nd clusters are composed by 6–7 and 5 Cys, respectively, in BLM642–1290 (3 and 3–4, respectively, in RecQ$^{E.coli}$). The remaining Cys that do not interact with Cd$^{2+}$ could be related to residues totally buried in the protein structure. The possible implication of the BLM zinc finger motif is addressed in the following section together with the issue of the Cadmium effect on the proper BLM/DNA interaction.

Cd$^{2+}$ binding to BLM primarily impairs its DNA-binding activity. To gain further insight into the mechanism of Cd$^{2+}$-induced inhibitions of DNA unwinding and ATPase activities, we examined DNA-binding activities of BLM$^{642–1290}$ and BLM$^{full-length}$ in the presence of increasing CdCl$_2$ concentrations by using a steady-state fluorescence anisotropy assay$^{31}$. We first determined DNA-binding isotherms curves (i) in the absence of Cd$^{2+}$ to define the experimental conditions in terms of protein concentrations (i.e. high enough above the $K_d$ to ensure DNA saturation) and (ii) in the presence of Cd$^{2+}$ and absence of any protein to determine the CdCl$_2$ concentration range in which the fluorescence anisotropy of the fluorescein-labeled DNA was not influenced by direct

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**Figure 1.** Effect of Cd$^{2+}$ on DNA unwinding (a) and ATPase (b) activities of BLM$^{642–1290}$ (left), BLM$^{full-length}$ (middle) and RecQ$^{E.coli}$ (right). (a) DNA helicase activity was revealed by a radioactive assay as described in Methods. 40 nM of proteins were pre-incubated in the presence of increasing concentrations of CdCl$_2$ for 2 min at 37 °C. The unwinding reaction was initiated by the addition of the DNA substrate (fork duplex substrate; see Supplementary Table S1) and was carried out for 20 min at 37 °C. NE: no enzyme; ∆: Heat denatured DNA substrate. (b) 200 nM of proteins were pre-incubated in the presence of increasing concentrations of CdCl$_2$ for 2 min at 37 °C before initiation of the ATPase activity. The ATPase activity was carried out for 10 min at 37 °C (see Methods for more details).

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**Table 1.** Experimentally determined IC$_{50}$ values characterizing inhibitions of ATPase and DNA-binding activities by Cd$^{2+}$.

| Activity          | Determined IC$_{50}$ values (μM) |
|-------------------|----------------------------------|
|                   | BLM$^{642–1290}$ | BLM$^{full-length}$ | RecQ$^{E.coli}$ |
| ATPase            | 7.3                | 6.7                | 65             |
| DNA-binding       |                    |                    |                |
| ssDNA (18-nt)     | 9.22               | 6.67               | 41.71          |
| ssDNA (24-nt)     | 10.22              | 6.63               | 40.22          |
| ssDNA (40-nt)     | 9.98               | 6.64               | 39.17          |
| dsDNA (24-bp)     | 8.17               | 6.84               | 44.12          |
Cd\(^{2+}\)-DNA interactions (data not shown). A protein concentration of 200 nM and CdCl\(_2\) concentrations up to 100 \(\mu\)M were found to correspond to optimal conditions and then, were used in subsequent experiments.

BLM\(^{\text{full-length}}\) and BLM\(^{642-1290}\) displayed similar inhibition profiles of the DNA-binding activities (Fig. 3). The Cd\(^{2+}\)-dependent inhibitions of the protein-DNA interaction were efficient, with IC\(_{50}\) values in the 6.6–10.2 \(\mu\)M range (Table 1). These values were consistent with those derived from the inhibition of helicase and ATPase activities. Moreover, the inhibition profiles were similar, regardless of the nature of the DNA substrate, single- (ss) or double-stranded (ds) DNA (Fig. 3a) or the DNA length, from 18- to 40-mer (Fig. 3b). However, the RecQ\(_{\text{E.coli}}\) DNA-binding activity was weakly affected by Cd\(^{2+}\) (IC\(_{50}\) = 39–44 \(\mu\)M), as observed for its ATPase or helicase activity, confirming that BLM is more sensitive to Cd\(^{2+}\) than RecQ\(_{\text{E.coli}}\).

The binding of BLM\(^{642-1290}\) to DNA was next studied in the presence of Cd\(^{2+}\) and various amino-acids such as Cys, His and Val. Among the different amino-acids, Cys is the residue forming by far the most stable complex\(^{32}\). Accordingly, when amino-acids were pre-incubated with both Cd\(^{2+}\)/DNA before addition of the protein, the BLM\(^{642-1290}\) DNA-binding activity was fully restored by Cys but only partially restored by His or Val (Supplementary Fig. S2). The counteracting effect of Cys was also observed on the ATPase activity of BLM\(^{642-1290}\) (data not shown). By contrast, the addition of amino-acids after pre-incubation of BLM\(^{642-1290}\) with Cd\(^{2+}\)/DNA did not restore the DNA-binding activity, regardless of the nature of the amino-acid (Supplementary Fig. S2). The absence of protective/competition effect by Cys in the latter case can be ascribed to irreversible binding of Cd\(^{2+}\) to BLM sulfydryl groups and EDTA only was able to reverse the Cd\(^{2+}\)-mediated inhibition of DNA-binding (see below).

