The first bromodomain of the testis-specific double bromodomain protein Brdt is required for chromocenter organization that is modulated by genetic background

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Mice homozygous for a mutation (Brdt ΔBD1/ΔBD1) lacking the first bromodomain of Brdt, a testis-specific member of the BET family of double-bromodomain containing proteins, are sterile and exhibit profound defects in chromatin remodeling during spermiogenesis. We have now observed that a prominent feature of the aberrant spermatid nuclei is a fragmented chromocenter, a structure comprised of peri-centromeric heterochromatin. There was a concomitant increase in the levels of heterochromatin protein 1 alpha (H1α), suggesting that the presence of multiple chromocenters was correlated with a spread of heterochromatin beyond the normal centromeric region. Brdt protein was normally present throughout the nucleus but was excluded from the chromocenter. A more densely staining region of Brdt protein appeared to separate sirtuin 1 (Sirt1) protein from contact with the chromocenter. Although still nuclear, this unique localization of Brdt protein was lost in Brdt ΔBD1/ΔBD1 mutant spermatids and Brdt and Sirt1 overlapped around the chromocenters. There was also ectopic localization of the H1 histone family, member N, testis-specific (H1Nnt) protein in Brdt ΔBD1/ΔBD1 mutant spermatids, which may be linked to the previously reported loss of polarized localization of peri-nuclear heterochromatin foci. The extent of chromocenter fragmentation was more severe and penetrant in mutant testes on a pure 129Sv/Ev background as compared to a pure C57Bl/6 background. Indeed, all aspects of the mutant phenotype were more severe on the 129Sv/Ev background. Contrary to previous studies in genetic models where fragmented chromocenters were observed in spermatids, the Brdt ΔBD1/ΔBD1 mutant spermatids do not undergo apoptosis (on either background). These observations suggest that the first bromodomain of Brdt is critical in the formation and/or maintenance of an intact chromocenter and implicate this structure in proper remodeling of the chromatin architecture of the sperm head.

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Introduction

Spermiogenesis is a unique developmental process in which the male genome must be properly packaged in order to retain its integrity once exiting the body. During spermiogenesis dramatic changes occur in both chromatin architecture and cell morphology. The condensation and elongation of the sperm nucleus involves a highly regulated but poorly understood replacement of nucleosomes with transition proteins and then protamines (Govin et al., 2004). The first striking chromatin reorganization to occur after the completion of meiosis is the formation of the chromocenter. Starting in stage I round spermatids the chromocenter is formed by the coalescence of the centromeric heterochromatin of each chromatid in the center of the nucleus, while their telomeres are associated mostly with the nuclear envelope (Haaf and Ward, 1995; Meyer-Ficca et al., 1998; Zalensky et al., 1995). The chromocenter remains intact throughout spermiogenesis and can still be found at the center of the mature sperm nucleus (Hoyer-Fender et al., 2000; Zalensky et al., 1995). It has further been shown that the chromatin is not randomly organized around the chromocenter, but rather whole chromosome domains are arranged parallel to the chromocenter (Haaf and Ward, 1995) This has led to the hypothesis that the chromocenter creates a defined nuclear topology and acts as a starting point for the sequential reorganization of chromatin during spermiogenesis (Meyer-Ficca et al., 1998). The mechanisms for establishing and maintaining the chromocenter are not known.

The presence of multiple or ‘fragmented’ chromocenters has been shown to be correlated with mis-localization of the testis specific histone H1 family member N (H1Nnt) (Catena et al., 2006). H1Nnt is normally specifically localized to the apical pole of round and elongating spermatids where it suppresses the formation of perinuclear...
heterochromatin foci under the acrosome (Martianov et al., 2005). In two mutant mouse models, Tbp1fl/fl and Hmgb2−/−, the presence of fragmented chromocenters is correlated with aberrant, bipolar localization of H1fnt (Catena et al., 2006). How aberrant localization of H1fnt affects latter stages of elongation and condensation is not known as the spermatids arrest and undergo apoptosis before the completion of spermiogenesis. In these two models, the process that gives rise to multiple or ‘fragmented’ chromocenters has not been investigated. Indeed, the fundamental question of whether an intact chromocenter is formed and is then fragmented or whether the individual centromeres fail to properly coalesce is not known.

Brdt is a testis-specific member of the BET family of chromatin interacting proteins, all of which contain two tandem bromodomains (Chua and Roeder, 1995), and binds to acetyl-H4, competes with the Sir2 deacetylase to stop the spread of transcriptional silencing at constitutive heterochromatin–euchromatin boundaries (Ladurner et al., 2003; Matangkasombut and Buratowski, 2003). There are four mammalian BET genes, Brd2, Brd3, Brd4 and Brdt, each of which is expressed in the testis but in distinct patterns (Shang et al., 2004). Brd2 and Brd4 are essential genes as null mutants of either are embryonic lethal (Houzelstein et al., 2002; Shang et al., 2009). As Brdt is testis-specific, a mutation in the gene that completely removed the first bromodomain (BD1), BrdtΔBD1, did not affect viability, but rather caused complete male sterility (Shang et al., 2007).

