A bromodomain-DNA interaction facilitates acetylation-dependent bivalent nucleosome recognition by the BET protein BRDT

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Abstract

Bromodomains are critical components of many chromatin modifying/remodeling proteins and are emerging therapeutic targets, yet how they interact with nucleosomes, rather than acetylated peptides, remains unclear. Using BRDT as a model, we characterized how the BET family of bromodomains interacts with site-specifically acetylated nucleosomes. We find that BRDT interacts with nucleosomes through its first (BD1), but not second (BD2) bromodomain, and that acetylated histone recognition by BD1 is complemented by a novel bromodomain-DNA interaction. Simultaneous DNA and histone recognition enhances BRDT's nucleosome binding affinity, specificity, and ability to localize to and compact acetylated chromatin. Conservation of DNA binding in bromodomains of BRD2, BRD3 and BRD4, indicates that bivalent nucleosome recognition is a key feature of these bromodomains and possibly others. Our results elucidate the molecular mechanism of BRDT association with nucleosomes and identify new structural features of BET bromodomains that may be targeted for therapeutic inhibition.

Introduction

The basic repeating unit of chromatin is the nucleosome, consisting of two copies each of four core histones (H2A, H2B, H3 and H4) wrapped in ~147 bp of DNA¹. This repeating unit, and the higher order structures it forms, serves to regulate DNA accessibility for tight control of all DNA-templated processes. Access is mediated by epigenetic modifications including histone posttranslational modifications (PTMs) and DNA methylation, which either directly influence chromatin structure, or recruit or repel chromatin effector proteins that harbor modification-specific DNA or histone binding modules, such as bromodomains²⁻⁴.

The bromodomain and extra-terminal (BET) family (BRD2, 3, 4 and BRDT in human) are multifunctional chromatin effector proteins, whose critical roles in transcription and chromatin biology have made them attractive therapeutic targets for a wide range of malignancies (recently reviewed in 5-7). These proteins have a domain architecture that is conserved from yeast to human, which features two N-terminal bromodomains separated by a linker of ~110 amino acids, a 'motif B' that mediates BET protein dimerization⁸, and a characteristic extra-terminal (ET) domain that acts as a protein-protein interaction module for recruiting co-factors involved in transcriptional regulation^{9,10}. Additionally, longer isoforms of BRD4 and BRDT (**Fig. 1a**) have an extended C-terminus that allows them to facilitate RNA polymerase II dependent transcription through interactions with the positive-transcriptional elongator (pTEFb) complex^{11,12}.

Bromodomains recognize acetylated lysine residues and have a highly conserved structural fold consisting of four α -helices (α_Z , α_A , α_B , α_C) forming a compact left-handed bundle. Variable loops connecting helices α_Z to α_A (ZA loop) and α_B to α_C (BC loop) shape the acetyl-lysine binding pocket, thus contributing to substrate specificity^{13,14}. Structure-based alignments and phylogenetic analysis of the 61 human bromodomains, which are found in 46 diverse proteins, divides them into eight distinct families¹⁴. The variability of the acetylated lysine pockets has recently allowed the development of inhibitors that are specific against various members of these families, particularly the BETs (recently reviewed in ¹⁵).

Bromodomains typically bind to acetylated lysine residues with relatively low affinity (micromolar) and relatively poor selectivity for single acetylated lysine residues within an isolated peptide¹⁴. However, specificity and affinity are frequently increased in the presence of

multiple modifications. For example, both bromodomains of BRDT, the testis-specific member of the BET family that is essential for spermatogenesis, show a preference for multiply-acetylated histone peptides^{16,17}. The first bromodomain of BRDT (BD1) preferentially binds histone H4 tail peptides acetylated at lysines 5 and 8 (H4K5_{ac}K8_{ac}), whilst the second bromodomain (BD2) has highest affinity for histone H3 tail peptides acetylated at lysines 18 and 23 (H3K18_{ac}K23_{ac}). In both cases, binding to individually acetylated peptides is either weaker or could not be determined¹⁶. Notably, the crystal structure of the BD1-H4K5_{ac}K8_{ac} peptide complex revealed that the single binding pocket of BD1 can simultaneously bind to both acetylated lysines of a K5_{ac}K8_{ac} peptide¹⁶.

There is increasing recognition that multivalency is a central component of chromatin biology. Chromatin binding proteins and complexes frequently use multiple low-affinity interactions to achieve affinities, specificities and dynamics that would otherwise be impossible through monovalent interactions¹⁸. These multivalent interactions can occur either through the combination of histone or DNA interactions within a single protein, or within a complex carrying several chromatin binding domains (recently reviewed in ^{19,20}). Typically these interactions have been studied using a 'divide and conquer' approach, particularly in studies of histone tail binding by chromatin readers that have predominantly used isolated histone peptides in place of nucleosomes.

To gain a more comprehensive picture of how BET bromodomains interact with chromatin, we sought to characterize how BRDT, with its tandem bromodomains, interacts with acetylated nucleosomes. Using a combination of biophysical methods including isothermal titration calorimetry (ITC) and methyl-TROSY NMR, we find that BRDT interacts with acetylated nucleosomes through BD1 only, whilst BD2 is unexpectedly unable to interact with acetylated histones within a nucleosome. We show that BRDT-BD1 binding to nucleosomes is bivalent and consists of simultaneous recognition of both acetylated histone tails and DNA. Importantly, we find that the newly identified bromodomain-DNA interaction is non-specific and facilitates recruitment of BRDT to bulk chromatin, where it can bring about chromatin compaction in response to histone hyperacetylation in cells. We find similar DNA binding in other members of the BET bromodomain family, suggesting that this bivalent mode of recruitment is not limited to BRDT. Our results are the first in-depth characterization of bromodomain association with

acetylated nucleosomes and highlight the importance of nucleosome components beyond the histone tail for bromodomain recognition.

Results

Nucleosome structure influences BRDT-bromodomain binding

The interactions of the BET bromodomains with chromatin have largely been studied by two approaches. Firstly, chromatin immunoprecipitation experiments and pull-downs have been used to correlate BET binding with the presence of particular histone modifications on chromatin *in vivo*^{14,21}. Alternatively, the bromodomains have been tested individually by *in vitro* experiments, such as peptide arrays and ITC, to investigate their binding preferences for specifically modified histone tail peptides^{14,16}. Here, we extend these studies and use BRDT to characterize the interaction of the BET bromodomains with *in vitro* reconstituted, site-specifically acetylated nucleosomes.

Based on previous characterization of the binding specificities of the two bromodomains of murine Brdt¹⁶, we utilized a histone semisynthesis approach and native chemical ligation to reconstitute four types of acetylated nucleosomes *in vitro*^{22,23}. These nucleosomes were either unmodified, acetylated on histone H4 (H4K5_{ac}K8_{ac}), acetylated on histone H3 (H3K18_{ac}K23_{ac}) or acetylated on both H3 and H4. The histone modifications did not interfere with octamer or nucleosome refolding (**Supplementary Fig. 1**) and therefore were used to investigate how nucleosome structure affects bromodomain binding to histone tails.

We tested two BRDT bromodomain constructs for their histone and nucleosome binding properties. The first construct (N-BRDT(1), residues 1-143) included the core bromodomain fold and the N-terminus of BRDT that has been shown to be essential for BRDT's ability to compact acetylated chromatin²⁴. The second construct, (BRDT(2), residues 258-383) was based on the murine Brdt-BD2 bromodomain crystal structure¹⁶.

As expected, isothermal titration calorimetry (ITC) showed that N-BRDT(1) requires acetylation of histone H4 for binding to histone peptides and nucleosomes. No binding was detected for unmodified samples (**Supplementary Fig. 2a, b**), whereas the dissociation constants between N-BRDT(1) and H4K5_{ac}K8_{ac} acetylated peptides and nucleosomes were found to be 13 μM and 2 μM, respectively (**Fig. 1b**). These data show that BRDT-BD1 interacts with nucleosomes with a >6-fold enhancement in affinity compared to acetylated H4 histone tail peptides alone, and indicates that the nucleosome structure is an important determinant of histone tail recognition.

