BRD4 interacts with NIPBL and BRD4 is mutated in a Cornelia de Lange-like syndrome

Citation for published version: Olley, G, Ansari, M, Bengani, H, Grimes, GR, DDD study, Bickmore, W, Pradeepa, MM & FitzPatrick, D 2018, 'BRD4 interacts with NIPBL and BRD4 is mutated in a Cornelia de Lange-like syndrome' Nature Genetics. DOI: 10.1038/s41588-018-0042-y

Digital Object Identifier (DOI): 10.1038/s41588-018-0042-y

Link: Link to publication record in Edinburgh Research Explorer

Document Version: Peer reviewed version

Published In: Nature Genetics

General rights
Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.
BRD4 interacts with NIPBL and BRD4 is mutated in a Cornelia de Lange-like syndrome

(82 characters)

Gabrielle Olley1§, Morad Ansari1§, Hemant Bengani1, Graeme R Grimes2, James Rhodes3, Alex von Kriegsheim2, Ana Blatnik1,4, Fiona J. Stewart5, Emma Wakeling5, Nicola Carroll9, Alison Ross6, Soo-Mi Park10, DDD study11, Wendy A Bickmore1§, Madapura M Pradeepa1,7#, David R FitzPatrick1.

1. MRC Human Genetics Unit, MRC Institute of Genetics and Molecular Medicine at the University of Edinburgh, Edinburgh EH4 2XU, UK
2. MRC Institute of Genetics and Molecular Medicine at the University of Edinburgh, Edinburgh EH4 2XU, UK
3. Department of Biochemistry, Oxford University, South Parks Road, Oxford UK
4. Cancer Genetics Clinic, Institute of Oncology, Ljubljana, Slovenia
5. Department of Medical Genetics, Belfast City Hospital, Belfast BT9 7AB UK
6. Department of Medical Genetics, Ashgrove House, Foresterhill, Aberdeen, AB25 2ZA UK
7. School of Biological Sciences, University of Essex, Colchester CO4 3SQ, UK
8. North West Thames Regional Genetics Service, London North West Healthcare NHS Trust, Harrow HA1 3UJ, UK
9. South-East Scotland Regional Genetics Services, Western General Hospital, Edinburgh EH4 2XU, UK
10. East Anglian Medical Genetics Service, Clinical Genetics, Addenbrooke’s Treatment Centre, Cambridge University Hospitals NHS Foundation Trust, Cambridge CB2 0QQ, UK
11. Wellcome Trust Sanger Institute, Hinxton, Cambridge CB10 1SA, UK

Key Words: BRD4; NIPBL; Cohesin; Cornelia de Lange syndrome; Enhancer; De Novo Mutation; Super-enhancer

§These individuals had equivalent contributions to this work
These individuals are each considered to be corresponding authors for this work.
We show that the clinical phenotype associated with BRD4 haploinsufficiency overlaps with Cornelia de Lange syndrome (CdLS) – most often caused by mutation of NIPBL. More typical CdLS was observed with a de novo BRD4 missense variant, which retains the ability to co-immunoprecipitate with NIPBL but which binds poorly to acetylated histones. BRD4 and NIPBL display correlated binding at super-enhancers and appear to co-regulate developmental gene expression. (66 words)

Super-enhancers are clustered cis-regulatory elements (CRE) controlling genes important for cell type specification. Super-enhancers are molecularly defined as genomic intervals with high levels of H3K27 acetylation and binding of both BRD4 and the mediator complex. A critical role for cohesin in super-enhancer function has been recently reported. Acute depletion of cohesin resulted in disruption of higher-order chromatin structure and disordered transcription of genes predicted to be under super-enhancer control. The most widely studied human cohesinopathy is Cornelia de Lange syndrome (CdLS), a severe multisystem neurodevelopmental disorder, which is associated with a generalized deregulation of developmental genes. Typical CdLS is caused by heterozygous or mosaic loss-of-function (LOF) mutations in the gene encoding NIPBL. NIPBL is recruited to sites of double-strand breaks in DNA and functions as a transcriptional activator but it is best known for its role in loading the cohesin complex onto DNA and is required for cohesin-mediated loop extrusion and TAD formation. Causative mutations in the genes encoding the core components of the cohesin ring (SMC1A, SMC3, RAD21) and the SMC3 deacetylase HDAC8 have been identified in CdLS-like conditions. However, individuals with de novo mutations in the chromatin associated/modifying proteins ANKRD11, KMT2A and AFF4, which have no known association with cohesin can also present with CdLS-like disease.