Brdt expression is restricted to pachytenie and diplotene spermatocytes and round spermatids; however, no obvious morphological defect was seen in these cells in BrdtΔBD1 mutant testes (Shang et al., 2007). Rather, the first obviously visible defects were observed in stage IX spermatids which fail to properly elongate and the heterochromatin foci normally observed at the nuclear envelope were absent. The severity of the BrdtΔBD1 phenotype in elongating spermatids and sperm varied between mice or even among tubules of a single testis. Some spermatids seemed to elongate fairly normally and some mice had epididymal sperm. Epididymal sperm were always grossly morphologically abnormal, with excess cytoplasm, misshapen heads, and deformed or truncated tails that often lacked the midpiece. The initial observations were made on BrdtΔBD1 mutant mice that were maintained on a mixed genetic background of C57BL/6 J (B6) and 129/SvEv (129), which we speculated may have contributed to this heterogeneity. We have therefore backcrossed the mutation onto genetically pure C57BL/6 J and 129/SvEv mice, and investigated the phenotype in greater detail in each background. We have uncovered striking defects in the chromocenters of the mutant spermatids, increased heterochromatin, and ectopic expression of specialized histones — all of which may contribute mechanistically to the abnormalities in chromatin remodeling.

Materials and methods

Backcrossing to pure 129 and B6 backgrounds

A mixed 129 Sv (129) × C57Bl/6 J (B6) male heterozygous for the BrdtΔBD1 mutant allele was mated with two pure strain 129 or B6 females. A single heterozygous mutant male offspring was then mated with those same two females. This was repeated for five more generations after which two new pure strain females were used for two more generations. One final cross of a heterozygous mutant female offspring with a pure 129 or B6 male was carried out to insure that the Y-chromosome was also from the pure background. This process resulted in progeny that were ten generations backcrossed, and we considered these mice to now carry the BrdtΔBD1 mutant allele on a pure background.

Histological analysis, immunohistochemistry and immunofluorescence

Histological analysis and periodic acid Schiff (PAS) staining were carried out according to our laboratory’s standard protocols as previously published (Chung and Wolgemuth, 2004; Shang et al., 2007). Bouin’s fixed sections were used for Hematoxylin and Eosin staining, PAS staining and also for anti-trimethyl-histone H3 (Lys9) (H3K9me3) immunostaining. The H3K9me3 immunostaining used rabbit polyclonal primary antibody (Upstate, cat#07-523) at a concentration of 1:200 and the Vectastain ABC Kit-Rabbit IgG (Vector Laboratories, Inc.). Four percent paraformaldehyde (PFA) fixed sections were used for immunofluorescence with H1fnt, H4, and H2AZ antibodies and immunostaining with Hmgb2 antibodies. The rabbit polyclonal anti-H1fnt primary antibody (Santa Cruz, cat# sc-136700) was used at 1:200, the goat polyclonal anti-H4 primary antibody (Upstate, cat# sc-8657) was used at 1:300, the rabbit polyclonal anti-H2AZ primary antibody (AbCam, cat# 4174–100) was used at 1:200. The rabbit polyclonal anti-Hmgb2 primary antibody (Epitomics, cat# T2134) was used at 1:200. The following secondary antibodies were used: Alexa Fluor-488 donkey anti-goat IgG (Molecular Probes, cat# A11055) at 1:300, Alexa Fluor-488 goat anti-rabbit IgG (Molecular Probes, cat #11008) at 1:300, and Alexa Fluor-594 goat anti-rabbit IgG (Molecular Probes, cat# 11012) at 1:300. The Vectastain ABC Kit-Rabbit IgG (Vector Laboratories, Inc.) was used for immunodetection of Hmgb2.

To investigate subcellular localization of Brdt, a single cell suspension of spermatogenic cells was made from wild-type and mutant testes by removing the tunica albuginea and placing the tubules in cold PBS. The tubules were manually sheared with scissors and by pipetting, and then passed through a 40 μm filter. The resulting suspension was dropped on slides, air dried, and then fixed in 4:1 methanol:acetone for 10 min. Two Brdt antibodies generated by our lab (Shang et al., 2007) were both used at a concentration of 1:300 and yielded identical results. The rabbit polyclonal anti-Sirt1 primary antibody (AbCam, cat#ab12193) was used at a concentration of 1:200. The following secondary antibodies were used: DyLight 594 Fab fragment donkey anti-rabbit IgG (Jackson Immunoresearch cat# 711-517-003 04) at 1:300 and Alexa Fluor-488 goat anti-rabbit IgG (Molecular Probes, cat#11008) at 1:300. The staining was done sequentially, first with anti-Sirt1, followed by DyLight 594, then anti-Brdt, then Aleza Fluor-488, followed by DAPI.