Taking into consideration that BLM contains a zinc finger that is composed of four conserved Cys\(^{5}\), we next investigated whether the BLM ZBD could be a target site for Cd\(^{2+}\). Mutation of any one of these Cys leads to the BS and inactivates BLM activity both in vitro and in vivo\(^{6}\). First, we wondered whether Zn\(^{2+}\) and Cd\(^{2+}\) would be able to bind to the same site and whether the replacement of Zn\(^{2+}\) by Cd\(^{2+}\) would lead to a conformational change of the BLM ZBD and, consequently, impairs BLM activity. To further investigate the mechanism of Cd\(^{2+}\)-mediated inhibition and the interplay between Cd\(^{2+}\) and Zn\(^{2+}\), the influences of both metals on the DNA-binding step were compared using the fluorescence anisotropy-based assay. Interestingly, Cd\(^{2+}\) but also Zn\(^{2+}\) inhibited the DNA-binding step of BLM\(^{\text{full-length}}\), BLM\(^{642-1290}\) and RecQ\(_{\text{E.coli}}\), however to different extents (Supplementary Fig. S3). Up to 50 \(\mu\)M, Cd\(^{2+}\) alone was consistently more potent than Zn\(^{2+}\) alone for inhibition. Beyond 50 \(\mu\)M, both metals inhibited BLM\(^{\text{full-length}}\) and BLM\(^{642-1290}\) in a similar manner. To note, no inhibition was observed for BLM\(^{642-1290}\) at low Zn\(^{2+}\) concentrations (<15 \(\mu\)M). In this concentration range, the inhibitory effect of Cd\(^{2+}\) was not reversed by addition of Zn\(^{2+}\). The Cd\(^{2+}/\text{Zn}^{2+}\) combination was even more efficient for inhibiting DNA-binding activity than Cd\(^{2+}\) alone, a typical synergistic inhibition phenomenon. Such a behavior

Figure 2. Study of the Cd\(^{2+}\)-protein stoichiometries for BLM\(^{642-1290}\) (left) and RecQ\(_{\text{E.coli}}\) (right) before (a) or after (b) treatment with 0.5 mM N-Ethylmaleimide (NEM). Increasing concentrations of CdCl\(_2\) (0–37.5 \(\mu\)M) were added to 0.5 \(\mu\)M of BLM\(^{642-1290}\) or RecQ\(_{\text{E.coli}}\) diluted in Tris-HCl buffer (50 mM, pH 8.0) supplemented with 50 mM NaCl and 1 mM DTT. The mixture was further incubated for 5 min at 25 °C. Protein-Cd\(^{2+}\) complexes were then discarded using Q-sepharose beads. The Cd\(^{2+}\)-protein stoichiometries were deduced from the determination of free Cd\(^{2+}\) remaining in solution, using the fluorescence-based Measure-iT Cadmium assay as described in Methods.

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was also observed with BLM\textsuperscript{full-length}, except that low Zn\textsuperscript{2+} concentrations were more efficient for inhibiting the DNA-binding step of BLM\textsuperscript{full-length} compared with BLM\textsuperscript{642–1290}. Regarding RecQ\textsubscript{E.coli} DNA-binding activity which was only partially inhibited by Zn\textsuperscript{2+} alone (85% of activity at 100 \(\mu\)M), the more potent effect of Cd\textsuperscript{2+} was not reversed by addition of Zn\textsuperscript{2+}.

Altogether, these results indicate that, under our experimental conditions, Cd\textsuperscript{2+} and Zn\textsuperscript{2+} do not bind competitively to the same site. It is important to note that, especially with Cys\textsubscript{4} ZBD (corresponding to BLM or RecQ\textsubscript{E.coli} ZBDs), Cd\textsuperscript{2+} forms much more stable complexes than zinc\textsuperscript{33}. Taking into account that a large excess of Zn\textsuperscript{2+} over Cd\textsuperscript{2+} could be required for efficient competition (a condition which is not compatible with the proper Zn\textsuperscript{2+}-dependent inhibition described above), we cannot definitively confirm or ruled out a targeting of the BLM ZBD by Cd\textsuperscript{2+}. However, it is unlikely that the ZBD alone accounts for the Cd\textsuperscript{2+}-dependent inhibition since BLM and RecQ\textsubscript{E.coli} are characterized by different susceptibilities to Cd\textsuperscript{2+}; this differential susceptibility appears to be more related to the larger number of targeted Cys in the case of BLM, as suggested by stoichiometry experiments. The relationship between the defect of DNA-binding and the binding of Cd\textsuperscript{2+} to surface Cys residues was further investigated below by studying the Cd\textsuperscript{2+} effect on the multimeric status of BLM.

