Mice deficient of Myc super-enhancer region reveal differential control mechanism between normal and pathological growth

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Abstract The gene desert upstream of the MYC oncogene on chromosome 8q24 contains susceptibility loci for several major forms of human cancer. The region shows high conservation between human and mouse and contains multiple MYC enhancers that are activated in tumor cells. However, the role of this region in normal development has not been addressed. Here we show that a 538 kb deletion of the entire MYC upstream super-enhancer region in mice results in 50% to 80% decrease in Myc expression in multiple tissues. The mice are viable and show no overt phenotype. However, they are resistant to tumorigenesis, and most normal cells isolated from them grow slowly in culture. These results reveal that only cells whose MYC activity is increased by serum or oncogenic driver mutations depend on the 8q24 super-enhancer region, and indicate that targeting the activity of this element is a promising strategy of cancer chemoprevention and therapy.

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Introduction

Deregulated expression of the MYC oncogene is associated with many cancer types (Reviewed in Albihn et al., 2010; Dang, 2012; Evan, 2012). MYC acts primarily as a transcriptional activator that increases expression of many genes required for RNA and protein synthesis above the level that is required in resting cells. In cancer cells, aberrantly elevated levels of MYC drive global amplification of transcription rates, providing the cells with necessary resources for rapid proliferation (see, for example Brown et al., 2008; van Riggelen et al., 2010; Ji et al., 2011; Lin et al., 2012; Sabò et al., 2014; Walz et al., 2014).

Transcription of the MYC gene is regulated by a diverse array of regulatory elements located both upstream and downstream of the MYC transcription start site (TSS). Variants in the MYC upstream region contribute to inherited susceptibility to most major forms of human cancer, and account for a very large number of cancer cases at the population level (Amundadottir et al., 2006; Gudmundsson et al., 2007; Yeager et al., 2007; Al Olama et al., 2009; Yeager et al., 2009). For example, the polymorphism rs6983267 linked to colorectal (Tomlinson et al., 2007) and prostate (Yeager et al., 2007) cancers contributes more to cancer morbidity and mortality than any other
known inherited variant or mutation, including the inherited mutations in classic tumor suppressors such as \( \text{RB} \), \( \text{TP53} \) and \( \text{APC} \). Through computational and experimental analyses, we and others have shown that the risk allele G of rs6983267 creates a strong binding site for the colorectal-cancer associated transcription factor Tcf7l2 (Pomerantz et al., 2009; Tuupanen et al., 2009). This binding site is located within the \( \text{Myc-335} \) enhancer element that is dispensable for mouse viability, but required for efficient Tcf7l2-driven intestinal tumorigenesis (Sur et al., 2012b).

More recently, another enhancer element, located 1.47 Mb downstream of \( \text{Myc} \) was shown to be required for formation of acute lymphoblastic leukemia (ALL) in mice (Herranz et al., 2014). However, in contrast to the \( \text{Myc-335} \) element, this element is also required for normal T-cell development. Thus, the mechanism by which individual \( \text{Myc} \) enhancer elements contribute to normal development and tumorigenesis is still unclear.

Several studies have shown that the 8q24 region contains a large number of additional enhancer elements (see, for example [Hallikas et al., 2006; Ahmadiyeh et al., 2010; Yan et al., 2013; Yao et al., 2014]) and super-enhancers that are active in many different types of human cancer (Hnisz et al., 2013; Lovén et al., 2013; Zhou et al., 2015). The \( \text{MYC} \)-associated super-enhancers are activated during the process of tumorigenesis (Hnisz et al., 2013), and downregulation of super-enhancer activity leads to selective inhibition of \( \text{MYC} \) expression (Lovén et al., 2013). Thus, \( \text{MYC} \)-associated super-enhancer activity is required for tumorigenesis, but the role of these elements in normal tissue morphogenesis and homeostasis has been unclear.

To address this problem, we have in this work generated multiple mouse strains deficient of regulatory elements upstream of the \( \text{Myc} \) promoter. Since this region contains multiple tumor type and tissue-specific enhancers and super-enhancers, for the sake of clarity we refer to the deleted region here as the ‘super-enhancer region’. By analysis of the mice, we found that the entire super-enhancer
region conferring multi-cancer susceptibility contributes to MYC expression in vivo, yet is not required for mouse embryonic development and viability. However, this region is required for the growth of normal cells in culture and cancer cells in vivo. As cultured cells are exposed to serum, which is a signal of tissue damage, this finding suggests that tumor cells and cells responding to damage signals share regulatory mechanisms that are dispensable for normal physiological growth control.