Quantification of fragmented chromocenters

PFA fixed histological sections for both genotypes on both backgrounds where prepared by the histology technician and labeled numerically, without identifying genotypes or strain background. The investigators then stained the sections with anti-H3K9me3 and PAS and proceeded with the quantitation of the numbers of chromocenters per round spermatid in 100 tubules, without knowledge of the genotype or background of the sections. A cell was counted as a round spermatid if it had a visible acrosome, as having a fragmented chromocenter if there were three foci of H3K9me3 staining, and as having a severely fragmented chromocenter if there were four or more foci. Statistical significance was assessed using a standard two-tailed T-test.

Immunoblot analysis

For immunoblot analysis of components in the testicular cell chromatin our previously published procedures were followed (Shang et al., 2007). Briefly, a single cell suspension for control and mutant 129 testes were made as described above. The cells were pelleted at...
2500 g for 10 min, and homogenized in modified RIPA buffer [50 mM Tris–HCl, pH 7.4, 150 mM NaCl, 1% NP-40, 0.5% sodium deoxycholate and protease inhibitor cocktail (Roche)], incubated for 30 min at 4 °C, and centrifuged at 12,000 g for 10 min at 4 °C. The pelleted chromatin was boiled for 5 min in 1× NuPAGE loading buffer (Invitrogen). Cellular lysates from whole 129 and B6 testes were made with standard RIPA buffer and boiled for 5 min in 1× NuPAGE loading buffer (Invitrogen). Chromatin preparations and lysates were separated on NuPAGE 4–12% Bis–Tris gels (Invitrogen), transferred to PVDF membranes and the blots were incubated with rabbit polyclonal anti-Fibrillarin antibody (GenScript, cat# A01462) at a concentration of 1:3000, goat polyclonal anti-Hp1α antibody (Abcam, cat# ab77256) at 1:3000, goat polyclonal anti-H4 antibody (Upstate, cat# sc-8657) at 1:5000, and rabbit polyclonal anti-Hmgb2 antibody (Epitomics, cat# T2134) at 1:3000.

Quantitative real-time PCR

To perform quantitative real-time PCR, whole testes from control 129 and B6 mice were homogenized in Trizol reagent (Invitrogen) and we extracted RNA according to the manufacturer's protocol. cDNA was synthesized using random hexamers and the TaqMan Reverse Transcription Kit (Applied Biosystems) and qRT-PCR performed with the Power SYBR Green master mix (Applied Biosystems) on a ABI Prism 7500 (Applied Biosystems). Each reaction was performed in triplicate and with RNA from three different mice from each background. The values were normalized to the expression of ribosomal protein, large, PO (Rplp0) as an internal control. The following primers were used:

- Hmgb2 Forward: GCCACAGCCCTTTACATGGG
- Hmgb2 Reverse: GCCACAGCCCTTTACATGGG
- Rplp0 Forward: CAAAGCTGAACAAAGGAAGAG
- Rplp0 Reverse: AATTAAGCAGGCTGACTTGGTTG
- Tbpl1 Forward: TTTGGTGCCAGACGTTTAGC
- Tbpl1 Reverse: GCCACAGCCCTTTACATGGG
- Rdh50 Forward: CAAGCTGAACAAAGGAAGAG
- Rdh50 Reverse: AATTAAGCAGGCTGACTTGGTTG

Sperm counts

Sperm were collected into non-capacitation medium (10 mM HEPES, 95 mM NaCl, 4.8 mM KCl, 1.2 mM MgSO4, 1.2 mM KH2PO4, 20 mM Sodium Lactate, 5 mM Sucrose, pH 7.4) at 37 °C from both caudal epididymides of individual animals as per our laboratory's standard protocol as previously published (Chung et al., 2011; Travis et al., 2001). Briefly, cauda were manually sheared with scissors and forceps. The solution was then passed through a 100 μm filter and medium was added up to 1 ml. For sperm suspensions from control and heterozygous mice from 129 animals and all cauda from B6 animals the sperm solution was then diluted 1:10 in medium and counted on a hemocytometer under an inverted light microscope. The 1:10 dilution was not needed to count the 129 mutant sperm. The individual counting the sperm did not know the genotype or background of the sperm to avoid bias.

Results

Mutant round spermatids exhibit fragmented chromocenters

In our initial characterization of the phenotype of BrdtΔBD1/ΔBD1 mutant mice on a mixed 129/B6 background, heterogeneity was noted in the testicular morphology between mutant mice, as well as among tubules in a single testis. To assess if genetic background contributed to this heterogeneity, we backcrossed the mutation onto pure 129 and B6 backgrounds. Histological analysis of the mutant testes on the 129 background revealed a heretofore unreported component of
the mutant phenotype—a striking increase in the presence of fragmented chromocenters in round spermatids. Since the earliest defect we had previously observed in the Brdt^{ΔBD1/ΔBD1} mutant testis occurred in stage IX elongating spermatids, we initially examined the testes of 24 day-old animals, as at this age no tubules would have developed beyond stage VIII. DAPI-stained control 129 testes revealed that the majority of spermatids contain a single focus of dense heterochromatin or one large focus (the chromocenter) and a smaller lighter secondary focus (the nucleolus) (Fig. 1A, wide arrows), along with a low basal level of fragmented chromocenters—defined as those spermatids that had three or more foci (Fig. 1A, arrowheads; Table 1). In striking contrast, the majority of round spermatids in mutant 129 testes contain severely fragmented chromocenters, with most spermatids exhibiting at least three and frequently four or more foci of heterochromatin (Fig. 1B, arrows; Table 1).