**Cd\textsuperscript{2+} induces BLM oligomerization.** During the course of our experiments, we observed that high Cd\textsuperscript{2+} concentrations (typically >30 \(\mu\)M) induced BLM precipitation in samples. Size-exclusion chromatography analysis showed that while BLM\textsuperscript{642–1290} alone was eluted as a monomer as previously reported\textsuperscript{34}, its elution profiles when pretreated with Cd\textsuperscript{2+} displayed three peaks, corresponding to molecular weights of 75, 150 and >205 kDa, respectively (Fig. 4a), indicating that Cd\textsuperscript{2+} induced protein oligomerization. We further studied the oligomeric status of BLM in the absence and presence of Cd\textsuperscript{2+} by Dynamic Light Scattering (DLS) (Fig. 4b). DLS analysis confirmed size-exclusion chromatography experiments. Indeed, without Cd\textsuperscript{2+}, BLM\textsuperscript{642–1290} was monodisperse in solution and characterized by a radius of 3.44 nm, compatible with a monomeric form (peak I). The addition of Cd\textsuperscript{2+} dramatically modified the profile of size distribution. The peak I disappeared and was replaced by two peaks (II+III) corresponding to large BLM multimers, probably non-specific cross-linked aggregates (with MW of 3.2 \(\times\) 10\textsuperscript{5} and 1.3 \(\times\) 10\textsuperscript{7} kDa, respectively). Addition of EDTA on Cd\textsuperscript{2+}-treated samples of BLM\textsuperscript{642–1290} dissociated higher-order oligomers leading to a recovery of the monomeric form as shown by size-exclusion chromatography (Fig. 4a) or DLS (Fig. 4b; peak III, radius 3.95 nm). Altogether, these results show that Cd\textsuperscript{2+} induces the formation of higher-order BLM oligomers that is reversible upon addition of EDTA.
BLM multimers/aggregates. Note that both Cd$^{2+}$ DNA-binding. In other words, the DNA-binding site should be completely hidden in the context of these large measured in the presence of Cd$^{2+}$ unwinding kinetic rate constant and the corresponding reaction amplitude characterizing BLM642–1290 (Fig. 6a),

Cys of the 1st cluster (surface residues as defined above) mainly mediate the Cd$^{2+}$ its own on the BLM642–1290 DNA-binding activity, fully counteracted the inhibitory effect of Cd$^{2+}$ DNA (Fig. 5a), in accordance with results shown in Fig. 3. To note, NEM that did not lead to significant effect on BLM642–1290 in the presence of 200 μM CdCl$_2$. BLM642–1290 alone is characterized by peak I. BLM642–1290 in the presence of Cd$^{2+}$ is characterized by two peaks (II + II'). BLM642–1290 in the presence of 200μM CdCl$_2$ + 2 mM EDTA is characterized by peak III.

Dual effect of DTT on the modulation of the BLM DNA-binding activity by Cd$^{2+}$ and reversibility with EDTA. DTT has been previously described either as a stimulating or a counteracting agent for Cd$^{2+}$-mediated inhibition$^{35,36}$. All the above-mentioned results were obtained under reducing conditions, i.e. in the presence of DTT. We then compared the Cd$^{2+}$ effect on the BLM$^{642–1290}$ DNA interaction under reducing and non-reducing conditions (Fig. 5). In the presence of DTT, Cd$^{2+}$ efficiently inhibited the binding of BLM$^{642–1290}$ to DNA (Fig. 5a), in accordance with results shown in Fig. 3. To note, NEM that did not lead to significant effect on its own on the BLM$^{642–1290}$ DNA-binding activity, fully counteracted the inhibitory effect of Cd$^{2+}$, suggesting that Cys of the 1st cluster (surface residues as defined above) mainly mediate the Cd$^{2+}$-dependent inhibition.

By contrast, under non-reducing conditions, Cd$^{2+}$ did not significantly inhibit the BLM$^{642–1290}$ DNA-binding activity (Fig. 5b), suggesting that Cd$^{2+}$-targeted Cys are most likely involved in intra- or inter-molecular disulfide bonds and react with Cd$^{2+}$ upon reduction only (in the form of sulfhydryl groups). The effect of Cd$^{2+}$ on the BLM$^{642–1290}$ DNA-binding activity was further studied under different reducing conditions by varying the DTT concentration. At 0.2 mM DTT (Fig. 5c, right), this effect was comparable to the one previously observed at 2 mM DTT (Fig. 5a), i.e. DTT promoted the Cd$^{2+}$-mediated inhibition of protein-DNA interaction, while no Cd$^{2+}$ effect was observed in the absence of DTT (Fig. 5c, left). Interestingly, under limited reducing conditions (0.02 mM DTT; Fig. 5c, middle), the addition of Cd$^{2+}$ led to an increase in the steady-state fluorescence anisotropy, instead of a decrease. This dual effect of DTT remains unclear. We can hypothesize that Cd$^{2+}$ differentially modulates self-assembly properties of BLM$^{642–1290}$ depending on the DTT concentration. As shown above, no free sulfhydryl of surface Cys is available for Cd$^{2+}$ binding in the absence of DTT and the presence of intra/inter-molecular disulfide bonds is compatible with DNA-binding. Under moderate DTT conditions (0.02 mM DTT), a part of Cys residues only could be in their reduced forms and Cd$^{2+}$ could promote the formation of higher-order BLM$^{642–1290}$ multimers of moderate sizes which remain compatible with DNA-binding but leading to higher steady-state anisotropy values due to the size of protein-DNA complexes (larger compared with the size of complexes obtained in the absence of DTT). By contrast, under strong reducing conditions (>0.2 mM DTT), most of surface Cys residues should be in the reduced state and thus, Cd$^{2+}$ promotes large multimers/aggregates, not competent for DNA-binding. In other words, the DNA-binding site should be completely hidden in the context of these large BLM multimers/aggregates. Note that both Cd$^{2+}$ effects (inhibition or stimulation) were abolished by addition of EDTA (Fig. 5c) in a dose-dependent manner (Supplementary Fig. S4). Consistent with our hypothesis, EDTA was shown to reverse Cd$^{2+}$-induced multimers by size-exclusion chromatography and DLS (Fig. 4).