Results

Functional mapping of the super-enhancer region upstream of Myc
To dissect functional significance of the 8q24 region during normal development, we generated series of Myc alleles in mice using homologous recombination in ES cells. These include the Myc-335 enhancer deletion allele we have described previously (Sur et al., 2012b), and deletions of two additional conserved enhancer elements, Myc-196 and Myc-540, both of which are active in mouse intestine and colorectal cancer cells. In addition, we generated a point mutation that inactivates a conserved CCCTC-Binding factor (CTCF) site 2 kb upstream of the Myc TSS. This site has previously been reported to be required for Myc expression (Gombert and Krumm, 2009), and to have insulator activity (Gombert et al., 2003) (Figure 1a). Each allele contained loxP site(s) in the same orientation to allow conditional knockouts of the enhancers, and to facilitate generation of large deletions and duplications by interallelic recombination (Wu et al., 2007). All alleles were bred to homozygosity, and resulted in generation of viable mice. Expression of Myc in the colon of Myc-196−/− and Myc-540−/− mice was not markedly altered, suggesting that these elements have little effect on regulation of Myc in the intestine under normal laboratory conditions (Figure 1b). Myc expression level was also normal in Myc-CTCFmut/mut mouse colon despite loss of CTCF and cohesin (Rad21) binding to the region proximal to the Myc promoter (Figure 1c).

Mice lacking the Myc super-enhancer region are viable and fertile
As the individual mutations and deletions had limited effect, we next decided to generate two large deletions in the Myc locus using interallelic recombination between the Myc-CTCFmut/loxP site and the loxP sites at Myc-335− and Myc-540−, yielding deletions of 365 kb (GRCm38/mm10 chr15:61618287-61983375) and 538 kb (chr15:61445326-61983375), respectively (Figure 2a). The resulting alleles, MycΔ2-367 and MycΔ2-540, were then segregated out from the corresponding duplications, and bred to homozygosity. Given the very large regions that were deleted (Figure 2b), we expected to see a strong phenotype. However, no overt phenotype was identified in the MycΔ2-367/Δ2-367 mice. The mice were born at the expected mendelian ratio, and both males and females were viable and fertile. Analysis of Myc expression, however, revealed a strong decrease in Myc expression in the colon and ileum of the mice (not shown).

The larger deletion, MycΔ2-540, could also be bred to homozygosity, and both males and females were viable. Given that the entire Myc regulatory region spans more than 2 Mb of DNA and is located on both sides of the Myc coding region (Rosenbloom et al., 2013; Sloan et al., 2016), the deletion is not expected to be equivalent to deletion of the Myc gene itself. Still, the viability of the mice is striking, since the region deleted contains regions linked to risk for myeloma, chronic lymphocytic leukemia and pancreatic, thyroid, bladder, prostate, breast, and colon cancers (Chung and Chanock, 2011; Sahasrabudhe et al., 2015; Mitchell et al., 2016; Zhang et al., 2016). To characterize the mice further, we analyzed histology and MYC expression in the tissues where these tumors originate from. This analysis revealed normal morphology of mammary gland, spleen, bladder, prostate and colon in MycΔ2-540/Δ2-540 mice (Figure 2c).

Loss of the super-enhancer region leads to tissue-specific changes in Myc expression
Although the MycΔ2-540/Δ2-540 mice exhibited a normal phenotype, Myc expression was altered in a tissue-specific manner in these mice. This is expected since this region contains individual tissue specific regulatory elements. The expression of Myc was strongly decreased in colon, small intestine and prostate of these mice (Figure 3a and not shown). Immunohistochemical analysis of MYC expression in intestine revealed strong decrease of nuclear staining, and loss of MYC expression
**Figure 1.** Cancer susceptibility region upstream of Myc contains several conserved enhancer elements that are dispensable for normal mouse development and MYC expression. (a) Comparison of Myc locus between human and mouse. The susceptibility regions for prostate cancer (PrCa), chronic lymphocytic leukemia (CLL), breast cancer (BrCa), colorectal cancer (CRC) and bladder cancer (BlCa) are marked. Red vertical lines mark the location of the Tcf7l2-binding CRC Myc enhancers in the two species. The lower panel shows the regional conservation probability predicted by PhastCons (hg19 assembly, UCSC) with non-overlapping sliding windows for the whole region and each enhancer locus with a size of 500 bp and 10 bp, respectively. (b) Deletion of Myc-196 and Myc-540 enhancer elements does not affect Myc expression in the colon as determined by qPCR analysis (Myc-196/−/− n = 2, Myc-540/−/− n = 3 and wild-type n = 5). See Figure 1—source data 1 for details. (c) Mutation of the Myc-CTCF site causes loss of CTCF and Rad21 binding at the Myc locus (top panel). Binding of CTCF and Rad21 at a control Actb locus is not affected. Red and black arrowheads denote binding sites at Myc and Actb loci, respectively; green: Myc-CTCFmut/mut, blue: wild-type. The gene body for Myc and Actb is shown below the respective panels. The qPCR analysis reveals that despite loss of CTCF/cohesin binding, the expression of Myc mRNA is not altered in the colon (for qPCR, Myc-CTCFmut/mut n = 4, wild-type n = 3). See Figure 1—source data 1 for details. Error bars denote one standard deviation.