As secondary spermatocytes also exhibit foci of heterochromatin (Cobb et al., 1999), it was critical to ensure that the cells in question were indeed round spermatids. We therefore performed PAS staining which permitted visualization of the acrosome. To concomitantly visualize the chromocenters, the sections were also stained with anti-H3K9me3 (Greaves et al., 2006). Cells with severely fragmented chromocenters did indeed have developing acrosomes and thus could only be spermatids (Fig. 1D, arrows).

**Brdt protein is nuclear but excluded from the chromocenter**

Our initial studies clearly showed that Brdt is a nuclear protein and associated with chromatin (Shang et al., 2007). Given the new observation of chromocenter abnormalities, we characterized the distribution of Brdt protein within sub-nuclear regions of spermatids using fluorescent immunostaining. Brdt protein localized throughout the nucleus as predicted, but did not extend into the dense region of the chromocenter (Fig. 2B). With these preparations, Brdt protein appeared to be more concentrated immediately surrounding the chromocenter (Fig. 2B, inset). In contrast, although the truncated protein produced in Brdt^{ΔBD1/ΔBD1} mutant spermatids localized to the nucleus, it is more evenly distributed throughout the nucleus (Fig. 2G, inset). This observation suggests that the BD1 is involved in determining the sub-nuclear association of Brdt to specific chromatin.

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**Table 1**

<table>
<thead>
<tr>
<th></th>
<th>B6+/+</th>
<th>B6^{ΔBD1/ΔBD1}</th>
<th>129+/+</th>
<th>129^{ΔBD1/ΔBD1}</th>
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<td>Total number of round spermatids (RS) per tubule of 100 tubules examined</td>
<td>128.4 ± 15.1</td>
<td>115.5 ± 11.2</td>
<td>123.8 ± 7.5</td>
<td>100.6 ± 9.7</td>
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<td>RS with non-fragmented chromocenters (avr. per tubule of 100 tubules examined)</td>
<td>195.3 ± 13.1</td>
<td>84.3 ± 6.9</td>
<td>104.3 ± 5.4</td>
<td>33.5 ± 4.0</td>
</tr>
<tr>
<td>RS with fragmented chromocenters (avr. per tubule of 100 tubules examined)</td>
<td>8.9 ± 2.0</td>
<td>31.2 ± 4.3</td>
<td>19.5 ± 2.1</td>
<td>67.1 ± 5.7</td>
</tr>
<tr>
<td>Percent of RS that have fragmented chromocenters (≥ 3 heterochromatin foci)</td>
<td>7%</td>
<td>23%</td>
<td>17%</td>
<td>67%</td>
</tr>
<tr>
<td>Percent of RS that have severely fragmented chromocenters (4 heterochromatin foci)</td>
<td>&lt; 1%</td>
<td>6%</td>
<td>2%</td>
<td>55%</td>
</tr>
</tbody>
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**Fig. 2.** Dense Brdt expression separates the chromocenter from Sirt1 protein. Fluorescent staining of control (A–E) and mutant (F–J) round spermatids with Brdt and Sirt1 antibodies and the merge of the two. (A–B) Brdt is present in the nucleus of round spermatids, with regions of denser staining surrounding the chromocenter from which it is excluded (inset merge of Brdt and DAPI). (C) Sirt1 is also present in the nucleus of round spermatids, and is also excluded from the chromocenter (inset in E). (D–E) Brdt and Sirt1 protein localization overlap except for the region directly surrounding the chromocenter where Brdt is strongly detected but Sirt1 is excluded (green fluorescence only). (F–G) Truncated Brdt protein is still present in the nucleus of mutant round spermatids, but the denser staining around the chromocenter is no longer seen (inset merge of Brdt and DAPI). (H) Sirt1 expression is still nuclear and excluded from the chromocenter. (I–J) Brdt and Sirt1 protein localization overlap completely (no exclusive green fluorescence), and both are present directly adjacent to the chromocenter. While there are still some regions of dense Brdt staining, it is no longer localized around the chromocenter. Scale bar, 12.5 μm.
Dense Brdt expression separates the chromocenter from sirtuin 1 protein