Surprisingly, our ITC data indicated that BRDT(2) cannot interact with acetylated nucleosomes, despite interacting with the equivalently acetylated histone H3 peptides (**Fig. 1c**). This suggests that the nucleosome structure prevents BRDT(2) from binding the H3 tail.

To confirm our ITC data and further characterize how BRDT interacts with nucleosomes, we adopted the methyl-TROSY NMR methodology that facilitates the investigation of large complexes²⁵. Here, ¹³C, ¹H methyl groups of isoleucine, leucine and valine in a perdeuterated background were used as sensitive probes for monitoring interactions between the bromodomains and acetylated nucleosomes.

Consistent with our ITC data, N-BRDT(1) interacted with (H4K5_{ac}K8_{ac}) acetylated nucleosomes (**Fig. 2a**), whereas BRDT(2) showed no interaction with H3K18_{ac}K23_{ac} acetylated nucleosomes (**Supplementary Fig. 2c**). For N-BRDT(1), overlaid ¹³C-¹H methyl-TROSY spectra showed chemical shift perturbations (CSPs) in residues surrounding the acetylated lysine binding pocket when tested with both (H4K5_{ac}K8_{ac}) acetylated peptides and nucleosomes (**Fig. 2a,b**). In addition, binding of N-BRDT(1) to acetylated nucleosomes also induced specific CSPs distinct from those caused by peptide binding alone (**Fig. 2a,b**). These perturbations occurred towards the opposite end of N-BRDT(1) (relative to the histone binding pocket), and suggest that BD1 may make additional contacts with the nucleosome outside of the histone H4 tail.

BRDT-BD2 is tethered to acetylated nucleosomes by BD1

We hypothesized that BRDT-BD2 may require BRDT-BD1 for recruitment to acetylated nucleosomes and thus may only bind when tethered by BD1. Therefore, we expressed and purified a longer BRDT construct encompassing both bromodomains (BRDT(1-2)) to verify whether both BD1 and BD2 interact with acetylated nucleosomes when linked.

We first characterized this construct by size-exclusion chromatography (SEC), analytical ultracentrifugation (AUC), NMR spectroscopy and small angle X-ray scattering (SAXS) (**Supplementary Fig. 3**) and found that BRDT(1-2) is a monomeric, elongated and flexible molecule with the two bromodomains at either end of an unfolded linker (**Supplementary Fig. 3**). Comparison of overlaid ¹H, ¹⁵N HSQC spectra (**Supplementary Fig. 3c**) and ¹³C-¹H methyl-

TROSY spectra (**Supplementary Fig. 4a**) of BRDT(1-2) with individual N-BRDT(1) and BRDT(2) data showed a good correspondence between peak positions. Our experimental data therefore give no indication of significant changes in the folds of these domains when linked, nor of dimerization between them, as has been proposed for other bromodomains of the BET family^{8,26,27}. New resonances, which appeared in the BRDT(1-2) ¹H, ¹⁵N HSQC spectrum (**Supplementary Fig. 3c**), were attributed to the linker; these resonances were predominantly grouped in the middle of the spectrum and have significantly higher intensities, suggesting that the linker is unstructured. Comparable peak intensities for the resonances of the bromodomains in the linked construct, when compared to those of N-BRDT(1) and BRDT(2) alone, indicate that the two domains rotate independently of each other.

We then tested how BRDT(1-2) interacts with acetylated peptides and nucleosomes using leucine and valine ¹³C-¹H methyl-TROSY NMR. Overlaid spectra of samples containing BRDT(1-2) and acetylated H3 or H4 histone peptides show CSPs for both BD1 and BD2 (**Supplementary Fig. 4a, lower panels**). The observed CSPs occur in almost identical resonances regardless of the peptide, as would be expected for both peptides targeting the histone binding pockets. These data therefore show that both bromodomains bind to both acetylated histone peptides and highlight the lack of specificity of individual bromodomains for acetylated histone peptides alone.

In contrast to the peptide binding, we found that only BD1 was able to interact with nucleosomes uniformly acetylated on both histones H3 (K18_{ac}K23_{ac}) and H4 (K5_{ac}K8_{ac}) (**Supplementary Fig. 4b**). Resonances from the linker and BD2 did not show evidence of interaction with the nucleosomes and appear to have remained flexible in solution. This result shows that tethering of the BD2 to nucleosomes is not sufficient to induce interaction.

To confirm that the experimental conditions were conducive to seeing binding between BD2 and nucleosomes, we repeated the experiments with acetylated histones from the purified octamer used for nucleosome reconstitution. BRDT(1-2) binding was tested under the same concentrations and buffer conditions as the nucleosome experiments (**Supplementary Fig. 4b**). Here, BD2 interacted with the histones, demonstrating that BD2 is unable to bind to acetylated H3 tails when presented in the context of a nucleosome.

BRDT interacts non-specifically with DNA through BD1

Given the proximity of the histone H4 tail to the DNA that encircles the histone octamer, we investigated whether BRDT-BD1 might interact with DNA using electrophoretic mobility shift assays (EMSAs). Strikingly, N-BRDT(1) showed robust binding to the 167 bp Widom DNA that we use for reconstituting nucleosomes (15 μ M) (**Fig. 3a**), whereas BRDT(2) did not interact with the DNA (**Supplementary Fig. 5**).

To investigate the binding specificity and stoichiometry of these interactions we tested N-BRDT(1) binding to 66 and 25 bp DNA oligonucleotides (**Fig. 3b**). These oligonucleotides had an unrelated sequence to the previously tested 167 bp DNA (sequences in Methods). N-BRDT(1) interacted with each of the DNAs, showing that it interacts with nucleic acids without sequence specificity. Quantification of the EMSAs showed that N-BRDT(1) interacts with 66 bp and 25 bp DNAs with dissociation constants of 29 and 52 μM, respectively. An apparent increase in affinity as the DNA length increases is expected for a non-specific DNA-binding protein presented with an increased number of potential binding sites in longer DNA. Accordingly, the shifts seen in the EMSAs suggest that at least two N-BRDT(1) molecules can interact with 66 bp DNA, whereas only a single shift is seen for 25 bp DNA (**Fig. 3b, arrowheads**).

Our ITC data (**Fig. 1b**; **Supplementary Fig. 2a**, **b**) suggested that N-BRDT(1) required acetylation of H4 to bind to nucleosomes, however binding of N-BRDT(1) to free DNA would suggest that it should also bind non-specifically to nucleosomes, independent of acetylation. To test this, we performed EMSA experiments with 167 bp Widom DNA and unmodified or acetylated nucleosomes. We found that N-BRDT(1) did indeed show weak binding to unmodified nucleosomes in a similar manner seen for DNA. The apparent discrepancy with the ITC result suggests that - due to limitations in the achievable sample concentrations - the interaction could not be detected under our experimental conditions.

As predicted from our ITC experiments (**Fig. 1b**), N-BRDT(1) binds with a significantly higher affinity and apparent specificity to nucleosomes containing acetylated histone H4 (**Fig. 3c**). In contrast, BRDT(2) showed no interaction with unmodified or acetylated nucleosomes (**Supplementary Fig. 5**), supporting our findings from NMR and ITC experiments.

To our knowledge, these data are the first demonstration of DNA binding by a bromodomain and show that BRDT-BD1 interacts non-specifically with DNA and nucleosomes. BRDT therefore has a significantly different mode of interacting with nucleosomes than previously envisaged, and may target bulk chromatin through low affinity, non-specific DNA interactions, priming it to interact tightly and specifically following histone hyperacetylation.

BD1 interacts with DNA and H4 through distinct interfaces

To further characterize the BD1-DNA interaction we analyzed the sequences of human BRDT BD1 and BD2 (**Supplementary Fig. 6a**), the X-ray crystal structure of BD1 (2RFJ¹⁴) and a homology model of BD2 (generated using the Phyre2 web server²⁸) (**Fig. 4a**). These analyses identified a positively charged patch in BD1 but not BD2 that correlates in location with residues showing NMR CSPs specific to nucleosome binding (**Fig. 2a,b**). We speculated that this region may be responsible for interacting with DNA and may explain the difference in DNA binding between BD1 and BD2.