Cd$^{2+}$-induced inhibition of BLM DNA unwinding activity was not reversed by addition of EDTA. We next addressed the question of whether EDTA could restore the unwinding activity of BLM as measured in the presence of Cd$^{2+}$ using the stopped-flow FRET assay. First, Cd$^{2+}$ significantly reduced both the unwinding kinetic rate constant and the corresponding reaction amplitude characterizing BLM$^{642–1290}$ per se since the Mg$^{2+}$ cofactor is required for this activity. As shown in Fig. 6b, although the kinetic rate constant was lower in the presence of EDTA, the reaction amplitude was only slightly affected. This result shows that BLM$^{642–1290}$ sustains a significant DNA unwinding activity even in the presence of EDTA. Nevertheless, the Cd$^{2+}$-dependent inhibition of BLM activity was not counteracted by EDTA (Fig. 6b), in contrast to that previously observed for the DNA-binding step. The fact that the helicase activity of BLM$^{642–1290}$ cannot be recovered...
Figure 5. Influence of DTT on the Cd\(^{2+}\)-mediated inhibition of DNA-BLM\(^{642-1290}\) interaction and protective effect of NEM. BLM\(^{642-1290}\)/DNA complexes were formed using 200 nM protein and 5 nM F-18 oligonucleotide in the presence (a) or absence (b) of 2 mM DTT. The concentration of CdCl\(_2\) and NEM were 50 \(\mu\)M and 0.5 mM, respectively. The fluorescence anisotropy was monitored at 25 °C after addition of F-18. The relative DNA-binding activity was calculated according to Eq. 2. (c) Differential effect of Cd\(^{2+}\) on the DNA-binding activity of BLM\(^{642-1290}\) as a function of DTT concentration and reversibility with EDTA. The time of addition for Cd\(^{2+}\)/EDTA is explicitly indicated by arrows. Protein and F-18 concentrations were 200 and 5 nM, respectively. The fluorescence anisotropy was monitored at 25 °C as a function of time. Left: no DTT; middle: 0.02 mM DTT; right: 0.2 mM DTT.

Figure 6. Cd\(^{2+}\)-induced impairment of helicase unwinding activity is irreversible. (a) Dependence of the kinetic rate constant and reaction amplitude as a function of CdCl\(_2\) concentration as measured by stopped-flow FRET assay. BLM\(^{642-1290}\) was first preincubated with varying concentrations of CdCl\(_2\) for 5 min at 25 °C. The DNA substrate (16-bp duplex with a 20-nt 3’ tail) was then added into the reaction mixture and the reaction was initiated by rapid mixing with 1 mM ATP. Insert: typical kinetics for DNA unwinding in the presence of various CdCl\(_2\) concentrations. (b) EDTA failed to restore Cd\(^{2+}\)-induced DNA unwinding. BLM\(^{642-1290}\) was first preincubated for 5 min at 25 °C in the absence or presence of 50 \(\mu\)M CdCl\(_2\). Each sample was further incubated for 5 min in the absence or presence of 2.5 mM EDTA, before addition of the DNA substrate and reaction initiation with ATP as explained in the legend of panel (a). BLM\(^{642-1290}\) alone (grey trace); BLM\(^{642-1290}\) + EDTA (red trace); BLM\(^{642-1290}\) + Cd\(^{2+}\) (blue trace); BLM\(^{642-1290}\) + Cd\(^{2+}\) + EDTA (green trace). Reactions were performed with 4 nM DNA and 60 nM protein in Tris-HCl buffer (25 mM, pH 7.5) supplemented with 50 mM NaCl, 2 mM MgCl\(_2\) and 1 mM DTT at 37 °C.
upon addition of EDTA, suggests that Cd²⁺ also inactivates (at least) one additional Cys or another residue, most likely playing a proper catalytic role for DNA unwinding, in an irreversible manner.