Sirtuin 1 (Sirt1) is the mammalian homologue of the yeast histone deacetylase Sir2 and like Sir2, is part of the machinery that keeps constitutive heterochromatin transcriptionally silenced (Palacios et al., 2010). Acetylated H3 lysine 9 is one of the targets of Sir2 deacetylase activity (Imai et al., 2000). Deacetylation of H3 by Sir2 can be a required step before tri-methylation of lysine 9 by histone methyltransferase Su(var)3-9 (Ekwall, 2004; Shankaranarayana et al., 2003). The chromodomain of heterochromatin protein 1 binds to tri-methylated H3K9, making this histone modification a characteristic hallmark of heterochromatin (Bannister et al., 2001; Lachner et al., 2001). The mammalian homologues of the methyltransferase, Su39h1, and the testis-specific methyltransferase Su39h2, are both present in round spermatids where they are restricted to the chromocenter (O’Carroll et al., 2000). Immunostaining to determine the localization of Sirt1 revealed that it was indeed expressed in round spermatids, but was completely excluded from the chromocenter (Fig. 2C, and insets in 2D, E). Double-staining for Sirt1 and Brdt showed that the dense region of Brdt staining around the chromocenter separated the region of Sirt1 localization from the chromocenter proper (Fig. 2D–E, green fluorescence only, insets in 2D,E). This created essentially a ‘buffer’ region between the chromocenter and Sirt1 protein. In mutant round spermatids, where there is no region of denser, peri-chromocentric Brdt staining, this buffer was lost and Sirt1 now also localized adjacent to the chromocenter (Fig. 2H–J, no exclusive green fluorescence).

Localization of the histone variant H2A.Z is unchanged in mutant spermatids with fragmented chromocenters

BET proteins have been implicated in delimiting heterochromatin–euchromatin boundaries in the yeast model wherein Bdfl is involved in the deposition of the histone variant Htz1 (Krogan et al., 2003b; Mizuguchi et al., 2004; Raisner et al., 2005; Zhang et al., 2005). The mammalian homologue of Htz1, H2A.Z, results in the spread of euchromatin in the yeast model wherein Bdf1 is involved in euchromatin boundaries in the yeast model wherein Bdf1 is involved (Ekwall, 2004; Shankaranarayana et al., 2003). The mammalian homologues of the methyltransferase, Su(var)39h1, and the testis-specific methyltransferase Su(var)39h2, are both present in round spermatids where they are restricted to the chromocenter (O’Carroll et al., 2000). Immunostaining to determine the localization of Sirt1 revealed that it was indeed expressed in round spermatids, but was completely excluded from the chromocenter (Fig. 2C, and insets in 2D, E). Double-staining for Sirt1 and Brdt showed that the dense region of Brdt staining around the chromocenter separated the region of Sirt1 localization from the chromocenter proper (Fig. 2D–E, green fluorescence only, insets in 2D,E). This created essentially a ‘buffer’ region between the chromocenter and Sirt1 protein. In mutant round spermatids, where there is no region of denser, peri-chromocentric Brdt staining, this buffer was lost and Sirt1 now also localized adjacent to the chromocenter (Fig. 2H–J, no exclusive green fluorescence).

Levels of heterochromatin are elevated in mutant testes

To understand the mechanisms by which loss of the first bromodomain of Brdt results in multiple chromocenters, we asked whether the fragmented chromocenters might actually reflect an increase in the overall amount of heterochromatin in mutant spermatids. Alternatively, if the aberrant structures were the result of a failure of the centromeres to coalesce properly, the amount of heterochromatin in the spermatids would be unchanged. In the Tbp1−/− mutant model, multiple chromocenters were visualized by staining for H3K9me2, a known
Fig. 4. H1fnt is ectopically localized in some 129 mutant round spermatids. (A–B) Fluorescent staining to localize H1fnt in day-24 histological sections reveals a single region of apical expression in control 129 round spermatids (A) and ectopic localization in mutant 129 round spermatids (B, open arrowheads). (C) Immunostaining of H1fnt in a mutant 129 adult histological section shows non-elongated mutant spermatids with H1fnt surrounding the nucleus (arrowheads). Scale bar, 25 μm.

Component of heterochromatin but the total levels of Hp1α in control and mutant chromatin were never measured (Martianov et al., 2002). We therefore extracted chromatin from both control and mutant testes of 129 mice and measured the levels of Hp1α protein. We observed a greater than three-fold increase in the levels of Hp1α protein in chromatin from the mutant testes (Fig. 5). In nuclei of round spermatids, HP1α is found both at centromeres and also in the nucleolus (Horakova et al., 2010). To determine whether increased levels of Hp1α were deposited at both sites, we measured levels of fibrillarin, a nucleolar structural protein (Fig. 5). Levels of fibrillarin were unchanged in the mutant testes, suggesting that the number of nucleoli is the same and thus that the increased levels of Hp1α reflected increased heterochromatin at the centromeric regions.