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The following source data is available for figure 1:

**Source data 1.** Myc transcript levels in wild-type and mutant mice in Figure 1b-c.

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Figure 2. Deletion of the 8q24 super-enhancer region is well tolerated during normal development and homeostasis. (a) Schematic representation of the 365 kb and 538 kb deletions. (b) MycΔ2-540/Δ2-540 deletion removes a region containing several active enhancer elements upstream of Myc as shown by ChIP-seq analysis of histone H3 lysine 27 acetylation (H3K27ac) and lysine four trimethylation (H3K4me3). The deletion also removes several Tcf7l2 ChIP-seq peaks. Signal from MycΔ2-540/Δ2-540 and wild-type mice are shown in green and blue, respectively. Red arrowheads and horizontal lines mark Figure 2 continued on next page
the different enhancer positions. (c) Haematoxylin/ Eosin stained sections of spleen, bladder, prostate, colon (Bar = 100 μm) and Carmine Alum stained whole mounts of mammary glands, Bars = 3 mm, 100 μm (inset) showing normal development and homeostasis of different organs in MycΔ2-540Δ2-540 mice. (d) MycΔ2-540Δ2-540 mice have a reduced number of B-cells compared to the wild-types. Left panel: FACS plots of a representative MycΔ2-540Δ2-540 and wild-type mouse spleen showing B-cell (B) population. Right panel: Scatter dot plot of total number of B cells in the spleen and bone marrow of wild-type (squares), n = 5 and MycΔ2-540Δ2-540 (filled circles), n = 5. Each point represents individual mouse. Line represents the median. See Figure 2—source data 1 for details. The number of CD4+ and CD8+ T-cells is not affected by the deletion (see Figure 2—figure supplement 1 and appendix 1).

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The following source data and figure supplements are available for figure 2:

Source data 1. B cell numbers in the wild-type and MycΔ2-540Δ2-540 mice in Figure 2d.
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Figure supplement 1. The loss of the Myc super-enhancer region results in a decrease in the number of B-cells, but no major defects in hematopoiesis.
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Figure supplement 1—source data 1. B and T-cell populations in the wild-type and MycΔ2-540Δ2-540 mice in Figure 2—figure supplement 1a.
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from the transit amplifying cell compartment. However, expression of MYC was still detected at the base of the crypt in the region where the intestinal stem cells are known to reside (Figure 3b). These results are consistent with the role of the deleted region in tumorigenesis of colon and prostate. To analyze the effect of decreased MYC expression on the proliferation in the transit amplifying

Figure 3. Tissue-specific effect of MycΔ2-540Δ2-540 deletion on MYC expression. (a) qPCR data showing the percentage of Myc expression in MycΔ2-540Δ2-540 relative to the wild-type in colon (Co) n = 4, prostate (Pr) n = 2, bladder (Bl) n = 5, spleen (Sp) n = 4 and mammary gland (Ma) n = 3. Red line marks the expression level (100%) in wild-type mice. Error bars indicate one standard deviation. See Figure 3—source data 1 for details. (b) Immunohistochemistry shows reduced expression of MYC (n = 3 for each genotype) protein in intestinal crypts of MycΔ2-540Δ2-540 mice without any significant effect on proliferation as indicated by Ki-67 (n = 2 for each genotype) immunostaining, Bar = 10 μm. Brown: IHC staining, Blue: Haematoxylin staining.

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The following source data is available for figure 3:

Source data 1. Myc transcript levels in MycΔ2-540Δ2-540 mice relative to the wild-types in Figure 3a.
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compartment, we performed immunohistochemistry (IHC) for the proliferation marker Ki-67. Both the wild-type and \textit{Myc}^{\Delta 2-540/\Delta 2-540} had similar proliferative activity in the intestinal crypts (Figure 3b).

In contrast to colon and prostate, \textit{Myc} expression was not markedly affected in the bladder, and was elevated in the spleen (Figure 3a). To analyze the cellular composition of the spleen, we performed flow cytometric analysis of markers for hematopoietic stem cells and lymphoid lineage cells. \textit{Myc}^{\Delta 2-540/\Delta 2-540} mice had a near normal hematopoietic compartment (Figure 2d). The only observed difference was a small reduction of B cells in the \textit{Myc}^{\Delta 2-540/\Delta 2-540} mice compared to the wild-type mice both in the spleen and the bone marrow. In contrast to the decrease in B-cells, the T cell numbers were not affected by the deletion (Figure 2—figure supplement 1a). This finding is consistent with the published data that regulatory elements controlling T-cell development and T-cell acute lymphoblastic leukemia are located 1.47 Mb downstream of the \textit{Myc} ORF (Herranz et al., 2014).