Discussion

The BLM helicase plays key roles in numerous cellular processes including DNA double-strand break repair (DSB), Holliday junction resolution and chromosome segregation[32–39]. In this context, the study of the toxic effect of Cd on BLM is of particular interest for understanding the pivotal role of BLM in keeping genome stability. Among Cd²⁺, Zn²⁺ and Hg²⁺, only Cd²⁺ has been reported to induce SCEs, a unique cytological feature of BLM-deficient cells[40]. Furthermore, recent studies highlight that Cd²⁺ targets major players of the DNA-repair machinery including proteins involved in the BER, NER or MMR pathways[21–23], although little is known about Cd²⁺ effects on proteins involved in the DSB pathway such as BLM. Here, we investigated the interplay between BLM and Cd²⁺ at the molecular level. We found two distinct molecular mechanisms accounting for the Cd²⁺-mediated inactivation of BLM. Cd²⁺ targets surface Cys in their reduced state and promotes the formation of large BLM multimers and then inhibition of the BLM-DNA interaction. The Cd²⁺-dependent multimerization and DNA-binding inhibition processes were found to be fully reversible upon addition of EDTA. However, the inhibition of the BLM helicase activity was irreversible suggesting another mechanism at the catalytic level, mediated by the targeting of a catalytic residue.

We found that Cd²⁺ was able to efficiently inhibit both helicase and ATPase activities of BLMfull-length and BLM642–1290 in the low micromolar concentration range. The corresponding IC₅₀ values were compatible with values derived from DNA-binding inhibition curves suggesting that the Cd²⁺-dependent inhibition of helicase and ATPase activities could be explained in part by inhibition of the DNA-binding step. The reversibility of the DNA-binding inhibition process by EDTA indicates that the Cd²⁺-dependent inhibition mechanism is not associated with an irreversible structural modification of the protein that could affect DNA-binding properties. It is important to note that, under experimental conditions where Cd²⁺ was maintained below 100 μM, fluorescence anisotropy experiments did not show any significant direct DNA-Cd²⁺ interaction, in accordance with previous studies[41]. Furthermore, we found that Cd²⁺ inhibits both BLM and RecQ<sub>E.coli</sub> helicases to different extents, with RecQ<sub>E.coli</sub> much less susceptible to Cd²⁺ (one order of magnitude) than BLM. This differential susceptibility reinforces the idea that Cd²⁺ does not directly target DNA in our activity or DNA-binding assays. Instead, this differential susceptibility appears to be related to the number of solvent-exposed Cys contained in the protein structure with RecQ<sub>E.coli</sub> having much less surface Cys (i.e. protected by NEM) than BLM[42–1290] (Fig. 2).

We also tested whether Cd²⁺ could replace Zn²⁺ in the ZBD based on several statements: (i) Cd²⁺ has been previously reported to react with thiol groups, particularly with Cys and glutathione that act as the first line of defense against Cd²⁺ in cells[36,42]. (ii) The antagonistic effect of Zn²⁺ on Cd²⁺ has long been documented and previous studies have shown that Cd²⁺-targeted sites in proteins correspond to zinc finger motifs and that the replacement of Zn²⁺ by Cd²⁺ may be reversible[43,44]. However, here, we failed to demonstrate that Zn²⁺ prevents the inhibitory effect of Cd²⁺ on BLM. Taking into account the higher affinity of Cd²⁺ over Zn²⁺ for Cys<sub>4</sub> ZBD, large excess of Zn²⁺ should be required to observe a protective effect against Cd²⁺; it was not possible to satisfy this condition since, instead to rescue BLM DNA-binding activity, Zn²⁺ on its own displayed a significant inhibitory effect although this effect was less important than Cd²⁺. If Cd²⁺ target the helicase ZBD, this is probably not the predominant effect for the following reasons: (i) BLM and RecQ<sub>E.coli</sub> are differentially affected by Cd²⁺, in accordance with their respective number of solvent-exposed Cys (Fig. 2 and Supplementary Table S2) and (ii) we have previously shown that Zn²⁺-coordination to BLM or RecQ<sub>E.coli</sub> ZBD is strictly required for correct protein folding (during production) but dispensable after for both DNA-binding and helicase activities[34]. The mechanism by which Zn²⁺ inhibits BLM remains unknown and it appears that Zn²⁺ differentially affects helicases of the RecQ family with Zn²⁺ having only a slight inhibition effect on the binding of RecQ<sub>E.coli</sub> to DNA (Supplementary Fig. S3). Consistent with our study on human BLM, it was previously shown that Zn²⁺ also significantly impairs the helicase activity of BLM yeast homologue, Sgs1[45]. It has also been shown that Zn²⁺ enhances the 3′ → 5′ exonuclease activity of another human RecQ helicase, the Werner protein, at the expense of the helicase activity[46–48].