To identify novel disease loci we studied 92 individuals with CdLS in whom no plausibly diagnostic variants could be identified in the known causative genes. In this group we identified 2/92 (2.2%) individuals with de novo mutations affecting BRD4. The first individual had a CdLS-like condition and a heterozygous 1.04 Mb deletion encompassing BRD4 (plus 28 other protein-coding genes (Family 4198; DECIPHER 281165)) (Fig. 1A, Supplemental Fig. 1&2). Targeted re-sequencing in the remaining 91 affected individuals
identified an individual, with more typical CdLS, who has a de novo missense mutation located in the second bromodomain (BD2) of BRD4 (3049; NM_058243.2 c.1289A>G, p.(Tyr430Cys))(Fig. 1B, Supplemental Fig. 5). Subsequently two affected individuals who were not part of the original cohort were identified with de novo frame-shift mutations in BRD4. The first of these was identified via ongoing screening of individuals with CdLS-like disorders (CDL038 NM_058243.2 c.1224delinsCA p.(Glu408Aspfs*4); Fig. 1B). The second indel variant was discovered through analysis of trio whole exome sequencing data generated by the Deciphering Developmental Disorders study (DDD; DECIPHER 264293) NM_058243.2 c.691del p.(Asp231Thrfs*9); Fig. 1B). The latter individual was recruited to DDD on the basis of intellectual disability, mild short stature and a ventricular septal defect but had not been suspected to have CdLS (Supplemental Table 1). On review of 7 reported heterozygous multigenic deletions encompassing BRD4 we found a significant phenotypic overlap with CdLS (Supplemental Fig. 3; Supplemental Table 1) with at least 2/7 fulfilling the established CdLS diagnostic criteria. Taken together; these data support BRD4 haploinsufficiency as the likely genetic mechanism for the CdLS-like phenotype.

It has been previously reported that mice carrying heterozygous LOF mutations in Brd4 show marked early postnatal mortality, severe prenatal onset growth failure, abnormalities of the craniofacial skeleton and reduced body fat; all features common in CdLS. Brd4 homozygous null embryos die soon after implantation. Heterozygous LOF mutations in only 12 other non-imprinted autosomal mouse genes have both postnatal lethality and postnatal growth retardation recorded as features in the Mouse Genome Informatics database (MGI); one of these being Nipbl (Supplemental Fig. 4A-C). 4 of these 13 haploinsufficient mouse genes have been implicated in super-enhancer function (Brd4, Nipbl, Chd7 and Crebbp) (Supplemental Fig. 4E).

BRD4, is a member of the bromodomain and extraterminal domain (BET) protein family with tandem bromodomains that ‘read’ acetylated lysine marks on chromatin. BRD4 binds mostly to hyper-acetylated genomic regions that encompass promoters and enhancers and BRD4 levels are particularly high at super-enhancers. BRD4 regulates transcription elongation by paused RNA polymerase II (Pol II) via mediating the release of Cdk9 activity, which results in phosphorylation of serine 2 of Pol II C-terminal domain (CTD). Tyr430, the
residue substituted in individual 3049 with the more typical CdLS phenotype (Fig. 1B), lies
within the third alpha helix (αB) of the second bromodomain (BD2) of BRD4; close to the
recognition site that mediates binding to acetylated lysine\(^{23}\). pTyr430Cys (Y430C) is a non-
conservative amino acid substitution which could plausibly impair the binding of BRD4 to
acetyl lysine. Indeed, compared to wild-type BD1 and BD2, a tagged BRD4 BD2 containing
the Y430C mutation shows reduced binding to acetylated histone peptides in vitro (Fig. 2A).
In mouse BRD4 the “human equivalent” missense variant would be p.Tyr431Cys; the
difference in amino acid numbering is the result of an “extra” proline in the poly-proline repeat
(position 215-217 in the human protein; Supplemental Fig. 6). To avoid confusion we will
use Brd4\(^{Y430C}\) as the mouse variant designation; we introduced this variant onto both alleles
\((Brd4^{Y430C/Y430C})\) of mouse embryonic stem cell (mESC) lines by Cas9-induced homology
directed repair (HDR). BRD4 immuno-precipitation (IP) in Brd4\(^{Y430C/Y430C}\) mESC shows
impaired binding to acetylated histones (H3K9ac and K3K27ac) (Fig. 2B, Supplemental Fig.
18).

Label-free quantitative (LFQ) mass spectrometry (MS) following IP was performed
using two different BRD4 antibodies on lysates from Brd4\(^{Y430C/Y430C}\) and wild-type mESCs
(Supplemental Table 4). This detected 1,082 proteins present in BRD4 IP from both cell
lines, 90 of which were absent in all IgG controls (Fig. 2C). Of these, BRD4 was the top hit
with three of the remaining 89 proteins being NIPBL, Rad21 (core cohesin ring component)
and Esco2 (SMC3 acetylase) (Fig. 2D, Supplemental Fig. 7). Other subunits of cohesin
(SMC1A, SMC3, STAG2, PDS5A, PDS5B) also showed evidence of enrichment
(Supplemental Fig. 8). The association of BRD4 with NIPBL was replicated using LFQ MS
on an independent Brd4\(^{Y430C/Y430C}\) mESC line created using the same genome editing
protocol. Reciprocal IPs using antibodies to NIPBL and SMC3 confirmed the BRD4
interaction with both NIPBL and the core cohesin ring (Fig. 2E, Supplemental Fig. 9&20). In
mESC Brd4\(^{Y430C/Y430C}\) shows a similar level of NIPBL association to wild-type Brd4,
suggesting that this interaction is unlikely to be mediated via co-binding to acetylated
chromatin (Fig. 2E, Supplemental Fig. 19).