To identify regulatory elements that could explain the effect in B-cells, we performed ChIP-seq analysis of chromatin from LSK-Flt3L cells and mature B-cells isolated from wild-type mice. This analysis identified two B-cell specific regulatory elements. The Myc 2–540 deletion results in loss of one of the elements, and moves the other element very close to the Myc TSS (Figure 2—figure supplement 1b). Although the exact regulatory mechanism is not clear and requires further study, the above data is consistent with a role of the super-enhancer region in development of chronic lymphocytic leukemia, which is primarily a B-cell malignancy. However, the decrease in B-cell number does not affect viability, and the \textit{Myc}^{\Delta 2-540/\Delta 2-540} mice are healthy and do not display an immune-deficient phenotype under normal ‘clean’ mouse housing conditions in the absence of known pathogenic microorganisms.

To compare the role of the 8q24 super-enhancer region in growth of cells in vivo and in cell culture, we isolated fibroblasts from the skin of adult \textit{Myc}^{\Delta 2-540/\Delta 2-540} and wild-type mice. Based on presence of active histone marks, and undermethylation of focal elements, the super-enhancer region is active in fibroblasts from both humans and mice (Figure 4a and Figure 4—figure supplement 1). However, the resident fibroblasts in the skin of \textit{Myc}^{\Delta 2-540/\Delta 2-540} mice appeared normal as judged by Vimentin expression (Figure 4b). Ki-67 staining (IHC) of skin sections showed comparable proliferation levels in wild-type and \textit{Myc}^{\Delta 2-540/\Delta 2-540} mice (Figure 4b). In contrast, most lines of fibroblasts (6 out of 7) isolated from \textit{Myc}^{\Delta 2-540/\Delta 2-540} mice grew slower in culture compared to fibroblasts isolated from wild-type mice (Figure 4c; p-value=0.0256, Mann-Whitney one tailed test).

**Deletion of the \textit{Myc} super-enhancer region affects MYC target gene expression only under culture conditions**

To understand the mechanism by which the deletion of the 8q24 super-enhancer region has a differential effect on growth during normal tissue homeostasis and growth under culture conditions, we subjected both the mouse tissues and cultured cells to RNA-seq analysis. Analysis of mouse tissues confirmed the changes in \textit{Myc} expression observed by qPCR (Figure 5a and Figure 5—figure supplement 1). Surprisingly, despite more than 80% decrease of \textit{Myc} expression in the colon, very few genes were downregulated in the tissues, and none of the significantly altered genes were known \textit{MYC} targets (Supplementary file 1). These results suggest that expression of canonical \textit{MYC} target genes is not sensitive to decreases in \textit{MYC} protein level during normal tissue homeostasis. In contrast to the in vivo situation, where \textit{Myc} is downregulated but key target genes are not affected, in cultured \textit{Myc}^{\Delta 2-540/\Delta 2-540} fibroblasts that grew slowly in culture, the downregulation of \textit{Myc} lead to a loss of expression of key target genes that drive cell growth and division. Upstream regulator analysis performed using Ingenuity Pathway Analysis revealed that the highest-ranked potential regulator for the identified gene set was \textit{MYC} (Figure 5b).

Measured by FPKM values, the cultured wild-type fibroblasts had higher \textit{Myc} mRNA levels than normal tissues, whereas the cultured null fibroblasts had \textit{Myc} levels that were comparable to or lower than those of normal wild-type tissues. The elevated \textit{Myc} levels in cultured cells are caused by serum stimulation, as \textit{Myc} mRNA levels are low in serum-starved fibroblasts and strongly induced by serum (Ref. [Dean et al., 1986] and our unpublished data). These results indicate that the 8q24 super-enhancer region is dispensable for normal tissue homeostasis under conditions where \textit{MYC} activity is relatively low. However, the region is required for induction of \textit{MYC} activity to levels that
are high enough to drive the expression of MYC target genes above their basal levels during pathological growth.

The Myc super-enhancer region is required for tumorigenesis in mice

We have shown earlier that deletion of a 1.7 kb Myc-335 enhancer sequence located at the 8q24 super-enhancer region is required for intestinal tumorigenesis in mice (Sur et al., 2012b). As the super-enhancer region deleted in MycΔ2−540/Δ2−540 mice carries risk also for other cancer types, including breast and bladder cancer, we tested the susceptibility of the MycΔ2−540/Δ2−540 mice to carcinogen induced bladder and mammary tumorigenesis. The MycΔ2−540/Δ2−540 mice were not resistant to N-Butyl-N(4-hydroxybutyl) nitrosamine (BBN) induced bladder tumors. Both wild-type (n = 8)
and Myc<sup>Δ2.540/Δ2.540</sup> (n = 8) mice developed urothelial changes ranging from hyperplasia to high grade invasive urothelial carcinoma after 5 months of BBN treatment. In contrast, comparison of median tumor-free survival times of wild-type and Myc<sup>Δ2.540/Δ2.540</sup> mice exposed to mammary-tumor inducing dimethylbenzaanthracene/ medroxypregesterone (DMBA/MPA) regimen revealed that the Myc<sup>Δ2.540/Δ2.540</sup> mice were partially resistant to mammary tumorigenesis (Figure 6a). The median tumor-free survival time for the wild-type and Myc<sup>Δ2.540/Δ2.540</sup> mice was 88 and >120 days, respectively. Although we cannot pinpoint the specific regions that contribute to breast tumorigenesis by analysis of the Myc<sup>Δ2.540/Δ2.540</sup> fibroblasts lines were analysed to confirm the downregulation of Myc. Ingenuity pathway analysis performed on one of these is shown.