It was previously shown that the binding of BLM and RecQ<sub>E.coli</sub> to DNA relies on similar mechanisms. Regarding their binding to dsDNA and ssDNA, both proteins involve distinct protein domains with ssDNA-binding sites located along the A1, A2, WH and HRDC domains, whereas the dsDNA-binding site is located near the ZBD[34,49]. Although different between BLM and RecQ<sub>E coli</sub>, the IC₅₀ values characterizing protein-DNA interaction inhibitions were similar between ss and dsDNA, reinforcing the idea that the Cd²⁺-dependent DNA-binding inhibition is not strictly related to the ZBD and probably occurs via a more general and common mechanism, regardless of the nature of DNA (ss or ds). Besides the targeting of ZBD, other mechanisms of Cd²⁺-mediated protein inhibitions involved in DNA-repair have been proposed. It was proposed that Cd²⁺ inhibits mismatch repair pathway by abrogating the ATPase activity of the MSH2-MSH6 complex, via a mechanism in which Cd²⁺ binds in a non-specific manner, leading to a stoichiometry of more than hundred Cd²⁺ per protein[29]. In the case of BLM, the number of target sites appears to be limited. In contrast to MSH2-MSH6 complexes that possess a high content of non-specific Cd²⁺ binding sites, we found that BLM[42–1290] is characterized by a much lower stoichiometry (11–12 Cd²⁺ per protein) and Cys sulfydryl groups are directly involved in this Cd²⁺-binding process. We have characterized two subclasses of Cys targeted by Cd²⁺, based on (i) the presence of two distinct slopes in stoichiometry plots, most likely accounting for differences in accessibility/affinity and (ii) the NEM protective effect. The first class (6–7 residues) is composed by Cys displaying high affinity for Cd²⁺, most likely located at the protein surface while the second class (5 residues) corresponds to Cys with lower affinity for Cd²⁺ and probably buried and located in the protein core.

Our data suggest that the first class of Cys is fully responsible for the Cd²⁺-dependent inhibition of BLM DNA-binding since NEM, which prevents the binding of Cd²⁺ to Cys belonging to this class only, fully rescues the DNA-binding activity in the presence of Cd²⁺ (Fig. 5). The Cd²⁺-mediated inhibition of DNA-binding was strictly
dependent on the presence of DTT suggesting the involvement of free sulphydryl groups of surface Cys in the binding of Cd\(^{2+}\). Most of the class 1 Cys should be engaged in disulfide bridges under non-reducing conditions, explaining the absence of any inhibitory effect of Cd\(^{2+}\) in the absence of DTT. In parallel, we found that Cd\(^{2+}\) promotes BLM higher-order multimers or aggregates starting from monodisperse samples of monomers. The mechanism behind at the molecular level is not yet clearly understood and still under investigation. This phenomenon was fully reversed by EDTA, in accordance with the counteracting effect of EDTA on BLM DNA-binding activity. To note, only reducing conditions (to ensure free sulphydryl groups) were compatible with the observation of Cd\(^{2+}\)-dependent inhibition, suggesting that DNA-binding sites of BLM protomers should be hidden upon the formation of these large non-specific multimers/aggregates. Interestingly, mild reducing conditions, which modulate the number of free -SH at the protein surface, led to an intermediary result between reducing condition (inhibition of DNA-binding by Cd\(^{2+}\) as measured by a decrease in the anisotropy value) and non-reducing condition (no effect of Cd\(^{2+}\) on the anisotropy value) (Fig. 5c): in this mild reducing condition, Cd\(^{2+}\) led to an increase in the anisotropy value, probably accounting for DNA-binding of organized multimers of moderate sizes. Their oligomeric status should be intermediary between monomers and large non-specific multimers/aggregates, with remaining solvent accessible DNA-binding sites. The different number of solvent-exposed Cys between BLM\(^{642-1290}\) and RecQ\(^{E.coli}\), 6–7 and 3, respectively, highlights the relationship between the number of Cys potentially targeted by Cd\(^{2+}\) and the extent of the Cd\(^{2+}\)-dependent DNA-binding inhibition. To note, these numbers of solvent-exposed Cys as determined experimentally agree well with the number of surface Cys based on 3D structures (7 and 3, respectively, without taking into account ZBD Cys; Supplementary Table S2). Nevertheless, the Cd\(^{2+}\)-dependent inhibition of the BLM unwinding activity was not reversed by EDTA, suggesting an additional catalytic mechanism of inhibition, unrelated to the DNA-binding step. According to the three-dimensional BLM structure\(^{1,32}\), we performed single point mutations of Cys residues implicated in DNA binding/unwinding (C895S and C901S). The two BLM mutants displayed modest reduced DNA-unwinding activity and their responses to Cd\(^{2+}\) were similar to the wild-type BLM (data not shown). This suggests that distinct residues (Cys or other amino-acids) or, alternatively, a combination of the two above-mentioned Cys could be responsible for the irreversible BLM inactivation by Cd\(^{2+}\). The underlying mechanism is currently under investigation.

Methods

Chemical reagents. CdCl\(_2\), ZnCl\(_2\), EDTA, DTT (dithiothreitol), NEM (N-Ethylmaleimide), ATP and Triton X-100 were purchased from Sigma. \([\gamma-32P]\) ATP was purchased from Perkin Elmer.