All aspects of the Brdt<sup>ΔBD1/ΔBD1</sup> mutant phenotype detected are background dependent

Fragmented chromocenters

Fragmented chromocenters were also observed in the round spermatids of homozygous mutants on the B6 background, however at a lower frequency (Fig. 1C, arrowheads). The number of spermatids with fragmented chromocenters was quantified for the various genotypes on both genetic backgrounds (Table 1). The basal level of fragmented chromocenters was significantly higher in the control 129 background than in the control B6 background (17% vs 7%; pValue ≤ 5.0 × 10⁻¹³). The number of cells exhibiting fragmented chromocenters in the Brdt<sup>ΔBD1/ΔBD1</sup> mutant on the 129 background was significantly increased compared to the controls (67% vs 17%; pValue ≤ 5.0 × 10⁻¹⁵), and also compared to the Brdt<sup>ΔBD1/ΔBD1</sup> mutant on the B6 background (67% vs 23%; pValue ≤ 5.0 × 10⁻⁷). Severely fragmented chromocenters were rarely seen in control spermatids of either the 129 (2%) or the B6 (<1%) backgrounds but were more frequent in the Brdt<sup>ΔBD1/ΔBD1</sup> mutants, significantly so in the 129 background (55% vs 6% pValue ≤ 5.0 × 10⁻¹³) (Table 1).

Spermatid elongation

Starting in stage IX, none of the spermatids in the 129 mutant testes elongate properly. Even in later stages of spermiogenesis, many spermatids remain round with no obvious change to the morphology of the nucleus (Fig. 6C arrow, and inset). Those cells that do begin to elongate remain aberrantly short and do not develop the characteristic hook that should be seen in later stages (Fig. 6E, and G). Compaction of the chromatin appears to be occurring as the size of the nucleus is greatly reduced when compared to round spermatids; however, without proper elongation this results in the presence of abnormal, very small round cells (Fig. 6C, E, G – arrowheads, inset in E and G). These cells are unlikely to be anything other than spermatids as the progression from the presence of round spermatids to the appearance of small, non-elongate spermatids can be followed directly in testes from mutant mice in the first wave of spermatogenesis that occurs during ensuing days of post-natal development. All of the various abnormalities described in the mixed background mutant testes (Shang et al., 2007) are now consistently seen in all tubules of strain 129 mice homozygous for the Brdt<sup>ΔBD1</sup> mutation (Table 2). Homozygous male mutants on the B6 background were still infertile but spermiogenesis proceeded more normally. Although aberrant elongation can still be seen in mutant tubules (Fig. 6D, F and H arrowheads, and inset in D and F), approximately 70% of the tubules contain some spermatids that appear to be elongating and compacting fairly normally and some later stage tubules look grossly morphologically normal (Fig. 6H). Some aberrantly short spermatids are present and 85% of all tubules still contain non-elongated small compacted spermatids (Fig. 6D, F, H arrowheads, and inset in D and F), but these cells are not prevalent (Table 2).

Disruption of cellular associations

We also observed that there was a loss of order of the cellular layers in the 129 mutant seminiferous epithelium which had not been reported in the mutant on a mixed background (Shang et al., 2007). For example, round and aberrantly elongating spermatids can be seen in the spermatocyte layer of the tubule (Fig. 6C, E, G – open arrows, inset in E) and round spermatids are sometimes found in the very center of tubules (Table 2). In contrast, on the B6
Fig. 6. The abnormalities in BrdTΔBD1/ΔBD1 mice are more severe and uniform on the 129 background. (A–H) H&E staining of histological sections of (A) control 129 stage X tubule, (B) control B6 stage X tubule, (C) mutant 129 stage X* tubule, (D) mutant B6 stage X tubule, (E) mutant 129 stage XII* tubule, (F) mutant B6 stage XII tubule, (G) mutant 129 stage V* tubule, and (H) mutant B6 stage V tubule. Grossly normally elongating spermatids are absent in the mutant 129 testes, but predominate the B6 mutant testes. Conversely, compacted but not elongated spermatids are present (C–H, arrowheads and D–G inset) in B6 testes, but they predominate the 129 testes (see Table 2 for quantification). Spermatids which have neither compacted nor elongating (C, arrow and inset) can be found only in the 129. In 129 testes some spermatids can be found in the spermatocyte layer of the tubules (C, E, and G, open arrows and E inset). Asterisk: although the severe disruption of the order of the seminiferous epithelium in the 129 mutant made precise staging difficult, we refer to approximately staged tubules with a roman numeral followed by an asterisk. Scale bar, 25 μm.
background, the loss of order of the seminiferous epithelium was observed at a low frequency, comparable to control testes, and these phenotypes were consistent with all tubules containing overtly properly elongating spermatids (Table 2).