The positively charged patch of BD1 is centered on the first α -helix (α Z) and features three prominent lysine residues (K37, K41 and K45) (**Fig. 4a**). We mutated these three lysines to serine both individually and in combination to test their effect on DNA binding.

We recorded 1D NMR spectra to ensure that the mutations had not affected the folding of the bromodomains and additionally tested each of the mutated proteins by ITC for binding to $H4K5_{ac}K8_{ac}$ peptides (**Supplementary Fig. 6b,c**). Neither mutation of the central lysine, nor triple mutation significantly altered the 1D NMR spectra or the acetylated histone H4 peptide binding affinity in ITC. This is in contrast to a previously characterized point mutation in the histone binding pocket (I155Y) that, despite maintaining the bromodomain fold, significantly reduced the binding affinity of N-BRDT(1) for acetylated histone peptides from 13 μ M to >600 μ M (**Supplementary Fig. 6b,c**)

Interestingly, individual mutation of each lysine reduced N-BRDT(1) binding to DNA, with the most pronounced effect seen by mutating the central lysine (K41S) (**Fig. 4b**). Double mutation of

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lysine residues 37 and 41 (2KS), or triple mutation of all three (3KS) abolished DNA binding (**Fig. 4b**).

The bromodomain is a small domain and given the spatial proximity between the histone binding pocket and DNA binding interface, we speculated that DNA binding may be influenced by histone tail binding. We performed quantitative EMSA titrations comparing N-BRDT(1) binding to 66 bp DNA +/- H4K5_{ac}K8_{ac} peptides. Peptide binding did not significantly affect N-BRDT(1)'s affinity for 66 bp DNA (33 and 29 μM, respectively; **Supplementary Fig. 7**), showing that the two binding sites function independently. Therefore, the increased binding affinity between BRDT-BD1 and nucleosomes, when compared to DNA or histone peptides alone, appears to occur largely through the entropic benefits of bivalency¹⁸, rather than through allostery or the direct effect of a composite binding interface, such as seen for the chromodomain of Chp1²⁹.

Next, we tested the affinity of our N-BRDT(1) DNA binding mutants for $H4K5_{ac}K8_{ac}$ nucleosomes, to quantify the role of DNA binding in nucleosome recognition (**Fig. 4c**). The single point mutation of the central lysine (K41S) reduced nucleosome binding affinity ~3.5-fold (7 μ M), whilst triple mutation of all three lysines further reduced the affinity to 9 μ M; approaching the affinity of N-BRDT(1) for $H4K5_{ac}K8_{ac}$ peptides alone (~13 μ M) (**Fig. 4c**). These results show that the BD1-DNA interaction contributes to nucleosome binding affinity. BRDT-BD1 thus interacts bivalently with nucleosomes via a combination of acetylation-specific histone recognition and non-specific DNA binding.

Nucleosome structure influences bromodomain specificity

We have demonstrated that BRDT-BD1 interacts with both acetylated histone tails and DNA simultaneously to augment its nucleosome binding affinity; however, we also wanted to address the issue of whether nucleosome structure contributes to binding specificity. Bromodomains are promiscuous for binding to acetylated lysine residues within a variety of sequence contexts ¹⁴. Although multiple acetylations within a single peptide enhance both the specificity and affinity of BET bromodomain binding, some (particularly BRD4-BD2) are still able to bind to a large variety of multiply acetylated histone peptides¹⁴.

To investigate whether nucleosome structure contributes to target specificity, we produced chimeric nucleosomes in which the acetylated histone H4 tail ($H4K5_{ac}K8_{ac}$) was ligated to the core of histone H3. This allowed us to test the binding affinity of N-BRDT(1) to nucleosomes containing the optimal acetylated histone H4 recognition motif ($H4K5_{ac}K8_{ac}$), but in an alternative position on the nucleosome.

Our ITC data indicate that N-BRDT(1) interacts with chimeric nucleosomes with a dissociation constant of 5 µM (**Supplementary Fig. 8**). Notably, this represents an ~2.5-fold enhancement in binding affinity over the interaction between N-BRDT(1) and the H4K5_{ac}K8_{ac} peptide, but an ~2.5-fold decrease in affinity compared to H4K5_{ac}K8_{ac} nucleosomes (**Fig. 1b**). We propose that DNA binding localizes the bromodomain to the chimeric nucleosome in the same manner as for the wild type (WT) nucleosome, thus enhancing the local concentration and affinity. However, in the case of the WT nucleosome, the position of the acetylated histone tail relative to the acetyllysine binding pocket (orientated by the DNA), is more favorable compared to the same acetylated sequence in the chimeric location. Therefore the affinity for the WT nucleosome is higher and an additional layer of specificity is generated.

This result shows that the peptide sequence surrounding acetylated lysines is not the only determinant of bromodomain specificity. Rather, the nucleosome structure and, specifically, the position of the acetylated lysines relative to DNA also influence bromodomain recognition. This demonstrates that the specificity of a chromatin reader can only accurately be evaluated with nucleosomes and/or chromatin and cannot be reliably judged at the level of peptides.

Other BET bromodomains interact with DNA

To investigate whether DNA binding is a general feature of the BET bromodomains or is specific to BRDT-BD1, we inspected the electrostatic surface charges of the human BET bromodomains and found that each member (with the exception of BRDT-BD2) also contains a positively charged patch (**Fig. 5a**). Although the residues involved are not identical to those in BRDT-BD1, the positional conservation of this patch suggests that this region may also interact with DNA. We therefore tested the other bromodomains for DNA binding by EMSA (**Fig. 5b**). Interestingly, the first bromodomain of BRD2 and the second bromodomains of BRD2, 3 and 4

all interacted with DNA, demonstrating that bromodomain-mediated DNA binding is a conserved feature amongst all members of the BET family.

Surprisingly however, the first bromodomains of BRD3 and BRD4 did not interact with DNA, despite the presence of the positively charged face. This result shows that inspection of the electrostatic surface potentials alone is not adequate for predicting an interaction between the bromodomains and DNA. The positive charge of this surface is therefore necessary but not sufficient for DNA interaction and specific residues and/or conformation must also be important.

Given the difference we find in the DNA binding ability of BRD4-BD1 and BRDT-BD1, we wondered if this was reflected in their nucleosome binding affinities. We therefore compared BRD4-BD1 binding to acetylated nucleosomes and histone peptides by ITC (**Fig. 5c**). Interestingly, in contrast to BRDT-BD1, BRD4-BD1 does not bind acetylated nucleosomes with higher affinity than acetylated histone peptides. This supports our hypothesis that the BRDT-BD1-nucleosome interaction is bivalent and is enhanced by simultaneous interaction with acetylated histone H4 and DNA. BRD4-BD1 does not interact with DNA, thus, the binding affinities for histone peptides and nucleosomes are the same.

BD1-DNA binding is important for chromatin compaction

Histone binding by BRDT-BD1 is essential for BRDT's ability to compact chromatin in an acetylation dependent manner (^{16,24}). To investigate whether BD1-mediated DNA binding has a similar functional significance, we assayed WT and mutant BRDT constructs for their ability to localize to, and compact, hyperacetylated chromatin in cells. Based on equivalent studies on murine Brdt^{16,24} we cloned a GFP-tagged human BRDT construct (aas 1-444; ΔC-sBRDT) that is predicted to be highly active in chromatin compaction. We then made triple lysine to glutamate (3KE) mutations that abolished N-BRDT(1) DNA binding (**Fig. 6a**), without interfering with acetylated histone peptide recognition (**Fig. 6b**). ITC data indicate that N-BRDT(1)-3KE has a 5-fold reduced affinity for H4K5_{ac}K8_{ac} acetylated nucleosomes when compared to WT N-BRDT(1) (**Fig. 6b**).