Recombinant proteins. RecQ\(^{E.coli}\) and BLM\(^{642-1290}\) helicases were expressed and purified as previously described\(^{54,55}\). BLM\(^{full-length}\) was expressed in Saccharomyces cerevisiae JEL-1 strain as previously described\(^{44}\) with some modifications in the purification protocol. Briefly, cells were thawed at 4 °C and cell disruption was performed using a French press in a Tris-HCl buffer (50 mM, pH 7.5) supplemented with 500 mM KCl, 10% sucrose, 1 mM DTT and protease inhibitors cocktail (Roche). After DNA fragmentation by sonication, the crude extract was subjected to centrifugation at 30,000 g for 45 min using a SS34 rotor (Sorvall). Filtered supernatant (using 0.45 μm filters) was loaded into 10 mL of Ni\(^{2+}\)-NTA agarose resin (Qiagen). Beads were washed with 100 mL of K\(_2\)HPO\(_4\)/KH\(_2\)PO\(_4\) buffer (20 mM, pH 7.4), 0.05% Triton X-100, 10% glycerol, 1 mM DTT (buffer A-20 mM) supplemented with 500 mM KCl and 20 mM imidazole, followed by 100 mL of buffer A-20 mM supplemented with 500 mM KCl and 50 mM imidazole. Proteins were then eluted with 100 mL of buffer A-20 mM supplemented with 500 mM KCl and 300 mM imidazole. Fractions containing BLM were pooled and directly loaded onto a 10-mL Biogel CHT hydroxyapatite column (Bio-Rad). The column was washed with 100 mL of Tris-HCl buffer (50 mM, pH 7.5) supplemented with 500 mM NaCl and 1 mM DTT, for 5 min at 25 °C. This condition is suitable to measure Cd\(^{2+}\)-protein -DNA complexes 32. When required, the concentration of NEM was 0.5 mM. Free Cd\(^{2+}\) in solution was measured by a fluorescence-based Measure-IT Cadmium assay (Invitrogen) according to the manufacturer’s protocol. To avoid any bias in the measurement of free Cd\(^{2+}\) due to equilibrium displacement (Cd\(^{2+}\)-protein -> Cd\(^{2+}\)-sensor), proteins were eliminated from the mixture by using Q-sepharose beads before the measurement. Free Cd\(^{2+}\) concentrations were deduced from calibration plots using CdCl\(_2\) solutions of known concentrations.

Oligonucleotides. The sequences of DNA substrates used for enzymatic or DNA-binding assays are shown in Supplementary Table S1. PAGE-purified fluorescein-labeled or unlabeled synthetic oligonucleotides were purchased from Eurogentec. Double-stranded DNA substrates were obtained by mixing equimolar amounts of complementary strands in 20 mM Tris-HCl, pH 7.5, 100 mM NaCl. The mixture was heated to 95 °C for 5 min and annealing was allowed by slow cooling to room temperature.

Cadmum binding assay. Cd\(^{2+}\) binding to helicases was assayed by incubating BLM\(^{642-1290}\) or RecQ\(^{E.coli}\) (0.5μM) with increasing concentrations of CdCl\(_2\) (from 0 to 37.5 μM) in 50 μL of Tris-HCl buffer (50 mM, pH 8.0) supplemented with 50 mM NaCl and 1 mM DTT, for 5 min at 25 °C. This condition is suitable to measure Cd:protein stoichiometries since both protein and Cd\(^{2+}\) concentrations were much higher than K\(_d\) values characterizing Cd-Cys complexes\(^{32}\). When required, the concentration of NEM was 0.5 mM. Free Cd\(^{2+}\) in solution was measured by a fluorescence-based Measure-IT Cadmium assay (Invitrogen) according to the manufacturer's protocol. To avoid any bias in the measurement of free Cd\(^{2+}\) due to equilibrium displacement (Cd\(^{2+}\)-protein -> Cd\(^{2+}\)-sensor), proteins were eliminated from the mixture by using Q-sepharose beads before the measurement. Free Cd\(^{2+}\) concentrations were deduced from calibration plots using CdCl\(_2\) solutions of known concentrations.
(we checked that interaction between free Cd and beads was negligible: fluorescence intensity values obtained after incubation of CdCl₂ solutions with beads were 95–100% of intensities measured with ”input” solutions). The number of protein-bound Cd²⁺ was estimated by subtracting the amount of free Cd²⁺ to the total amount of Cd²⁺.

Size-exclusion chromatography and dynamic light scattering (DLS) experiments. The size-exclusion chromatography experiment was performed according to Xu et al.²⁰. Briefly, the chromatography was performed at 25 °C, using an FPLC system (GE healthcare), on a Superdex 200 (analytical grade) column equilibrated with buffer S (20 mM Tris-HCl, pH 7.4, 500 mM NaCl, 1 mM DTT and 5% glycerol (v/v)) +/- 2 mM EDTA. 20 μl of untreated or Cd²⁺-treated BLM (+/- 2 mM EDTA) was loaded on the column (typically in the 2–6 μl concentration range) and was eluted with buffer S +/- 2 mM EDTA, at a flow rate of 0.4 ml/min; the absorbance was continuously monitored at 280 and 260 nm. The standard molecular markers (Sigma) used for calibration were eluted under identical experimental conditions.