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The following figure supplement is available for figure 5:

**Figure supplement 1.** Scatter plot comparing the median of FPKM values of gene transcripts in colon of wild-type (n = 4) and Myc<sup>Δ2.540/Δ2.540</sup> (n = 4) mice.

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**Figure 5.** Differential effect of Myc<sup>Δ2.540/Δ2.540</sup> deletion on MYC target gene expression. (a) Scatter plot comparing the average Fragments per kilobase of exons per million fragments mapped (FPKM) values of gene transcripts in colon and spleen of wild-type (n = 4) and Myc<sup>Δ2.540/Δ2.540</sup> (n = 4) mice. Genes showing significant (q < 0.05) differential expression are marked in red (Myc) or green (other genes). For median FPKM values of gene transcripts see **Figure 5—figure supplement 1** (b) Upstream regulator analysis of RNA-seq data shows that the highest ranked potential regulator affected in the slow growing Myc<sup>Δ2.540/Δ2.540</sup> fibroblasts is MYC. The activation z-scores are to infer the activation states of predicted upstream regulators. The overlap p-values were calculated from all the regulator-targeted differential expression genes using Fisher’s Exact Test. Two independent Myc<sup>Δ2.540/Δ2.540</sup> fibroblasts lines were analysed to confirm the downregulation of Myc. Ingenuity pathway analysis performed on one of these is shown.

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**Figure supplement 1.** Scatter plot comparing the median of FPKM values of gene transcripts in colon of wild-type (n = 4) and Myc<sup>Δ2.540/Δ2.540</sup> (n = 4) mice.

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Apc<sup>min</sup> background. In this study the wild-type mice had on an average 56 polyps at around 4 months of age (n = 5) when they had to be euthanized for ethical reasons similar to what we reported previously. The Apc<sup>min</sup>; Myc<sup>42–540/42–540</sup> looked healthy and had on an average 2.4 polyps even at 6 months of age (n = 5) compared to an average of 14.33 polyps reported for the Apc<sup>min</sup>; Myc<sup>-335</sup>/C0/C0 null mice at 4 months of age (Figure 6b). Together with our earlier findings, these results indicate that loss of the 8q24 super-enhancer region makes mice resistant to both genetically and chemically induced tumors.

We further tested the requirement of this region for the proliferation of cancer cell lines in cultures. We found that the corresponding region (hg19: chr8:128226490–128746456) was also required for GP5d colon cancer cell growth, as indicated by progressive loss of cells bearing a CRISPR/Cas9 induced deletion of the region during co-culture with unedited cells in the population (Figure 6c).

**Figure 6.** Myc<sup>–2 to –540 kb</sup> genomic region is required for the growth of cancers in vivo and cancer cells in vitro. (a) Tumor-free survival plots showing resistance of Myc<sup>42–540/42–540</sup> mice to development of DMBA/MPA induced mammary tumors. p-value=0.0002 (Mantel-Cox Log-rank test). See Figure 6—source data 1 for details. (b) The Myc<sup>–2 to –540 kb</sup> deletion results in fewer polyps than the Myc<sup>-335</sup> deletion alone. p-value=0.00019 (Students T-test, 2-tailed). Apc<sup>min</sup> mice were of 4 months of age (n = 5) and Apc<sup>min</sup>; Myc<sup>42–540/42–540</sup> mice were 6 months old (n = 5) at the time of analysis. Filled circles correspond to individual mice and red color denotes the median. See Figure 6—source data 1 for details. Bar equals 5 mm. (c) Crispr-Cas9 mediated deletion of region corresponding to Myc<sup>42–540/42–540</sup> in human GP5d colon cancer cells, results in a loss of the edited cells over time. Top panel shows the active enhancer elements in GP5d cells within this region as determined by ChIP-seq analysis of histone H3 lysine 27 acetylation (H3K27ac). The sites of sgRNAs (black lines) and genotyping primers (blue arrows) used are indicated (not to scale). Red arrows mark the enhancer regions used in this study. Bottom panel shows the PCR-genotyping of the MYC locus and the control IGH locus showing the specific loss of the cells with the edited MYC locus over time. GAPDH was used as internal control. The right panel in each set shows absence of any deletion in the non-transfected cells (day 2). 100 bp ladder DNA molecular weight marker is shown (M).

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The following source data is available for figure 6.