**Epididymal sperm**

We had previously reported that there was heterogeneity in the presence of epididymal sperm on a mixed background: some animals had considerable numbers of epididymal sperm while others had very nearly empty epididymides (Shang et al., 2007). Histological examination of the 129 mutant animals consistently revealed that all three major parts of the epididymis were virtually devoid of sperm, with only occasional sloughed off spermatocytes and round spermatids and cytoplasmic bodies and very small and round spermatids (Fig. 7A). In contrast, on the B6 background sperm are consistently present in the three regions of the epididymis (Fig. 7A). Morphologically, there was still a great deal of heterogeneity. Some sperm looked fairly ‘normal’, with close to the proper hook shape to the head and containing midpieces and full tails (Fig. 7C arrow). Others however, were highly aberrant and resembled the 129 mutant sperm (Fig. 7C). Correspondingly, caudal epididymal sperm counts were extremely low. Mutant 129 epididymides had an average of $4.02 \times 10^5$ total sperm (roughly 4 sperm per slide), which was 10-fold below the bottom threshold for fertility ($4 \times 10^6$) (Meistrich, 1982; Reid et al., 1981) and 75-fold below the 129 control average ($3.03 \times 10^7$) (Fig. 7B). The few sperm that were present in the 129 epididymis appear very abnormal with excess cytoplasm, misshapen heads, and deformed tails and midpieces (Fig. 7C). Interestingly, the average sperm count for the heterozygous *Brdt*ΔBD1/ΔBD1 animals on the 129 background was only $1.96 \times 10^7$ or about two-thirds that of the wild-type. Using a standard two-tailed t-test this was a significant loss of sperm at a pValue of 0.005. This is the first observed heterozygous phenotype; however, this number of sperm is still well above the threshold of fertility. In *Brdt*ΔBD1/ΔBD1 mutants on the B6 background, caudal sperm counts yielded an average of $2.41 \times 10^6$ sperm, also below the threshold for fertility but almost six-fold more than the 129 mutant (Fig. 7B). This difference in the number

![Fig. 7. Differences in epididymal sperm content of BrdtΔBD1/ΔBD1 mutants on 129 and B6 backgrounds.](image-url)

Table 2

Summary of morphological abnormalities in *Brdt*ΔBD1/ΔBD1 mutant mice. *Brdt*ΔBD1/ΔBD1 mutant mice on the 129 background never have normally elongating spermatids and all tubules contain very small condensed but not elongated spermatids. *Brdt*ΔBD1/ΔBD1 mutant tubules on the B6 background often contain normally elongating spermatids, but only 15% of all tubules look grossly morphologically normal. Defects in the order of the seminiferous epithelium occur at a greater frequency in 129 mutant tubules than in control tubules.

<table>
<thead>
<tr>
<th></th>
<th>B6+/+</th>
<th>B6ΔBD1/ΔBD1</th>
<th>129+/+</th>
<th>129ΔBD1/ΔBD1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percent of stage IX–XII tubules containing any normally elongating spermatids</td>
<td>100%</td>
<td>73%</td>
<td>100%</td>
<td>0%</td>
</tr>
<tr>
<td>Percent of all tubules containing very small compacted but not elongated spermatids</td>
<td>0%</td>
<td>85%</td>
<td>0%</td>
<td>100%</td>
</tr>
<tr>
<td>Percent of stage IX–XII tubules containing round spermatids</td>
<td>0%</td>
<td>0%</td>
<td>0%</td>
<td>12%</td>
</tr>
<tr>
<td>Percent of all tubules containing condensed spermatids in the spermatocyte layer</td>
<td>2%</td>
<td>9%</td>
<td>4%</td>
<td>23%</td>
</tr>
<tr>
<td>Percent of all tubules containing round spermatids in the tubule lumen</td>
<td>2%</td>
<td>3%</td>
<td>1%</td>
<td>23%</td>
</tr>
</tbody>
</table>
of sperm between the two mutants was statistically significant by a two-tailed t-test at a p-value of $0.5 \times 10^{-6}$.

Hmgb2 is expressed at higher levels in B6 round spermatids

Two other genes, Tbp1l and Hmgb2, have been shown by targeted mutagenesis to be required for proper chromocenter formation and/or maintenance (Catena et al., 2006). To ask whether either protein may be involved in the difference in the levels of fragmented chromocenters detected between the two backgrounds in both control and Brdt$^{\Delta BD1/\Delta BD1}$ mutant spermatids, we investigated their expression in testes from the two backgrounds. Quantitative real-time PCR revealed that expression of Tbp1l was comparable in the two backgrounds, but that Hmgb2 expression was nearly 4-times higher in the B6 testes (Fig. 8A). Immunoblotting with anti-Hmgb2 antibody showed that this mRNA difference was also reflected in the protein levels (Fig. 8B). Immunostaining of histological sections of testes from both strains placed side by side on one slide, confirmed that anti-Hmgb2 antibodies localized more intensely in both round spermatids and pachytene spermatocytes in the B6 testis as compared to 129 testes (Fig. 8C).