Fluorescence recovery after photobleaching (FRAP) experiments show that WT human and murine ΔC -sBRDT constructs have similar fluorescence recovery half-lives ($T_{1/2} = 5$ -7 seconds) following trichostatin A (TSA) induced histone hyperacetylation. This suggests that the human protein interacts with acetylated chromatin in a similar manner to the murine protein (**Fig. 6c**). In contrast, human and murine ΔC -sBRDT proteins with mutations in their histone binding pockets (I115Y and P50A/F51A/V55A (PFV1)^{16,24}, respectively), and a DNA-binding deficient human protein (ΔC -sBRDT-3KE) showed a significantly faster recovery half-life ($T_{1/2} = \sim 1.5$ seconds); indicative of reduced association with acetylated chromatin (**Fig. 6c**). Double mutation of both the histone and DNA binding interfaces of human ΔC -sBRDT (3KE/I115Y) did not further decrease the fluorescence recovery half-life. Mutation of either site alone is therefore sufficient to prevent acetylated chromatin association in cells (**Fig. 6c**). Control FRAP experiments with and without TSA confirm that human ΔC -sBRDT 3KE, I115Y and double mutants are similarly unresponsive to TSA-induced histone hyperacetylation (**Supplementary Fig. 9**).

In accordance with the findings above, ΔC -sBRDT-3KE, I115Y and the double mutant are also compromised in their ability to compact TSA-induced hyperacetylated chromatin in cells, when compared to the WT protein (**Fig. 6d**). Instead, ΔC -sBRDT with the 3KE and/or I115Y mutations remains diffuse in the nucleus following TSA treatment. These results demonstrate that BRDT(1)-mediated bivalent recognition of nucleosomes – both through DNA and acetylated histone tails - is critical for BRDT's ability to associate with, and remodel, acetylated chromatin. Furthermore, the data support our finding that BRDT-BD2 cannot associate with nucleosomes, by demonstrating that BD2 in unable to recruit BRDT to hyperacetylated chromatin in the absence of fully functional BD1.

Discussion

The BETs are a highly conserved protein family involved in diverse functions including transcriptional regulation and chromatin remodeling. Through their bromodomains, they are known to interact with acetylated histone tails and have a rare ability to stay associated with chromatin throughout the cell cycle³⁰⁻³². Here, we have used BRDT as a model for studying the interactions of the BET bromodomains with acetylated nucleosomes and find that DNA plays a critical role.

BRDT-BD1 interacts with doubly-acetylated histone H4 tails with a low micromolar affinity and associates with chromatin *in vivo* in an acetylation-dependent manner^{16,24}. We find that BRDT-BD1 also interacts with DNA and this interaction stimulates the binding of the bromodomain to acetylated nucleosomes *in vitro* and acetylated chromatin in cells. Manual alignment of a BRDT-BD1-H4K5_{ac}K8_{ac} peptide crystal structure alongside the structure of the nucleosome (1AOI¹) suggests how binding affinity and specificity may be enhanced (**Supplementary Fig. 10**). Orientation of BD1 with its DNA-binding interface towards the DNA positions the acetyl-lysine binding pocket in the correct orientation for interaction with the histone H4 tail. The distance of ~20 Å between H4R17 leaving the surface of the nucleosome and the pre-aligned BD1 acetyl-lysine pocket would be bridgeable for the extended histone tail, allowing BD1 to recognize H4K5_{ac}K8_{ac}.

Under this scenario, the non-specific interaction with DNA would serve to localize BD1 to nucleosomes and would allow BRDT to scan chromatin for acetylated histone H4. Stable BD1 binding would then be dependent on recognition of the appropriate acetylated lysine residues. BRDT would thus be stabilized on nucleosomes in a bivalent manner, similar to that of the PWWP domain of LEDGF, which simultaneously binds H3K36me3 and DNA to enhance its nucleosome binding affinity^{33,34}.

A corollary of this model is that the relative position/spacing of acetylated lysines - relative to the nucleosome core - would influence bromodomain target specificity. We were able to demonstrate that this is the case by showing that BRDT-BD1 binds to WT and H3-chimeric nucleosomes containing acetyl-lysine residues in identical sequence contexts, but different

positions, with differing affinities. Thus, the nucleosome structure imposes an added layer of specificity to bromodomain binding.

In stark contrast to BRDT-BD1, and predictions based on peptide binding studies, we find that BD2 neither interacts with DNA nor with acetylated nucleosomes. Whereas conservation of the contiguous positively charged patch on the rest of the BET bromodomains implies conservation of function, BRDT-BD2 has a considerably more negative electrostatic surface potential (**Fig. 5b**). This may actively repel BRDT-BD2 from DNA, thus preventing an interaction with acetylated histone H3 tails in the context of the nucleosome, even when BRDT-BD2 is physically tethered to the nucleosomes by BRDT-BD1 binding to acetylated H4.

Although nucleosome binding can be enhanced by tandem domains bivalently binding nucleosomes (2-3 fold enhancement for bromo-PHD of BPTF³⁵; 3-11 fold enhancement for two PHD fingers of CHD4³⁶), this is not the case for BRDT. Unlike the structured spacer element that connects the PHD and bromodomain of BPTF³⁵, or the relatively short linker between the PHD fingers of CHD4³⁶, we find that the linker connecting the bromodomains of BRDT is long (~110 aa) and disordered (**Supplementary Fig. 3**). Furthermore, it remains so upon BRDT binding to nucleosomes (**Supplementary Fig. 4b**). With this long flexible linker the two domains are effectively independent in solution, precluding any binding enhancement through 'prepaying' entropic costs of positioning the domains for binding¹⁸. Despite this, tethering by would still increase the relative local concentration and would therefore be expected to increase the likelihood of an interaction, if it were not inhibited. This has been shown to be the case for BRD4, where a construct encompassing BRD4-BD1 and BD2 was found to bind to nucleosomes acetylated on both histones H3 and H4 with a 2.6-fold increased affinity over BRD4-BD1 alone³⁷.

Previous research has shown that BRDT localization and its ability to compact chromatin in somatic cells are primarily mediated by BRDT-BD1, whilst BRDT-BD2 is, at least in part, dispensable 16,24. Our results give a molecular explanation to the observed biological data and indicate that BRDT-BD2 may have an alternate biological function. One possible function, which would depend on flexible tethering of BD2 to nucleosomes by BD1, would be the recruitment of an acetylated non-histone protein to chromatin. BRDT would not be unique in this

function; other BET proteins are also known to use their bromodomains for interactions with acetylated non-histone proteins ³⁸⁻⁴².

Possible candidates for BRDT-BD2 recruitment would be the transition proteins (TPs) and protamines (Prms); both of which are known to be acetylated⁴³⁻⁴⁵, and both of which are critical for BRDT-mediated post-meiotic genome repackaging during spermatogenesis¹². TPs and Prms depend on BRDT for nuclear localization in elongating spermatids, and loss of BD1 leads to their accumulation in the cytoplasm, preventing histone replacement¹². Although BRDT has a nuclear localization sequence, this data indicates that chromatin binding by BRDT-BD1 is required for nuclear retention of BRDT, TPs and Prms and therefore that BRDT interacts, either directly, or indirectly, with TPs and Prms – potentially via BRDT-BD2.

In contrast to BRDT-BD1 and BD2, BRD4-BD1 has unambiguously been shown to bind to histone peptides^{14,31}, mononucleosomes³⁷ and chromatin³¹, but also to other non-histone acetylated proteins (reviewed in 5). We find that BRD4-BD1 does not interact with DNA, which led us to compare BRD4-BD1 and BRDT-BD1 binding to acetylated nucleosomes by ITC. Although BRDT-BD1 and BRD4-BD1 have similar binding affinities for H4K5_{ac}K8_{ac} peptides, their binding affinities for nucleosomes differed significantly (~6-fold). This is largely because BRD4-BD1 does not show any enhancement of binding affinity for nucleosomes over isolated peptides.