**Source data 1.** Survival time and intestinal polyp numbers for mice in Figure 6a-b.

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Discussion

The region around the MYC gene carries inherited risk towards multiple major forms of cancer. On aggregate, this region contributes more to inherited cancer than any other locus in the human genome. The risk alleles for different cancer types are located in multiple distinct linkage disequilibrium blocks, indicating that different variants contribute to different cancer types. Several of these regions containing risk variants have been implicated in regulation of MYC expression (Hallikas et al., 2006; Sur et al., 2012b; Herranz et al., 2014; Uslu et al., 2014), suggesting that a large number of enhancers within this region can drive tumorigenesis. Some of the identified elements have also been shown to have roles in normal development (Herranz et al., 2014; Uslu et al., 2014).

To study the role of the 8q24 region more systematically, we have in this work deleted several individual enhancer elements, and also analyzed the effect of larger deletions on normal development and carcinogenesis in mice. Our analysis of mice lacking a 538 kb region upstream of the Myc gene suggests that enhancer elements within this region cooperatively enhance Myc expression. Deletion of individual enhancers in this region has only a weak (Sur et al., 2012b) or no effect on Myc expression in the mouse intestine in contrast to the deletion of the entire super-enhancer region, which leads to severe decrease in Myc expression in multiple tissues.

Myc deficient mouse embryos die due to placental defect at E9.5. The embryos are also smaller in size than wild-type embryos (Davis et al., 1993). However, when Myc is deleted only in the epiblast, the embryos grow normally and survive until E11.5, when they die due to defects in hematopoiesis (Dubois et al., 2008). None of these defects are observed in mice homozygous for the deletion of super-enhancer region. The 8q24 super-enhancer region is thus dispensable for MYC function both in the placenta and during early hematopoiesis. In our mouse colony, the super-enhancer region deficient mice also do not display the size or weight differences reported for Myc heterozygous mice that have a 50% reduction in MYC activity (Trumpp et al., 2001). These results indicate that tissue-specific enhancers that reside outside of the deleted regions drive sufficient MYC expression in the tissues that contribute to the phenotypes observed in Myc-/- and Myc-/+ mice. Consistently with this, several hematopoietic enhancers have been identified in the region 3' of the MYC ORF (Hnisz et al., 2013; Shi et al., 2013; Herranz et al., 2014).

Myc heterozygous mice also display increased longevity and enhanced healthspan (Hofmann et al., 2015). Although the deletion of the super-enhancer region that contains tissue-specific enhancers regulating MYC expression is not equivalent to a heterozygous deletion of the Myc gene in the whole body, the Myc-/-2-540/+2-540 mice could be an interesting model for identification of the tissues that contribute to the longevity phenotype.

Despite decreased levels of MYC in multiple adult tissues, the mice lacking the super-enhancer region are viable, fertile and display normal tissue morphology in all the tissues we investigated. They display no overt phenotype and do not have marked defects in cell proliferation. The mice are, however, resistant to intestinal tumorigenesis, and DMBA-induced mammary tumors, indicating that this region is important for tumorigenesis also in mice. Our data thus shows that despite the central role of this region in tumorigenesis (Sur et al., 2012b; Lovén et al., 2013), it is dispensable for normal tissue development and homeostasis under laboratory conditions. Whereas this result may appear very surprising, it is consistent with the original identification of this region using genome-wide association studies (GWAS). GWAS has a high power to identify common variants, and most variants that are common have only a limited effect on physiological functions. This is because a variant that has strong positive or negative effect is rapidly fixed or lost, respectively. Thus, GWAS are specifically biased to find variants that have a relatively large effect on disease, but a small effect on fitness.

Most genes in mammals do not have haploinsufficient phenotypes. Such buffering could be due to mechanisms that maintain constant expression level irrespective of gene dose. However, a simpler buffering mechanism involves either expressing a gene at a very low level where it has no effect, or at a high level where it can contribute its functions even if its expression level is decreased due to transcriptional noise or loss-of-function of one allele. A similar two state mechanism where physiological transcription factor (TF) activity levels in the relevant cell types are either too low to drive any target genes (off state), or high enough to activate all important targets (on state) could also mechanistically explain why most heterozygous null mutations of TF genes have no apparent phenotype.
Figure 7. Model showing the activity of the Myc super-enhancer region during normal homeostasis (left) and cancer (right). During normal tissue homeostasis (left), Myc enhancers are not strongly active, and MYC activity is relatively low. The MYC expression level is insufficient to recruit enough MAX proteins to MYC/MAX heterodimers to drive strong induction of the MYC target genes, which instead remain under the control of basal transcription factors (BTF). Under conditions of normal rapid proliferation as seen during embryonic development or during pathological insults in the adult during immune stimulation and wound healing, levels of MYC expression increase, and MYC super-enhancers become activated, leading to the recruitment of MAX proteins and the strong induction of MYC target genes.

Levels of MYC expression range from low to high, with targets classified as those with high affinity MYC binding sites (red arrows) or low affinity MYC binding sites (gray arrows). In normal adult tissue homeostasis, MYC activity is low, and targets remain under the control of basal transcription factors. In cancer, MYC activity is high, and targets are induced by MYC/MAX heterodimers.