In order to further assess the functional consequences of the BRD4 missense variant
we generated F0 mouse embryos following zygote injections of reagents to induce Cas9-
mediated HDR. As judged from digital sectioning from optical projection tomography, the
morphology of $\text{Brd}^4\text{Y}^{430C/+}$ and $\text{Brd}^4\text{Y}^{430C/430C}$ F0 mouse embryos is indistinguishable from
wild-type embryos. We also generated apparently non-mosaic F0 embryos homozygous for a
15bp in-frame deletion (NM_020508.4 c.1288_1302del; p.(Cys430_Asn434del;)
Supplemental Fig. 10); designated as $\text{Brd}^4\text{C}429\_\text{N}433\text{del}/\text{C}429\_\text{N}433\text{del}$ to maintain consistency with
human nomenclature) showing significant growth restriction at 13.5 dpc as their only obvious
phenotype (Supplemental Fig. 11). We derived mouse embryonic fibroblasts (MEF) from
13.5 dpc F0 $\text{Brd}^4\text{Y}^{430C/430C}$, $\text{Brd}^4\text{C}429\_\text{N}433\text{del}/\text{C}429\_\text{N}433\text{del}$ and control mouse embryos. The MEFs
together initially established from $\text{Brd}^4\text{C}429\_\text{N}433\text{del}/\text{C}429\_\text{N}433\text{del}$ embryos did not survive in long-term
culture, but we were able to generate sufficient cells for western blot analysis. Both wild-type
and $\text{Brd}^4\text{Y}^{430C/430C}$ MEF lines expressed comparable levels of BRD4 protein. However, in
$\text{Brd}^4\text{C}429\_\text{N}433\text{del}/\text{C}429\_\text{N}433\text{del}$ MEFs the BRD4 band was undetectable (Supplemental Fig. 12)
using an antibody raised against a peptide representing amino acid numbers 1312-1362 in
the mature peptide. The apparent null status of these cells may be the result of rapid
degradation of the abnormal protein or an artifact due to change in the epitope. The latter
may also explain the survival of these homozygous embryos past implantation.

BRD4 Chromatin IP (ChIP) from $\text{Brd}^4\text{Y}^{430C/430C}$ MEF showed reduced binding to the
promoters and super-enhancers of known BRD4 targets compared with wild-type MEF
(Figure 2F). To assess regions of common binding, we performed BRD4 ChIP-seq on wild-type mESC and compared this to publicly accessible NIPBL and BRD4 ChIP-seq data from
mouse (Supplemental Fig. 13) and human (Supplemental Fig. 14) ESCs. We used the
intersection of the BRD4-bound and NIPBL-bound genomic intervals from the ChIP-seq data
to create a set of high-confidence shared binding sites. By comparing different functional
genomic categories, mESC super-enhancers show the highest level of enrichment with
heterochromatin being the least enriched (Figure 2G&H, Supplemental Fig.15). To look for
any common functional effect on gene expression we then generated array-based
transcriptome data from control, $\text{Brd}^4\text{Y}^{430C/430C}$ and $\text{Nipbl}^{+/\text{ }}$ MEFs (Supplemental Fig. 16). Of
the >18000 genes probed on the microarray, 3049 have a transcription start site within 1Mb of
a defined MEF super-enhancer (Super Enhancer Archive). These super-enhancer-
associated genes showed significantly higher level of differential expression in both $\text{Brd}^4$ (p =
0.002) and Nipbl (p = 0.006) mutant cells compared to genes that are not super-enhancer associated. There is also a significant overlap in the specific genes that show differential expression in both BRD4 and NIPBL mutant MEFs (Supplemental Fig. 17).

CdLS can be considered a transcriptomopathy, presumed to result from loss of cohesin-dependent chromatin loops or a cohesin-independent NIPBL-mediated transcriptional activity. Our identification of de novo heterozygous loss of function mutations in BRD4 in a CdLS-like disorder, together with the functional genomic data presented above, suggests that CdLS may be more specifically defined as a disorder of super-enhancer function. Delineation of any direct or indirect physical interaction and/or functional co-dependency of BRD4 with NIPBL can now reasonably become a topic of investigation.

Acknowledgements

We thank the CdLS Foundation of UK and Ireland and particularly the families of the affected children for their time and support for the research. GO, MA, HB, WAB, PM, DRF were funded by the MRC University Unit award to the University of Edinburgh for the MRC Human Genetics Unit. AvK’s work was supported by a Carnegie Trust Research Incentive Grant 70382. The Deciphering Developmental Disorders study presents independent research commissioned by the Health Innovation Challenge Fund (grant HICF-1009-003), a parallel funding partnership between the Wellcome Trust and the Department of Health, and the Wellcome Trust Sanger Institute (grant WT098051). The views expressed in this publication are those of the authors and not necessarily those of the Wellcome Trust or the Department of Health. The study has UK Research Ethics Committee approval (10/H0305/83, granted by the Cambridge South REC, and GEN/284/12 granted by the Republic of Ireland REC). The research team acknowledges the support of the National Institute for Health Research through the Comprehensive Clinical Research Network.

Author Contributions

WAB, MMP and DRF conceived the study. DRF, WAB and MMP wrote the manuscript. All authors have read and commented on the manuscript. GO, MA, HB, NC, MMP and DDD generated the molecular biology and animal model data. AvK generated and analysed the
mass spectrometry data. FJS, EW, AR, SMP provided expert clinical interpretation and
details of the phenotype for each affected individual. AB performed the metanalysis of the
reported deletion cases. JR provided expert technical advice and cell reagents. GRG
performed the genomic and transcriptomic informatic analysis.