Based on our data, we propose three models for how BET bromodomains interact with nucleosomes (**Fig. 7**). BRDT-BD1 has a non-specific DNA binding activity which allows positioning of the bromodomain on the nucleosome for interaction with acetylated histone H4 tails. This bivalent interaction thus enhances the binding affinity and specificity of BRDT-BD1 for nucleosomes (**Fig. 7a**). BRDT-BD2 does not interact with DNA or nucleosomes (**Fig. 7a**), likely due to electrostatic repulsion from the DNA preventing BRDT-BD2 from accessing the histone H3 tail. Thus, it cannot enhance the binding of BRDT to nucleosomes. Finally, BRD4-BD1 retains a positively charged interface on a polarized surface but does not interact with DNA with an affinity measurable in our experiments. This domain interacts with H4K5_{ac}K8_{ac}, however shows no enhancement of binding stimulated by the nucleosome context. Unlike BRDT however, BRD4 binding to nucleosomes is enhanced by simultaneous binding of both of its bromodomains³⁷ (**Fig. 7b**).

Like BRD4-BD1, we show that BRD3-BD1 does not interact with DNA, whilst BRD2-BD1, BRD2-BD2, BRD3-BD2 and BRD4-BD2 do. In future work it will be interesting to assess the importance of these findings in nucleosome binding, transcriptional regulation and chromatin remodeling by the BET proteins *in vivo*. Furthermore, the BET bromodomains are just 8 of 61 human bromodomains. Our work provides the first description of DNA binding by a member of the bromodomain family; it will be intriguing to see whether bivalent binding of DNA and acetylated histone tails by bromodomains is a conserved feature of the wider bromodomain family of chromatin 'reader' modules.

In summary, we have used BRDT as a model to study the binding of BET bromodomains to sitespecifically acetylated nucleosomes. ITC experiments revealed that BRDT-BD1 has an enhanced affinity for acetylated nucleosomes over acetylated histone peptides, whereas BRDT-BD2 does not interact with acetylated nucleosomes, in contrast to predictions based on peptide binding studies. Using a range of biophysical methods and mutational analyses, both *in vitro* and in cells, we show that BRDT-BD1 bivalently interacts with acetylated nucleosomes and chromatin through concomitant interaction with histone H4 and DNA, facilitating BRDT-mediated acetylation-dependent chromatin compaction. We show that bromodomain-mediated DNA binding is conserved amongst members of the BET family, indicating that bivalent nucleosome recognition through simultaneous DNA and histone tail binding is an important component of nucleosome recognition by BET bromodomains and possibly in bromodomains beyond the BET family. Our results emphasize the importance of studying chromatin reader's interactions with nucleosomes rather than isolated peptides or DNA. Accordingly, this study provides important insight into the molecular mechanism of BET association with chromatin and shows that features outside of the bromodomains' histone binding pockets are crucial for interactions with nucleosomes. Knowledge of these features may pave the way for enhanced targeting of specific BET bromodomains for therapeutic purposes.

Methods

Expression and Purification of human BET Constructs

BRDT constructs N-BRDT(1) (aas 1-143), BRDT(2) (aas 258-383) and BRDT(1-2) (aas 1-383) were cloned using restriction-free cloning⁴⁶ as N-terminally His-TEV-tagged fusion proteins into pETM11 expression vectors. Original sequences were amplified from cDNA clones provided by Sino Biological Inc (catalogue number: HG11602-M).

BET bromodomains BRD2(2) (aas 348-455), BRD3(1) (aas 24-144), BRD3(2) (aas 306-416), BRD4(1) (aas 44-168) and BRD4(2) (aas 333-460), were gifts from Nicola Burgess-Brown provided through Addgene (plasmid numbers: 53626, 38940, 38941, 38942 and 38943, respectively). Codon-optimized BRD2(1) (aas 48-184; Eurofins) was cloned using restriction-free cloning⁴⁶ as an N-terminally His-TEV-tagged fusion protein into the pETM11 expression vector.

Plasmids were transformed into the *Escherichia coli* strain BL21-CodonPlus (DE3)-RIL (Stratagene), and the bacteria were grown in Luria-Bertrani (LB) medium supplemented with 50 μg ml⁻¹ kanamycin and 25 μg ml⁻¹ chloramphenicol at 37°C. Expression of recombinant proteins was induced by addition of 0.4 mM IPTG at an OD₆₀₀ of 0.6 followed by overnight incubation at 18 °C. The bacterial cell pellets were resuspended in lysis buffer (50 mM Tris [pH 8.0], 500 mM NaCl, Complete EDTA-free Protease Inhibitor Cocktail (Roche), DNase1 (Roche), 0.25 mM DTT) before being lysed by sonication and clarified by centrifugation at 25,000 r.p.m. Proteins were purified from the soluble fraction using nickel-NTA agarose (Qiagen), followed by TEV protease cleavage of the N-terminal His-tag and additional purification over Ni-NTA resin to remove the tags and His-tagged TEV protease. Finally, proteins were further purified by size exclusion chromatography using Superdex 75 or 200 columns (GE Healthcare) as appropriate. The purity and structural integrity of the purified proteins was monitored by SDS-PAGE and ¹H NMR (Supplementary Fig. 11).

NMR Labelled BRDT Constructs

For NMR experiments, the deuterated proteins were obtained by expression in M9 media containing 100% D₂O; specific ¹³C, ¹H labelling of ILV-methyl groups was achieved following

the protocol developed by the Kay laboratory ⁴⁷. Proteins were purified as described for the unlabeled equivalents.

Nucleosome Preparation

Full-length, unmodified histones

Histones H2A/H2B - Codon-optimized (supplied by Entelechon – now Eurofins), full-length human histones H2A/H2B were co-expressed from a pCDF-DUET vector transformed into the Escherichia coli strain BL21-CodonPlus (DE3)-RIL (Stratagene). Bacteria were grown in Luria-Bertrani (LB) medium supplemented with 100 µg ml⁻¹ streptomycin and 25 µg ml⁻¹ chloramphenicol at 37° C and induced at $OD_{600} = 0.6$ with IPTG (0.25 mM). Cells were harvested after 4 h by centrifugation. Cell pellets were resuspended in cold lysis buffer (20 mM Tris [pH 8.0], 100 mM NaCl, 0.1 mM EDTA, 10 mM β-mercaptoethanol and Complete EDTA-free Protease Inhibitor Cocktail (Roche)) and the cells lysed using an Emulsiflex-C3 homogenizer (Avestin) and clarified by centrifugation at 25,000 r.p.m. Clarified cell lysate was filtered using a 0.45 µm syringe filter (Merck Millipore) and injected onto combined 5 ml HiTrap Q HP and HiTrap Heparin HP columns (GE Healthcare) pre-equilibrated in lysis buffer. Columns were washed with one column volume of lysis buffer prior to detaching the HiTrap Q HP column. The HiTrap Heparin HP column was subsequently washed with 20% elution buffer (20 mM Tris [pH 8.0], 2 M NaCl, 0.1 mM EDTA, 10 mM β-mercaptoethanol) before eluting the H2A/H2B complex with a 20 CV gradient into 100% elution buffer. The pooled and concentrated fractions containing the H2A/H2B dimer were further purified by size exclusion chromatography over a Superdex 75 column (GE Healthcare). Protein purity was monitored by SDS-PAGE. The H2A/H2B complex was extensively dialysed into 1 mM DTT prior to lyophilisation and storage at -80°C.

Histones H3 and H4 - Codon-optimized (supplied by Entelechon – now Eurofins), full-length human histone H3 and histone H4 (kindly provided by T.Bartke) were individually expressed from pETM-13 and pETM-21b(+) vectors transformed into the *Escherichia coli* strain Rosetta (DE3) pLysS (Novagen). Bacteria were grown in Luria-Bertrani (LB) medium supplemented with 50 μg ml⁻¹ kanamycin (H3) or ampicillin (H4) and 25 μg ml⁻¹ chloramphenicol at 37°C and

induced at $OD_{600} = 0.6$ with IPTG (0.25 mM). Cells were harvested after 4 h by centrifugation. Histones were purified essentially as previously described ⁴⁸.