Figure 7 continued on next page.

Dave et al. eLife 2017;6:e23382. DOI: 10.7554/eLife.23382
Our analysis of the role of MYC in normal colon is consistent with such a simple buffering model (Figure 7). However, it should be noted that this buffering mechanism does not operate in all tissues and under all conditions. For example, Myc gene dose has effects on mouse size, longevity and hematopoiesis (Davis et al., 1993; Trumpp et al., 2001; Dubois et al., 2008; Hofmann et al., 2015). In addition, the level of expression of the Myc gene has quantitative effects on cell proliferation under pathological conditions such as activation of T-cells (Heinzel et al., 2017). These results indicate that in some situations, MYC is expressed at a level where cell growth responds linearly to small changes in MYC levels (Figure 7, middle panel). However, the lack of an overt phenotype in our model under normal physiological conditions in the absence of infection or tissue damage suggests that growth during normal tissue homeostasis in at least some adult tissues does not linearly respond to changes in MYC levels. The lack of an overt phenotype should not, however, be taken to mean that the mice have no phenotype at all. As the super-enhancer region contains several highly conserved DNA segments, and affects cell growth in culture, we expect that it will also affect responses to tissue damage or some other perturbation that we have not investigated here. Therefore, further studies are needed to determine the role of the super-enhancer region in various chronic and acute models of injury and infection.

Based on our data and the earlier literature we propose that under normal physiological conditions in the intestine, the Myc gene regulatory system is in the off state, and a basal level of expression of the MYC target genes is maintained by a MYC-independent mechanism. The target genes are thus only sensitive to an increase in MYC levels. Consistently, an 80% decrease of Myc mRNA expression does not lead to a proliferation defect, or major changes in expression of known MYC target genes. In contrast, in tumors the system is locked to an on state, where MYC targets are driven to a maximal level by MYC, and the targets are now only sensitive to a decrease in MYC activity (Figure 7).

The requirement of MYC in tumor cells appears absolute. In transgenic animal models, overexpression of MYC leads to deregulated proliferation and tumor development in multiple tissues (Felsher and Bishop, 1999; Pelengaris et al., 1999; D’Cruz et al., 2001; Jain et al., 2002; Shachaf et al., 2004). Furthermore, inhibition of MYC almost invariably causes growth arrest of cancer cells both in culture and in vivo (Soucek et al., 2002, 2004; Hart et al., 2014). Despite the importance of MYC for cancer growth, it appears that the role of MYC in controlling growth during adult tissue homeostasis is limited. In the adult tissues, MYC is expressed in rapidly proliferating compartments of the body like the intestinal crypts and skin. Deletion of Myc in these compartments does not result in prominent proliferation defects (Wilson et al., 2004; Baena et al., 2005; Bettes et al., 2005; Muncan et al., 2006). Although there is still controversy regarding MYC requirement for the intestinal homeostasis, in the skin MYC is dispensable under normal adult proliferation and homeostasis in vivo (Oskarsson et al., 2006). It is however required for Ras mediated tumorigenesis and growth of fibroblasts and keratinocytes in vitro (Mateyak et al., 1997; Oskarsson et al., 2006). Taken together, these results suggest that MYC is required for pathological proliferation, but is less important and in many cases dispensable for normal homeostasis of tissues in the adult. Our results are consistent with these observations.

Prior to our study it was not clear whether the MYC dependence of cancer cells in vivo and normal cells in culture is due to shared regulatory mechanisms. Our results have uncovered striking mechanistic similarities between growth of normal cells in culture, and growth of cancer cells in vivo by showing that MYC expression depend on the same genetic elements in cultured normal cells and in cancer cells. The similarity between tumor cells and cultured normal cells also suggest that many
potential drugs that block cancer cell growth may have been inadvertently discarded due to their negative effects on growth of normal cells in culture, even when they might not have affected normal tissue homeostasis in vivo.

Our results show that the MYC super-enhancer region that carries multi-cancer susceptibility in humans contributes to the formation of multiple tumor types also in mice. Despite its role in tumor formation, it is dispensable for normal development and homeostasis. Loss of the super-enhancer region leads to low MYC expression, but the lowered expression does not translate to changes in expression of MYC target genes in the intestine. Thus, the MYC/MA\(x\)/MNT system (Grandori et al., 2000) that drives cell growth and proliferation is robustly set to an off state during normal homeostasis, whereas in cancer, the system is locked to a pathological on state. This also explains how physiological growth control can be robust to small perturbations and transcriptional noise. Taken together, our results reveal an important difference between the transcriptional states of normal and cancer cells, and suggest that therapeutic interventions that decrease the activity of the Myc super-enhancer region would be well tolerated.