**URLs**

Super Enhancer Archive; http://www.bio-bigdata.com/SEA/
DECIPHER; http://decipher.sanger.ac.uk
UCSC Genome Browser; https://genome.ucsc.edu
Mouse Genome Informatics database; http://www.informatics.jax.org
Deciphering Developmental Disorders Study; https://www.ddduk.org

The authors declare that they have no competing financial interests

**Data Accessibility Statement**

Results of array-based comparative genomic hybridization from individual II:1 (family 4198)
are available on DECIPHER database (ID 281165). The DDD trio-based exome data used to
identify the de novo frame-shift mutation in individual 264293 is available from European
Genome-phenome Archive (EGA) under accession EGAD00001001848
Figure Legends:

Figure 1. *BRD4* mutations in CdLS and CdLS-like disorders. **A.** Pedigree drawing of proband with CdLS-like disorder associated with a *de novo* 1.04 Mb microdeletion of 19p (red bar), the location of which is shown on the Log$_2$ ratio plot of the array-based comparative genomic hybridization. Below on the left is a key to the symbols using in the pedigree images. To the right of this is the RefSeq gene content of the deleted region with the location of *BRD4* indicated in orange. The genes colored green are known disease genes associated with phenotypes that are not consistent with the clinical presentation in this case. Details of each is given in the supplementary notes. **B.** Pedigree drawings and facial photographs of probands with intragenic mutations of *BRD4*, with cartoon of BRD4 protein indicating the position of each of the variants in relation to the first bromodomain (BD1), the second bromodomain (BD2) and the N-terminal extra terminal domain (NET). The position of an inherited p.His304Tyr variant (orange text) reported in a single family with inherited cataracts is indicated and discussed in supplementary notes.
Figure 2. Binding of BRD4 wild-type and BRD4 p.Tyr430Cys variant to histone and non-histone proteins. 

A. Specificity of binding to acetylated histone tail peptides of wild-type BRD4 Bromodomain 1 (BRD4 BD1 WT), wild-type BRD4 Bromodomain 2 (BRD4 BD2 WT), and BRD4 p.Tyr430Cys mutant BD2 (BRD4 BD2 Y430C).  

B. Cropped immunoblot of endogenous BRD4 IPs and rabbit normal IgG (control) from Brd4 wild-type and Brd4Y430C/Y430C mESCs. Input =1% of mESC nuclear extract. Antibodies detect BRD4, H3K9ac, H3K27ac, H4K8ac and H3.  

C. Heatmap of the label-free mass spectrometry quantitative output values (average of triplicates) assigned to each protein following IP from BRD4 wild-type (WT) and BRD4-Y430C (MUT) mESC using IgG only control or Abcam/Bethyl antibody against BRD4.  

D. A plot of the log Andromeda scores assigned to the 90 proteins which are absent in the IgG controls and present in both cell lines using both BRD4 antibodies. Horizontal scatter aids visibility of each open circle and has no data correlate.  

E. Cropped immunoblot of reciprocal IP using BRD4 and NIPBL antibodies in Brd4 wild-type and Brd4Y430C/Y430C mESCs. Antibodies detect BRD4, NIPBL and SOX2.  

F. Percentage (%) input bound for BRD4 ChIP-qPCR across genomic regions in WT and BRD4 Y430C mutant MEFs (error bars = standard error of the mean from n=2 biological replicates).  

G. Forest plot of the log2 odds ratio with confidence intervals (CI) of different functional genomic categories within intersecting regions from BRD4 and NIPBL mESC ChIP.  

H. UCSC Genome Browser graphic showing colocalisation of BRD4 and NIPBL ChIPseq peaks over the super enhancer (blue bar) at Klf4 locus. H3K27ac, H3K4me1, H3K122ac and super enhancer tracks are previously published.
References:
14. Ansari, M. et al. Genetic heterogeneity in Cornelia de Lange syndrome (CdLS) and


**Online Methods Section**

**Methods**

**Patient ascertainment**

All the clinical research activity relating to this report has been in accordance with World Medical Association Declaration Of Helsinki on the Ethical Principles For Medical Research Involving Human Subjects. The research was conducted using protocols approved by UK multicenter ethics committees under the references; 04:MRE00/19 (MRC HGU) and 10/H0305/83 (DDD). Two of the affected individuals (4198 II:1 & 3049 II:1 Figure 1) are part of a larger cohort of patients with a diagnosis of CdLS or possible CdLS, referred by experienced clinical geneticists or pediatricians to the MRC Human Genetics Unit for research genetic analysis. The third (CDL038) was referred to the DNA Diagnostic Laboratory in NHS Lothian with a CdLS-like disorder. The final affected individual (264293) was identified using the trio whole exome sequence data generated by the Deciphering Developmental Disorders study.

**Array comparative genomic hybridization**

Array comparative genomic hybridization (aCGH) was performed using the Nimblegen 135k microarray platform (Roche Nimblegen) as described previously. Results were compared with the Database of Genomic Variants and polymorphic CNVs excluded.

**Droplet digital PCR**

A pair of oligonucleotide primers and the matching 5′ FAM-labelled Universal Probe Library (UPL) probe (# 25) (Roche) were designed to target coding exon 17 of the BRD4 gene using ProbeFinder software version 2.50 (Roche).