Truncated histones H3 and H4 for native chemical ligation

Codon-optimized human histones H3 and H4 (supplied by Entelechon – now Eurofins) were used for cloning H3(A25C) and H4(A15C) mutants into a pETM13 vector using restriction free cloning 46 . These constructs contain an N-terminal methionine initiation codon immediately followed by the alanine-to-cysteine mutation, thus lack residues 2-24 and 2-14, respectively. The *E. coli* methionyl-aminopeptidase removes the N-terminal methionine thus exposing the cysteine at the N-terminus. Plasmids were transformed into the *Escherichia coli* strain Rosetta (DE3) pLysS (Novagen), and bacteria were grown in Luria-Bertrani (LB) medium supplemented with 50 μ g ml $^{-1}$ kanamycin and 25 μ g ml $^{-1}$ chloramphenicol at 37°C. Cells were induced at OD $_{600}$ = 0.6 with IPTG (0.25 mM) and harvested after 4 h by centrifugation. The truncated histones were purified essentially as previously described 48 but in the absence of reducing agents. Purified histones were dialysed into water and lyophilized.

Native Chemical Ligation (NCL)

For ligations, truncated histones H3(A25C) and H4(A15C) were incubated with acetylated histone peptides carrying a C-terminal thioester in ligation buffer (200 mM NaPO₄ [pH 7.5], 0.5 mM TCEP, 6 M guanidine HCl, 100 mM sodium 2-mercaptoethanesulfonate) for 24 h at 25°C. The reactions were stopped by adding DTT to a final concentration of 100 mM.

Ligated H3 was diluted in SAU-0 (7 M urea, 20 mM sodium acetate [pH 5.2], 1 mM EDTA, 5 mM β -mercaptoethanol) to dilute the guanidine HCl concentration to < 200 mM and the protein was loaded on to a HiTrap SP HP column (GE Healthcare) pre-equilibrated in SAU-200 (200 mM NaCl). A 5 CV gradient to 25% SAU-600 (600 mM NaCl) followed by 10 CV at 25% eluted the vast majority of unligated H3. Ligated H3 was eluted with a 5CV gradient to 50% SAU-600 followed by a step to 100% SAU-600 to remove any remaining protein. Protein purity was checked by 18% SDS-PAGE and impure fractions were pooled for a second round of purification.

Truncated H4(A15C) and H3(A25C)(for chimeric nucleosomes) were ligated to a peptide consisting of a His-tag, TEV-cleavage site and the H4 N-terminal sequence (1-14) and therefore could be purified using Ni-NTA agarose (Qiagen). Ligated H4 was extensively dialysed into binding buffer (100 mM NaPO₄, 10 mM Tris, 6 M guanidine HCl, pH adjusted to 8.0) and then incubated with pre-equilibrated Ni-NTA resin. Ligated H4 was eluted with elution buffer (100 mM NaPO₄, 10 mM Tris, 8 M urea, pH adjusted to 4.5) and dialysed into 1 mM DTT prior to lyophilisation. Protein purity was checked by 18% SDS-PAGE.

167 bp Widom DNA

A plasmid containing 80 repeats of 167 bp with a centered Widom 601 sequence⁴⁹, was amplified in XL1 Blue cells (Stratagene). Plasmid DNA was purified by Gigaprep (Qiagen) using the standard protocol. Purified plasmid was digested using AvaI (NEB) to isolate 167 bp repeats, which were purified from the vector backbone by size exclusion chromatography with an XK 16/70 Superose 6 pg (GE Healthcare) gel filtration column. Purified DNA was precipitated by addition of 0.7 volumes of isopropanol and 0.3 M sodium acetate and pelleted by centrifugation at 20,000 g for 1 hour at 4°C. DNA was resuspended in TE buffer (10 mM Tris [pH 8.0], 1 mM EDTA) and the purity checked on a 2% agarose gel.

Nucleosome Reconstitution

Histone octamers were refolded from purified histones and assembled into nucleosomes with 167 bp DNA by salt deposition as previously described ⁴⁸. Chimeric nucleosomes contained chimeric H3-H4K5_{ac}K8_{ac} and wild type H2A, H2B and H4, and thus contained both modified and unmodified H4 tail sequences. Reconstitution conditions were optimized by titration and nucleosomes checked by 5% native PAGE. N-terminal His-tags on ligated H4 of H4K5K8ac nucleosomes and H3-H4K5_{ac}K8_{ac} chimeric nucleosomes were removed by TEV cleavage (2 hours, 30°C) of reconstituted nucleosomes (**Supplementary Fig. 1**). Cleaved nucleosomes were purified by incubation with Ni-NTA agarose and collection of flow-through.

Isothermal Titration Calorimetry (ITC)

ITC was carried out at 20°C with an ITC 200 Microcalorimeter (GE Healthcare) following dialysis of purified BRDT proteins and nucleosomes into interaction buffer (20 mM Tris [pH

8.0], 150 mM NaCl, 1 mM EDTA and 1 mM TCEP). For BRDT and BRD4 BD1 peptide binding experiments, histone peptides (450 μM) were titrated into BD1 proteins (25-30 μM). For BRDT-BD2 peptide binding experiments, histone peptides (2.5 mM) were titrated into BRDT-BD2 proteins (90 μM). For nucleosome binding experiments, BET bromodomain proteins (200-260 μM) were titrated into nucleosomes (6-11 μM). Peptides were supplied by Peptide Protein Research Ltd and were solubilized in interaction buffer. ITC data was analyzed with using the MicroCal OriginTM software package after correction for heats of dilution. A summary of all ITC presented in this paper can be found in **Supplementary Table 1**.

Nuclear Magnetic Resonanace (NMR)

NMR experiments were performed on Bruker Avance III 600 and 800 MHz spectrometers equipped with HCN triple-resonance cryo-probes. Protein assignments were obtained using a combination of standard triple resonance experiments⁵⁰. Nucleosome interaction was monitored by $^{13}\text{C-}^{1}\text{H}$ HMQC experiments recorded for 20 h on labeled BRDT constructs titrated into acetylated histones or nucleosomes (10 μ M) at 34°C. NMR data were processed by NMRPipe⁵¹ and analyzed with NMRView⁵².

Electrophoretic Mobility Shift Assays (EMSAs)

For the binding reaction, a master mix containing DNA and reaction buffer (20 mM Tris [pH 7.5], 100 mM NaCl, 5 mM DTT, 0.5 mM EDTA) was prepared and mixed with dilutions of BRDT constructs to the final concentrations indicated. Binding was performed for 30 min at 4°C. BRDT-DNA complexes were resolved by PAGE (5%-7% polyacrylamide, 3% glycerol, and 0.5× TAE) at 100 V for ~ 1.5 h (4°C). Experiments utilizing 167 bp DNA were stained with ethidium bromide and imaged using the AlphaImager HP imaging system (Proteinsimple). EMSA gels using ³²P-labeled 25 and 66 bp dsDNA oligos were dried and exposed to a phosphorimager screen. Unbound DNA was quantified as a proportion of total signal/lane using a Typhoon imager with ImageQuant software and data plotted against protein concentration to calculate the binding affinities indicated.

DNA sequences used for EMSA binding studies:

167bp Widom DNA:

Forward -

tegggggegcectggagaateceggtgeegaggegcteaattggtegtagacagetetageacegettaaacgeacgtaege getgteecegggttttaacegeeaaggggattaetecetagteteeaggeacgtgteagatatatacatectgtgeatgtae

Reverse -

ccgagtacatgcacaggatgtatatatctgacacgtgcctggagactagggagtaatccccttggcggttaaaacgcgggggacagc gcgtacgtgcgtttaagcggtgctagagctgtctacgaccaattgagcggcctcggcaccgggattctccagggcggccc

66 bp DNA:

cgatatagtgtaacggctatcacatcacgctttcaccgtggagaccggggttcgactccccgtatc

25 bp DNA:

cgaagtggccgagtggtctatggcg

Fluorescence Recovery after Photobleaching (FRAP)

FRAP experiments were performed as previously described¹⁶. In brief, 1.5 μg of each of the GFP-BRDT constructs was transfected into Cos7 cells using lipofectamine 2000. Cells were treated with TSA (50 ng ml⁻¹) and incubated at 37 °C in 5% CO2 for 16 h. FRAP analysis was performed using a Zeiss microscope (LSM710 NLO-LIVE7-Confocor3) equipped with a 488 nm laser and a LP505 filter, on ten independent cells. A circular region was bleached for 1.22 s; fluorescence and t1/2 recovery were measured with the software Zen. Cos7 cells (ATCC[®] CRL1651TM) were authenticated and supplied by the American Type Culture Collection (ATCC). This cell line has been tested for mycoplasma contamination with the MycoAlert Mycoplasma Detection Kit (Lonza).