DLS measurements were performed using a DynaPro NanoStar instrument (Wyatt Technology, France) equipped with a thermostated cell holding filtered (0.1 μm filters) solutions in disposable cuvettes (UVette, Eppendorf). The protein concentration was 1.5 μM in a Tris-HCl buffer (50 mM, pH 8.0, 200 mM NaCl, 1 mM DTT) (total volume, 50 μl). The scattered light was collected at an angle of 90°. Recording times were typically between 3–5 min (20–30 cycles in average of 10 s each). The analysis was performed with the Dynamics 7.0 software using regularization methods (Wyatt Technology, France). The molecular weight was calculated from the hydrodynamic radius using the following empirical equation 1:

\[ \text{Mw} = (1.68 \times R_H^{2.34}) \]

where Mw and RH represent the molecular weight (kDa) and the hydrodynamic radius (nm), respectively.

ATPase activity assay. The ATPase activity was assayed by measuring the release of free phosphate during ATP hydrolysis.³⁴ The reaction was carried out for 10 min at 37 °C in an ATPase reaction buffer (Tris-HCl 50 mM, pH 8.0) supplemented with 50 mM NaCl, 3 mM MgCl₂, 0.1 μg/ml BSA and 1 mM DTT. The reaction was initiated by the addition of helicase (200 nM) into a reaction mixture containing 3 μM DNA (25-nt ssDNA) and 2 mM [γ-³²P] ATP in the absence or presence of increasing CdCl₂ concentrations (final volume, 50 μl). The reaction was stopped by transferring 35 μl of the reaction mixture into a hydrochloric solution of ammonium molybdate. The liberated radioactive γ³²Pi was extracted with a solution of 2-butanol-benzene-acetone-ammonium molybdate (750:750:15:1) saturated with water. A volume of 400 μl was removed from the organic phase and radioactivity was quantified using a liquid scintillation counter (Beckman LS 5000CE).

Helicase assay. 1) Radioactive assay: DNA helicase reaction was carried out at 37 °C in a reaction mixture containing 25 mM Tris-HCl, pH 8.0, 50 mM NaCl, 3 mM MgCl₂, 0.1 μg/ml BSA, 1 mM DTT, 2 mM ATP. To address the Cd²⁺ effect on helicase activity, helicases were preincubated without or with Cd²⁺ for 2 min at 37 °C. The unwinding reaction was initiated by addition of 10 fmol of the 32P-labeled partial duplex DNA substrate (3000 cpm/μg) and the reaction mixture was further incubated for 20 min at 37 °C. The reaction was quenched by adding 5 μl of loading buffer containing 50 mM EDTA, 0.5% SDS, 0.1% xylene cyanol, 0.1% bromophenol blue and 50% glycerol. The reaction products were analyzed by gel electrophoresis using a 12% polyacrylamide gel.

2) Stopped-flow fluorescence measurements: A stopped-flow FRET assay was used for measuring the unwinding kinetic rate constant of BLM, using doubly labeled DNA substrates, with fluorescein and hexachlorofluorescein as a donor and acceptor, respectively.²⁰,³⁰ The set-up and kinetic data analysis were described in Liu et al.³⁰. The standard reaction was performed with 4 nM DNA substrate and 60 nM protein in 25 mM Tris-HCl, pH 7.5, 50 mM NaCl, 2 mM MgCl₂, 1 mM DTT at 37 °C.

DNA-binding assay. The influence of Cd²⁺ on the DNA-binding activity of BLM was assayed by measuring the steady-state fluorescence anisotropy parameter²⁶-⁵⁸, using a Beacon 200 polarization instrument (PanVera, Madison, Wi), equipped with a temperature-controlled cuvette, according to Xu et al.³¹. Briefly, the 3'-fluorescein-labeled DNA, free in solution and bound to BLM, are characterized by fast (low anisotropy value) and slow (high anisotropy value) rotational diffusion, respectively. The relative change in the anisotropy value allows the calculation of the fractional saturation function. Recombinant RecQ from BLMfull-length or BLM642-1290 were preincubated for 10 min at 25 °C with increasing concentrations of CdCl₂ in a Tris-HCl buffer (50 mM, pH 8.0, 50 mM NaCl, 1 mM DTT) before addition of the 3'-fluorescein labeled double- or single-stranded DNA (5 nM in a total volume of 150 μl). Fluorescence anisotropy was measured under real-time condition (steady-state fluorescence anisotropy values were recorded every 8 s). The effect of Cd²⁺ on the fractional saturation function (A) (also called relative DNA-binding activity) was calculated using equation 2:

\[ A(\%) = (A_x - A_0)/(A_y - A_0) \times 100 \]

where A₀ and A represent the fluorescence anisotropy values for a given concentration of protein in the presence or absence of CdCl₂, respectively. A₀ represents the anisotropy value characterizing the fluorescently labeled DNA alone.

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Author Contributions
N.B., E.D. and X.-G.X. conceived and designed the study. X.-G.X. and E.D. wrote the paper. W.Q., N.B. and B.Z. carried out biochemical experiments. E.H. performed DLS experiments.

Additional Information
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