Each 20 µl ddPCR reaction consisted of 40 ng of genomic DNA, 1X ddPCR SuperMix for probes (No dUTP) (Bio-Rad Inc.), forward and reverse primers at 1 µM each, UPL probe #25 at 250 nM, and 1X 5′ VIC-labelled RNase P TaqMan Copy Number Reference assay (Thermo Fisher Scientific). Droplet generation using the QX200 droplet generator (Bio-Rad Inc.)
followed by amplification, 95\(^\circ\)C for 10 minutes, 40 cycles of 94\(^\circ\)C for 30 seconds and 57\(^\circ\)C for 323
60 seconds, and a final incubation at 95\(^\circ\)C for 10 minutes, were performed as per
manufacturer's instructions (Bio-Rad Inc.). Following completion of the PCR, plates were read
using the QX200 droplet reader (Bio-Rad Inc.). Analysis of droplet counts, amplitudes and
DNA copy number were performed with QuantaSoft software (Bio-Rad Inc.) for channel 1 =
FAM and channel 2 = VIC.

Mutation analysis by DDD Trio Exome Sequencing, Ion AmpliSeq PCR-Ion PGM, and
Sanger sequencing

As part of a DDD Complementary Analysis Protocol #35 VCF files on the first 4293 trios with
whole exome sequence were searched for candidate de novo mutations in *BRD4*. Only one
possible *de novo* disruptive variant was identified in *BRD4*. This variant was validated as de
novo using the approach mentioned below. No other plausible cause for the developmental
disorder was apparent on trio based whole exome analysis.

An AmpliSeq panel encompassing the coding exons of *BRD4* and nine other candidate genes
was designed using the Ion AmpliSeq Designer tool (Life Technologies, IAD41056). Library
preparation and sequencing on the Ion PGM platform (Life Technologies), followed by
sequence alignment and variant calling on software NextGENe version 2.3.3 (Soft Genetics)
were performed as described previously\(^2\). A total of 92 individuals were screened, who had
previously scored as negative for mutations in *NIPBL*, *SMC1A*, *SMC3*, *HDAC8* and *RAD21*,
and large-scale genomic deletions/duplications. The same panel also applied to subsequent
clinical referrals to the NHS DNA diagnostic laboratory in Edinburgh was used to identify one
further *de novo* heterozygous loss of function mutation in an individual who had a CdLS-like
phenotype.

Any significant variants were confirmed by Sanger sequencing and analysed using Mutation
Surveyor software version 3.30, as described previously\(^2\). The *BRD4* sequence identifier,
NC_000019.10 was used in the analysis. Sequence variant nomenclature is reported
according to the *BRD4* transcript variant, NM_058243. Primer sequences and PCR conditions
are available upon request.
Plasmids, expression and purification of proteins

Human BRD4 BD1 and BD2 plasmids were kindly gifted by Prof Stefan Knapp (Nuffield Department of Clinical Medicine, Oxford). Proteins were expressed and purified at the Edinburgh Protein Production Facility (EPPF) as described previously.22

Site-directed Mutagenesis

The point mutation c.1289A>G, predicted to result in the protein variant p.Tyr430Cys (Y430C) was introduced into the BRD4-BD2 and FLAG-mBRD4 constructs using the QuikChange II XL Site-directed Mutagenesis kit (Agilent Technologies) following the manufacturer's instructions. The presence of the desired mutations was confirmed by Sanger sequencing.

CRISPR/Cas9 construct design

Guide RNA (gRNAs) 1 and 2 were designed across p.Tyr430 using online tool DNA 2.0. The wild-type and mutant repair templates (chr17:32,220,150-32,220,271; GRCm38) were synthesized by IDT as 122 bp Ultramer ssODN bearing the desired sequence change. For genome editing in mouse embryonic stem cells (mESCs) gRNAs 1 and 2 were cloned into PX461 (Addgene plasmid #48140) and PX462 (Addgene plasmid #62987) respectively. For genome editing in mouse embryos both gRNAs were cloned into PX461 and the full gRNA template sequence was amplified from the resulting PX461 clone using universal reverse primer and T7 tagged forward primers. The gRNA PCR template was used for in vitro RNA synthesis using T7 RNA polymerase (NEB), and the RNA template subsequently purified using RNeasy mini kit (Qiagen) purification columns. Cas9n mRNA was procured from Tebu Bioscience.

Genome editing in mouse embryonic stem cells (mESCs)

To generate mESCs carrying the p.Tyr430Cys missense variant in BRD4, 46C cells were cotransfected with gRNAs 1 and 2 (0.5 µg/ml) and the mutant repair template (0.5µg/2ml) using Lipofectamine® 3000 Transfection Reagent (ThermoFisher) as per the manufacturer's
instructions. After 48 hours, successfully transfected cells were selected for: firstly by puromycin treatment, and subsequently by FACS based on GFP expression. Resulting GFP and puromycin positive cells were plated at 500 cells/10cm$^2$. After 1 week, colonies were picked and plated in duplicate as 1 colony/well of a 96 well plate. Genomic DNA was extracted from the colonies and sequenced by Sanger sequencing. Wild-type clones and clones homozygous for the p.Tyr430Cys variant were expanded and frozen for later use.