Data availability

All relevant data reported in this paper are available from the authors upon request.

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Author Contributions

T.C.R.M. and H.G. performed biochemical purifications; T.C.R.M performed EMSA

experiments; B.S. carried out NMR experiments; B.S., T.C.R.M. and T.C. analyzed NMR data;

V.R. carried out and analyzed ITC experiments; S.K. and S.C. designed and performed FRAP

experiments. C.W.M. and T.C. conceived the study. T.C.R.M wrote the manuscript with input

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from all authors.

Competing financial interests

The authors declare no competing financial interests.

Figures

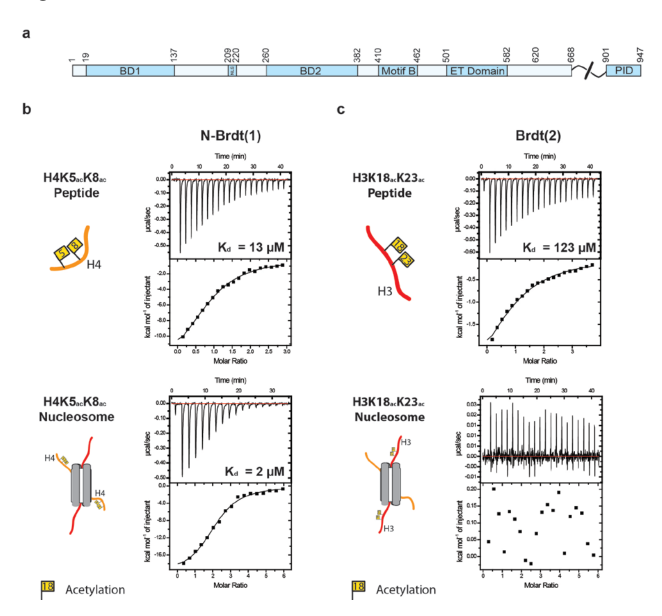


Figure 1. The nucleosome structure augments acetylated histone tail binding by BRDT-BD1 but prevents interaction with BRDT-BD2 (a) Domain layout of human BRDT (b) ITC profiles for N-BRDT(1) interactions with either acetylated histone H4 tail peptides or acetylated nucleosomes (both H4K5_{ac}K8_{ac}), as indicated. (c) ITC profiles for BRDT(2) interactions with either acetylated histone H3 tail peptides or acetylated nucleosomes (both H3K18_{ac}K23_{ac}), as indicated.

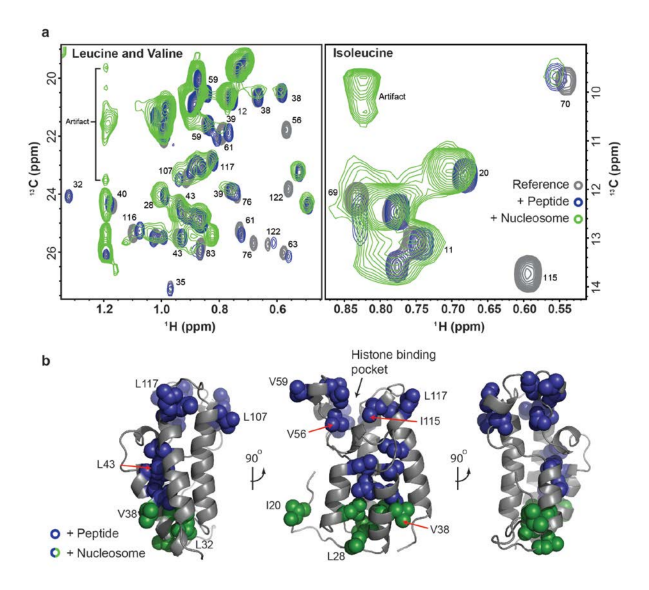


Figure 2. BRDT interacts bivalently with nucleosomes through BD1 (**a**) Overlaid ¹³C-¹H methyl-TROSY spectra of isoleucine, leucine and valine labeled N-BRDT(1) alone or in the presence of H4K5_{ac}K8_{ac} peptides or H4K5_{ac}K8_{ac} nucleosomes. (**b**) Residues showing chemical shift perturbations (CSPs) in (**a**) upon binding of nucleosomes (blue and green) or peptides only (blue) displayed on a homology model of human BRDT-BD1 (generated using the Phyre2 web server²⁸).

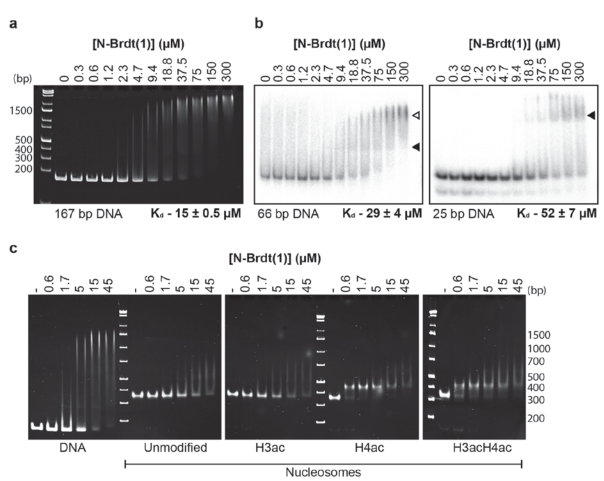


Figure 3. BRDT interacts non-specifically with DNA through BD1. (a) Electrophoretic Mobility Shift Assay (EMSA) of a titration of N-BRDT(1) incubated with 167 bp double stranded DNA containing a centralized Widom 601 sequence⁴⁹. DNA (0.2 μM) was mixed with N-BRDT(1) (0.3-300 μM) in a final volume of 6 μl and incubated for 30 minutes prior to native-PAGE electrophoresis (4°C) and visualization with ethidium bromide staining. Unbound DNA was quantified using a Typhoon imager with ImageQuant software and the obtained data used to calculate the binding affinity indicated. (b) EMSA titrations of N-BRDT(1) interacting with either 66 or 25 bp radio-labelled DNA, as indicated. Shifted bands containing BRDT(1)-DNA complexes with one (filled arrowhead) or two (unfilled arrowhead) N-BRDT(1) molecules are indicated. Affinities shown were calculated from quantification of unbound DNA as detailed in (a). (c) EMSA titrations of N-BRDT(1) interacting with either 167 bp Widom DNA, unmodified nucleosomes or nucleosomes carrying acetylated H3(K18_{ac}K23_{ac}) and/or H4(K5_{ac}K8_{ac}) tails, as indicated.