Discussion

This study reports heretofore unknown chromatin defects in Brdt$^{\Delta BD1/\Delta BD1}$ mutant spermatids and also illustrates the strongly genetic background-dependent nature of the phenotype. Our earlier analysis of the Brdt$^{\Delta BD1/\Delta BD1}$ mutant phenotype on a mixed 129/B6 background focused on defects in spermatid elongation and chromatin condensation (Shang et al., 2007). Upon breeding the mutation onto a genetically pure 129 background, a striking and highly penetrant phenotype was revealed — that of a high proportion of round spermatids with multiple foci of dense heterochromatin, indicative of fragmented chromocenters. There were still no apparent defects during meiosis, raising the possibility that ΔBD1 may represent a hypomorphic allele. It is interesting to recall that in the yeast model, mutations in Bdf1 that were only in the first bromodomain did not have an effect on sporulation (Chua and Roeder, 1995). Rather it was BD2 and the ET domain that were required. It is possible that we have yet to detect a subtle defect in pachytene spermatocytes, and we are continuing to investigate meiosis in the mutant testis. However it is equally possible that the truncated Brdt protein is sufficient for proper meiosis and progression to round spermatids.

Although genes involved in chromatin organization, such as Tbp1l and Hmgb2, are required for proper formation and/or maintenance of the chromocenter (Catena et al., 2006), the mechanisms involved are as yet unknown. The question of whether fragmented chromocenters result of a spreading of centromeric heterochromatin or alternatively, a failure of the centromeres to coalesce into one area is also not known. What also emerges from the studies on our Brdt$^{\Delta BD1/\Delta BD1}$ mutant is that the term “fragmented chromocenter” is somewhat misleading. That is, multiple heterochromatin foci are seen as early as Stage I in our mutant, and the frequency of their appearance does not change throughout round spermatid development. It therefore does not seem that a single chromocenter is forming and then fragmenting apart, but rather that an intact singular chromocenter does not form. We propose that going forward, this phenomenon should be termed ‘multiple chromocenters’ because although the centromeric heterochromatin has partially coalesced, it did not form one unique chromocenter.

In our Brdt$^{\Delta BD1/\Delta BD1}$ mutant, the increased levels of HP1α in chromatin suggest that there is an expansion of heterochromatin. No such increase in the levels of HP1α was reported in either Hmgb2$^{-/-}$ or

![Fig. 8. Hmgb2 is expressed at higher levels in the B6 testis.](image-url)
We observed ectopic expression of H1fnt in a small but distinct subset of step VII–VIII Brdt\textsuperscript{BD1/ΔBD1} mutant spermatids, very similar to what was seen in the Tbp11 and Hmgb2 knockout models which also exhibit multiple chromocenters. We further noted that Brdt\textsuperscript{BD1/ΔBD1} spermatids that failed to elongate often had nuclei entirely surrounded by H1fnt. Since the apical localization of H1fnt blocks the formation of foci of heterochromatin at the nuclear envelope under the acrosome (Martianov et al., 2005), this ectopic distribution throughout the periphery of the nucleus could explain, at least in part, the previously observed loss of such foci in the spermatids of our Brdt\textsuperscript{BD1/ΔBD1} mutant (Shang et al., 2007). It has been speculated that such foci may connect chromatin to the cytoskeleton and could couple chromatin condensation with morphological changes in the cytoplasm and the spermatid head. Indeed, excess cytoplasm, and defective head morphology are two the most striking defects of Brdt\textsuperscript{BD1/ΔBD1} sperm.

The connection between the presence of multiple chromocenters, aberrant localization of H1fnt and abnormal sperm morphology is further supported by our studies of the Brdt\textsuperscript{BD1/ΔBD1} mutation on pure backgrounds. Multiple chromocenters are more prevalent in 129 mutant spermatids than B6 or mixed mutant spermatids, which explains why we did not observe this defect in our initial analysis. The defect in spermatid elongation is also much more severe and penetrant on the 129 background. The loss of polarity of H1fnt localization might reflect an overall loss of polarity in the nucleus that impedes proper elongation. Further, ectopic H1fnt localization is probably the cause of the loss of perinuclear heterochromatin foci. It has been hypothesized that the foci connect chromatin to the actin network in the sperm head and may couple chromatin condensation to overall morphological change (Martianov et al., 2005). Thus their loss may disconnect condensation from proper elongation. Both these aspects of H1fnt mis-localization tie chromocenter formation to elongation. It is important to note that completely morphologically normal sperm are never produced. This suggests that Brdt has additional functions beyond chromocenter formation, such as transcriptional or post-transcriptional regulation of genes that are required for elongation and sperm head and body formation. We are currently investigating such a possible role for Brdt with microarrays and Chip-Seq approaches.

The dynamic chromatin changes that occur during spermiogenesis are still poorly understood, but we have clearly shown the Brdt, and specifically its first bromodomain is essential for this process to proceed normally. The chromocenter is a central organizing structure for the chromatin of the spermatid and mature spermatocyte and it is now apparent that without complete Brdt function the proper architecture of the sperm nucleus cannot be achieved. The Brdt\textsuperscript{BD1/ΔBD1} mutant is unique relative to other mutations affecting sperm chromatin organization in that many spermatids survive, and as a result it may be ideal for further studies of the interconnections of chromatin architecture and sperm head formation. Further study of the mechanism of chromocenter formation and Brdt's role therein will help to understand the highly complex and ordered restructuring of the chromatin of the sperm nucleus.

Acknowledgments

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References