**Genome editing in mouse embryos and generation of mouse embryonic fibroblast (MEFs)**

To generate mouse embryos carrying the p.Tyr430Cys variant in BRD4, injections were performed in single cell mouse zygotes. Injection mix contained Cas9 mRNA (50 ng/µl), gRNAs 1 and 2 (25 ng/µl) and each repair template DNA (75 ng/µl). The embryos were later harvested for analysis at 13.5 dpc stage of embryonic development. MEFs were isolated from limbs of individual E13.5 embryos by mincing in 1 ml of medium (DMEM, 10% FCS, 50 U/ml penicillin and 50 mg/ml streptomycin,). Resulting suspensions were grown at 37°C, 5% CO$_2$ and 3% O$_2$, and non-adherent cells removed after 24 hours. MEFs from embryos with unedited $\text{Brd4}$ alleles, clean homozygous knock-in for p.Tyr430Cys in $\text{Brd4}$ ($\text{Brd4}^{\text{Tyr430Cys/Tyr430Cys}}$) and homozygous knock-in for an in-frame deletion ($\text{Brd4}^{\text{Cys430_Asn434del/Cys430_Asn434del}}$) alleles were used for further experimentation.

**Generation of heterozygous loss-of-function Nipbl MEFs:**

Mice with $\text{Nipbl}$ floxed allele (a kind gift from Heiko Peters, University of Newcastle) were crossed with Cre745 mice (a kind gift from DJ Kleinjan, University of Edinburgh), containing a CAGGS-Cre construct in which Cre recombinase is under control of a chicken b-actin promoter to excise $\text{Nipbl}$ exon 1. Embryos were collected at 13.5 dpc. MEFs were isolated from heterozygous $\text{Nipbl}$ knockout embryo limbs by mincing in 1 ml of medium (DMEM, 10% FCS, 50 U/ml penicillin and 50 mg/ml streptomycin,). Resulting suspensions were grown at 37°C, 5% CO$_2$ and 3% O$_2$, and non-adherent cells removed after 24 hours.
Histone tail peptide arrays

A modified histone peptide array (Active motif, #13005) experiment was performed as described previously\textsuperscript{23}. Briefly, the array was blocked in TBST buffer (10 mM Tris/HCl pH 8.3, 0.05% Tween-20, 150 mM NaCl) containing 5% non-fat dried milk at 4°C overnight. The membrane was washed with TBST for 5 min, and incubated with 10 nM purified His-tagged BRD4 BD1 or wild-type (WT) and p.Tyr430Cys (Y430C) BD2 domains, at room temperature (RT) for 1 hour in interaction buffer (100 mM (0.5 µg/3 ml) KCl, 20 mM HEPES pH 7.5, 1 mM EDTA, 0.1 mM DTT, 10% glycerol). After washing in TBST, the membrane was incubated with mouse α-His (Sigma, H1029, 1:2,000 dilution in TBST) for 1 hour at RT. The membrane was then washed 3 times with TBST for 10 min each at RT, and incubated with horseradish peroxidase conjugated α-mouse antibody (1:10,000 in TBST) for 1 hour at RT. The membrane was submerged in ECL developing solution (Pierce, #32209), imaged (Imagequant, GE Healthcare) and the data quantified using array analyzer software (Active motif).

Nuclear extract co-immunoprecipitation

30 x 10\textsuperscript{6} wild-type and p.Tyr430Cys (Y430C) BRD4 mESCs were trypsinised, pelleted and resuspended in 5 ml ice-cold swelling buffer (10 mM Hepes, pH 7.9, 1.5 mM MgCl\textsubscript{2}, 10 mM KCl, 0.5mM DTT, Complete Mini EDTA-free protease inhibitor (Roche)) for 5 minutes on ice. Nuclei were pelleted by centrifugation at 2,000 rpm for 5 minutes at 4°C. The resulting nuclear pellets were sonicated in 2 ml RIPA buffer (50 mM Tris pH 7.5, 150 mM NaCl, 1% NP-40, 0.5% sodium deoxycholate, benzonase and Complete Mini EDTA-free protease inhibitor (Roche)), using a Bioruptor\textsuperscript{®} Plus sonication device (Diagenode) at 4°C, 30 seconds on, 30 seconds off. It was noted that prolonged (1 hour) exposure to the detergents in RIPA buffer affected the interactions of BRD4 as measured by mass spectrometry. Nuclear extracts were cleared by centrifugation at 13,000 rpm for 10 minutes at 4°C. Protein A Dynabeads (life technologies) were blocked prior to antibody coupling by washing 3 times with 5% BSA in PBS. Antibodies were coupled to the beads at 5 mg/ml by rotation for 1 hour at 4°C. Equivalent nuclear protein amounts were incubated with antibody coupled beads for 1 hour at 4°C. Beads were washed and pulled down proteins analysed by mass spectrometry or
western blot. Antibodies used: BRD4 (Bethyl A301-985A100), SMC3 (Bethyl 0300-060A), NIPBL (Bethyl A301-779A) and normal rabbit IgG (Santa-Cruz, sc-2027).