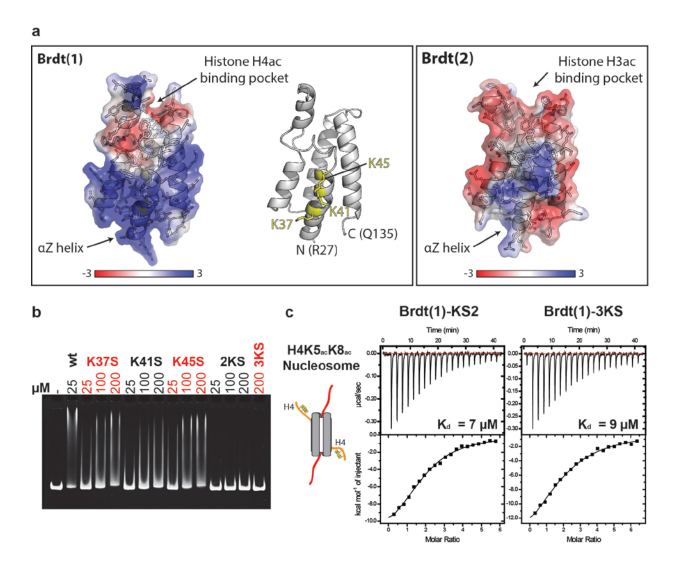


Figure 4. Residues involved in BRDT-BD1 DNA binding are distinct from those required for histone H4 recognition. (a) Crystal structure of human BRDT-BD1 (PDB code – $2RFJ^{14}$) and homology model of human BRDT-BD2 (generated using the Phyre2 web server²⁸), as indicated. Structures are shown as cartoons with side chain sticks and partially transparent electrostatic surfaces generated using PBD2PQR and APBS ($^{53-55}$). The positively charged patch centers on the αZ helix, which contains lysines K37, K41 and K45 (highlighted in yellow). (b) EMSA of N-BRDT(1) lysine mutants (concentrations as indicated) binding to 167 bp double stranded DNA (1 μM). Mutant '2KS' is a double mutant K37S, K41S. Mutant '3KS' contains all three point mutations. (c) ITC profiles for N-BRDT(1) mutants interactions with H4K5_{ac}K8_{ac} acetylated nucleosomes, as indicated.

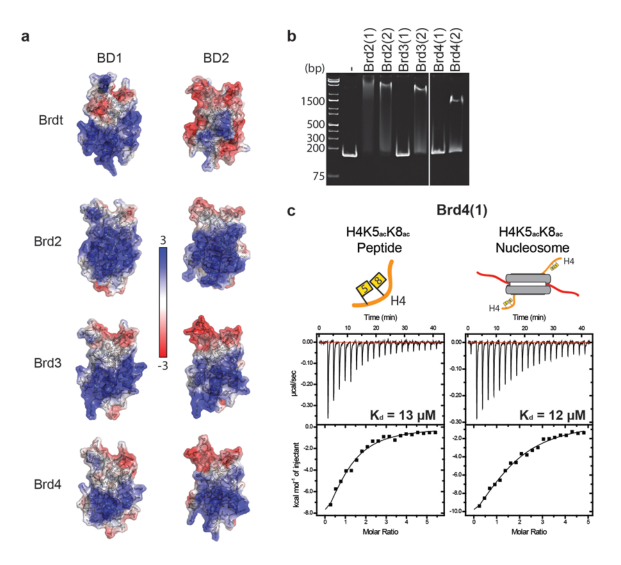
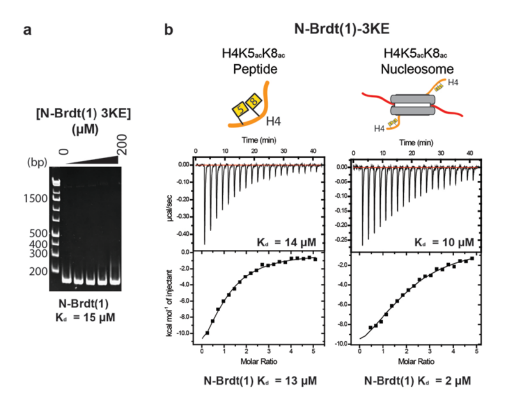
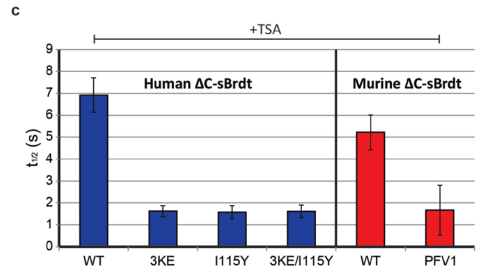


Figure 5. Other BET bromodomains interact with DNA. (a) Structures of the human BET bromodomains (PDB codes: BRDT-BD1 (2RFJ); BRD3-BD1 (2NXB); BRD3-BD2 (2OO1); BRD4-BD1 (2OSS, N-terminally truncated to R58 for comparison with other BD1 structures); BRD4-BD2 (2OUO)¹⁴; BRD2-BD1 (1X0J)²⁶; BRDT-BD2 structure is a homology model generated using the Phyre2 web server²⁸; BRD2-BD2 (2DVV)). Structures are shown as cartoons with side chain sticks and partially transparent electrostatic surfaces generated using PBD2PQR and APBS⁵³⁻⁵⁵. (b) EMSA of BET bromodomains (as indicated) interacting with 167 bp double stranded DNA. DNA (0.5 μM) was mixed with BET bromodomains (100 μM) in a final volume of 4 μl and incubated for 30 minutes prior to native-PAGE electrophoresis at 4 °C and visualization with ethidium bromide staining (C) ITC profiles for BRD4(1) interactions with H4K5_{ac}K8_{ac} peptides and nucleosomes, as indicated.





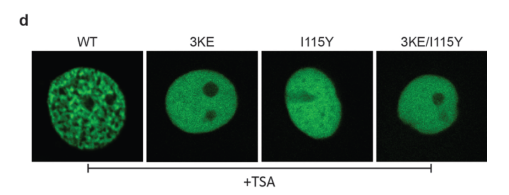


Figure 6. DNA binding by BRDT(1) is important for BRDT localization and chromatin compaction in cells. (a) EMSA of N-BRDT(1) lysine mutants (concentrations: 3 μM, 12.5 μM, 50 μM and 200 μM) binding to 167 bp double stranded DNA. DNA (0.4 μM) was mixed with N-BRDT(1) 3KE in a final volume of 5 μl and incubated for 30 minutes prior to native-PAGE electrophoresis at 4° C and visualization with ethidium bromide staining. (b) ITC profiles for N-BRDT(1) 3KE interactions with either acetylated histone H4 tail peptides or acetylated nucleosomes (both H4K5_{ac}K8_{ac}), as indicated. (c) Fluorescence recovery after photobleaching (FRAP) analysis of human and murine Δ C-sBRDT constructs in the presence of TSA-induced histone hyperacetylation. Cos7 cells were transfected by vectors expressing GFP-tagged WT and mutant Δ C-sBRDT constructs (as indicated) and cells were treated with the histone deacetylase inhibitor TSA (100 ng ml⁻¹) to induce histone hyperacetylation. A decrease in fluorescence recovery half-life (t_{1/2}) indicates an increase in protein mobility, and reduced chromatin association. Error bars show the standard error of the mean (s.e.m.) (d) Confocal microscopy images of representative transfected cells following TSA treatment.

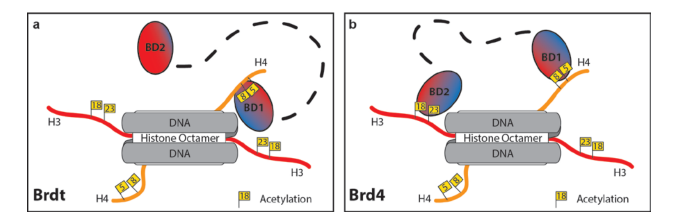


Figure 7. Schematic models of BRDT and BRD4 bromodomains interacting with acetylated nucleosomes. (a) BRDT interacts with acetylated nucleosomes via its BD1 domain. Binding may be initiated through non-specific interactions with DNA which allow BRDT to localize to chromatin. Specificity is generated through recognition of tandem acetylated lysine residues (K5_{ac}/K8_{ac}) on the histone H4 tail, whilst the affinity of the interaction is enhanced by bivalent interaction with both the histone tail and DNA. In contrast, BRDT-BD2 does not interact with acetylated nucleosomes and therefore is flexibly tethered to nucleosomes via BRDT-BD1. BRDT-BD2 may function to recruit as-yet-unknown acetylated non-histone proteins to the chromatin. (b) BRD4 interacts with acetylated nucleosomes via both its BD1 and BD2 domains. Our results indicate that BRD4-BD1 binds to nucleosomes through the acetylated histone H4 tail and does not additionally interact with DNA. Bivalent binding of BRD4 through both bromodomains has previously been shown to enhance BRD4 binding affinity for nucleosomes by 2.6-fold³⁷.