**Western blots**

For western blot analysis beads were washed 5 times with RIPA buffer, bound proteins eluted by boiling in 1X NuPage LDS buffer (ThermoFisher Scientific) with 1X NuPage reducing agent (ThermoFisher Scientific) for 5 minutes and separated on a 3-8% tris-acetate gel (reciprocal BRD4/SMC3/NIPBL IPs and MEF cell lysates) or 4-12% bis-tris gel (BRD4 IPs for acetylated histone binding) (ThermoFisher Scientific). Following electrophoresis, proteins were transferred to nitrocellulose membranes (ThermoFisher Scientific) using iBlot 2 Dry Blotting System (ThermoFisher Scientific) for 7 minutes (when probing for proteins <250 kDa only) or to PVDF membranes by wet transfer for 90 minutes (when probing for proteins >250 kDa) and incubated with primary antibodies overnight at 4°C. Membranes were washed 3 times in TBST and probed with HRP-conjugated secondary antibody (anti-Rb/anti-goat, 1:10,000) for 1 hour at RT. After 3 more washes in TBST, membranes were incubated with Pierce™ ECL Western Blotting Substrate (ThermoFisher Scientific) for 5 minutes and imaged using (Image-quant, GE Healthcare). Antibodies used: BRD4 (Bethyl A301-985A100, 1:3,000), SMC3 (Bethyl 0300-060A, 1:1,000), H3K27ac (Genetex GTX128944, 1:1,000), H4K8ac (Abcam ab15823, 1:1,000), H3K9ac (Abcam, ab10812, 1:500), H3 (Abcam, ab1791, 1:5,000), NIPBL (Bethyl A301-779A, 1:1000), SOX2 (Abcam ab97959, 1:1000) Actin-b (Abcam ab8229, 1:500).

**Mass spectrometry**

For analysis by mass spectrometry, beads were washed 3 times with Tris-saline buffer, and excess buffer removed. Immunoprecipitations were digested on beads, desalted and analysed on a Q-Exactive plus mass spectrometer as previously described\(^2\). Proteins were identified and quantified by MaxLFQ\(^2\) by searching with the MaxQuant version 1.5 against the Mouse proteome data base (Uniprot). Modifications included C Carbamlylation (fixed) and M oxidation (variable). Bioinformatic analysis was performed with the Perseus software suite.
Chromatin immunoprecipitation-quantitative PCR

Primary MEFs isolated from 13.5 dpc embryos were cultured for 3-4 passages in DMEM media supplemented with 15% FCS, 1% Pen/Strep, L-Glutamine, non-essential amino acids and Sodium pyruvate. Cells were harvested by trypsinizing and fixed immediately with 1% formaldehyde (Thermo Fisher Cat. 28906) (25°C, 10 min) in PBS, and stopped with 0.125 M Glycine. Chromatin immunoprecipitation (ChIP) was performed as described previously\textsuperscript{26}. Briefly, cross linked cells were re-suspended in Farnham lysis buffer (5 mM PIPES pH 8.0, 85 mM KCl, 0.5% NP-40, Complete Mini EDTA-free protease inhibitor (Roche)) for 30 minutes and centrifuged at 228 g for 5 minutes at 4°C. Nuclei were resuspended in RIPA buffer (1X PBS, 1% NP-40, 0.5% sodium deoxycholate, 0.1% SDS (filtered 0.2 -0.45 micron filter unit) + Complete Mini EDTA-free protease inhibitor (Roche)) and sonicated using a Bioruptor\textsuperscript{®} Plus sonication device (Diagenode) at full power for 40 minutes (30 seconds on, 30 seconds off) to produce fragments of 100-500 bp. 3 µg of each antibody was incubated with Protein A Dynabeads (ThermoFisher Scientific, 10001D) in 5 mg/ml BSA in PBS on a rotating platform at 4°C for two hours. An arbitrary concentration of 50 µg chromatin was incubated with antibody bound Dynabeads in a rotating platform at 4°C for 16 hours. Beads were washed 5 times (5 minutes each) on a rotating platform with cold LiCl wash buffer (100 mM Tris pH 7.5, 500 mM LiCl, 1% NP-40, 1% Sodium deoxycholate) and one time with RT TE buffer. ChIP complexes were eluted with elution buffer (1% SDS, 0.1 M NaHCO$_3$) and Input and ChIP samples were incubated at 65°C for 5 hours to reverse the crosslinks. 2 µl of RNase A (20 mg/ml) was added and samples were incubated at 37°C for 1 hour before 2 µl of Proteinase K was added and samples were incubated for 2 hours at 55°C. DNA was purified using QIAquick PCR Purification Kit (Qiagen, Cat. 28104), and analysed by qPCR for primers described in Supplementary Table 2. Antibodies used: rabbit IgG (Santa Cruz sc-2025), BRD4 (Bethyl A301-985A100).

Chromatin immunoprecipitation-sequencing (ChIP-Seq)

Wild-type mESCs were cultured in GMEM media supplemented with 10% FCS, 1% Pen/strep, L-glutamine, non-essential amino acids, sodium pyruvate and 1,000 U/ml
LIF. Cells were harvested by trypsinizing and fixed immediately with 1% formaldehyde (Thermo Fisher Cat. 28906) (25°C, 10 minutes) in PBS. This reaction was quenched with 0.125 M Glycine. ChIP was carried out as above. After purification, DNA was eluted in 20 µl and libraries were prepared for ChIP and input samples as previously described\textsuperscript{27}. Samples were sequenced at BGI (Hong Kong; 50-base single-end reads) using the HiSeq 4000 system (Illumina).

**Transcriptome analysis**

RNA was extracted from Brd4\textsuperscript{Tyr430Cys/Tyr430Cys} and Nipbl heterozygous null MEFs using the RNeasy Mini Kit (Qiagen) as per the manufacturer’s instructions. 1 µg of RNA was hybridised to a SurePrint G3 Mouse GE 8x60K microarray (Agilent; G4852A) and scanned on a Nimblegen scanner as described previously\textsuperscript{28}.

Quantile normalisation and background correction (method normexp) of the microarray data was carried out using the bioconductor package limma\textsuperscript{29}. Gene level expression was calculated by averaging the probes signal that mapped to the same gene (Gene Symbols mapped to Probe identifiers obtained from GEO, GPL13912, Agilent-028005 SurePrint G3 Mouse GE 8x60K Microarray). The normalised signal for technical replicate samples was averaged prior to Differential expression (DE) analysis.

**Transcriptome analysis statistics**

Gene level differential expression was conducted using the bioconductor package limma\textsuperscript{29}. Briefly, a linear model was fitted to each gene. Then empirical Bayes moderation was applied to the linear model fit to compute moderated t-statistics, moderated F statistic, and log-odds of differential expression. The Benjamini & Hochberg method was used to correct the p-values for multiple testing. Genes were identified as significantly differentially expressed if the FDR q value < 0.1.

To test whether genes (n= 3049/19113) with a transcription start site within 1Mb of MEF Super Enhancers (SE) are more highly ranked relative to other expressed genes in terms
differential expression (t-statistic) we performed a Mean-rank Gene Set Test (geneSetTest, bioconductor package limma).

A hypergeometric test was performed on the differentially expressed gene sets for Brd4 and Nipbl to determine if DE genes were significantly enriched between the two groups.

**ChIP-seq analysis**

Bowtie 2 (version 2.2.6) was used to map reads to mouse (mm9) and human (hg19) genomes (options bowtie2-align-s --wrapper basic-0)^30. To calculate the correlation of NIPBL and BRD4 with other histone modifications (see Supplementary Table 3), the correlation of the ChIP-seq binding profiles across the genome was calculated. DeepTools (version 2.3.5) multiBamSummary was used to calculate the coverage of mapped reads in 150 bp sequential bins across the mm9/hg19 genome (options --binSize 150bp, --ignoreDuplicates,--black, ListFileName,--extendReads 150,--mappingquality 30)^31. Genomic bins within Blacklisted regions and chrX and chrY were excluded from the analysis. Genomic bins were also restricted to regions of open chromatin using DNase I hypersensitive sites identified by the ENCODE project^32 (see Supplementary Table 3).

The genome wide coverage matrix was imported into R and Pearson's R was calculated. Correlation scores were visualised as a heatmap using the R package pheatmap (options; euclidean distance and complete clustering method)^33.

**Peak calling**

To call BRD4 and NIPBL bound regions we used the MACS peak caller (2.1.1). For BRD4 peaks we used the parameters broadPeaks and an FDR cut-off of 0.1.

For NIPBL we used the public ChIP-Seq dataset NIPBL in V6.5 (C57BL/6-129) murine ES cells (GEO ID, GSM560350) and, the accompanying whole cell extract dataset (GEO ID, GSM56035) as background.
For peak calling we used MACS with the parameters narrowPeaks and an FDR cut off of 0.1.

To perform intersections on genomic ranges, such as peaks regions, we used bedtools intersect (2.26.0)\textsuperscript{34}.

Any peaks that intersected with the mm9 genome blacklist regions or mapped to non-canonical chromosomes were removed from subsequent analysis.

**Genomic region enrichment**

To determine the preference of co-localised NIPBL and BRD4 binding to specific chromatin states we performed fisher enrichment analysis on a chromatin state map in mouse embryonic stem cells (ChromHMM, mm9). This state map has annotated the genome into six major chromatin states including: active promoter, poised promoter, strong enhancer, poised or weak enhancer, insulator, repressed, transcribed and heterochromatin.

In addition we looked at enrichment of co-localised NIPBL and BRD4 binding sites with Super Enhancers (SE) regions found in mESC line E14 using data from SEA: Super-Enhancer Archive\textsuperscript{35}.

We used the consensus set of SE regions from two mES E14 replicates to define SE regions.

**Genomic region enrichment statistics**

To calculate enrichment of peaks we used bedtools (2.26.0) fisher test. The fishers odds ratio was converted to Log2 scale and plotted using R forest plot package.

**URLs**

Database of Genomic Variants; [http://dgv.tcag.ca/dgv/app/home](http://dgv.tcag.ca/dgv/app/home)

Ion AmpliSeq Designer tool; [http://www.ampliseq.com](http://www.ampliseq.com)

Blacklisted regions; [https://sites.google.com/site/anshulkundaje/projects/blacklists](https://sites.google.com/site/anshulkundaje/projects/blacklists)

ChromHMM, mm9; [https://github.com/gireeshkbogu/chromatin_states_chromHMM_mm9](https://github.com/gireeshkbogu/chromatin_states_chromHMM_mm9)


pheatmap: Pretty Heatmaps (2015) v1.0.8. [https://CRAN.R-project.org/package=pheatmap](https://CRAN.R-project.org/package=pheatmap)
578 References

34. Ramírez, F et al., Nucleic Acids Res. 44(W1), W160-5 (2016)
37. Wei, Y. et al., Nucleic acids research. 44(D1), D172-D179 (2016)