Materials and methods

Mouse strains

We generated cKO Myc-196 and cKO Myc-540 strains with loxP sites flanking the regions chr15:61445326–61447611 and chr15:61789274–61791107, respectively (Taconic). These mice were crossed to Ella-cre mouse strain (Jackson Laboratory) to generate mice with enhancer deletions. Myc-CTCF\textsuperscript{mut} mouse strain was generated by mutating the CTCF-binding site at chr15:61983375–61983647 TGGCCAGTAGAGGACAC to TGGAACGTCTTGAATGC. In order to generate large deletions at the Myc locus (Myc<\textsuperscript{a,2-367} and Myc<\textsuperscript{a,2-540}) Myc-367<sup>–</sup> and Myc-540<sup>–</sup> were crossed to Myc-CTCF\textsuperscript{mut} that were also heterozygous for the Rosa26-Cre (Taconic). The Myc-540<sup>–</sup>, Myc-196<sup>–</sup> and Myc-CTCF\textsuperscript{mut} carry one loxP site at the respective loci (chr15:61445326, chr15:61618287 and chr15:61983375). The loxP site on chr15:61983375 is located immediately 5′ of the mutant CTCF binding site. We obtained compound heterozygotes carrying the chr15:61445326 or the chr15:61618287 loxP site together with the loxP site on chr15:61983375 and the Rosa26-Cre. The compound heterozygotes were screened by PCR for the interallelic recombination and the resultant deletion and duplication of the intervening sequence. Mice mosaic for the deletion and duplication were backcrossed to the C57Bl/6 mice in order to segregate the chromosomes carrying the deletion. The F1 heterozygotes were intercrossed to generate mice with homozygous large deletions. Myc-335 strain has been previously described (Sur et al., 2012b). All mice used in the study were on a C57Bl/6 genetic background. All mouse experiments were conducted in accordance with the local ethical guidelines, after approval of the protocols by the ethics committee of the Board of Agriculture, Experimental Animal Authority, Stockholm South, Sweden (Dnr S50/13, S11/15 and S16/15). The sequences of the different primer pairs used for genotypings are given in Supplementary file 2.

Mammary gland whole mount analysis

Inguinal mammary glands were removed from 8 week old virgin females and spread on glass slides. These were fixed for 4 hr in Carnoy’s fixative and subsequently stained O/N with Carmine Alum. The whole mounts were rinsed and dehydrated through increasing series of ethanol and cleared in xylene before mounting with the pertex mounting medium.

Quantitative PCR analysis

qPCR was performed as described previously (Sur et al., 2012b). Essentially, total RNA was isolated from whole tissue by homogenizing in RNA Bee reagent (ambios AMS Biotechnology) followed by RNA isolation using Qiagen’s RNA MiniElute kit according to manufacturers’ protocols. 0.5–1 μg of total RNA was reverse transcribed using high capacity reverse transcription kit in a 20 μl reaction (Applied Biosystems). Quantitative PCR in triplicates was performed using the SYBR select master mix (Applied Biosystems) on the LightCycler 480 instrument (Roche). For normalization, mouse β-actin transcripts were used as internal controls. Following primer pairs were used for quantitative PCR analysis.
**Myc-Fw**: 5'-GGGGCTTTGCCTCCGAGCCT-3', **Myc-Rev**: 5'-TGAGGGGCATCGTGGCT-3', **β-actin-Fw**: 5'-CTGTGCAGTCCGGTCCACCG-3', **β-actin-Rev**: 5'-CATGCCGGAGCCGTTGTCGAC-3'.

**RNA-sequencing**

NEBNext Ultra Directional RNA library Prep kit (NEB) was used for preparing the samples for RNA-seq together with the NEBNext Poly(A) mRNA magnetic isolation module (NEB) according to manufacturers protocol. In the case of tissues 1–2 mg and for cultured fibroblasts 200 ng of total RNA was used as starting material. For library preparation, adapters and index primers from NEBNext Multiplex Oligos for Illumina kit were used. The RNA-seq library was sequenced on a HiSeq2000 (Illumina). Sequencing reads were mapped to the mouse reference genome (NCBI37/mm9) using Tophat2 (version 2.0.13; RRID:SCR_013035) (Kim et al., 2013). Cuffdiff (version 2.2.1; RRID:SCR_001647) was used for differential gene expression analysis and for graphical representation, CummeRbund package (version 2.8.2; RRID:SCR_014568) (Trapnell et al., 2012) was used. The upstream regulator analysis was performed on all the significant differentially expressed genes (Cuffdiff q-value <0.05) using QIAGEN’s Ingenuity Pathway Analysis (IPA, QIAGEN Redwood city, www.qiagen.com/ingenuity; version 24718999, updated 2015-09-14; RRID:SCR_008653).

**ChIP-seq**

ChIP-seq was performed as described in (Sur et al., 2012b; Yan et al., 2013) with the following modifications: Adult 8–10 week old mice were euthanized and colon was removed, rinsed with cold PBS and cut into fine pieces. Tissue was crosslinked with 1.5% formaldehyde and cultured cells were crosslinked with 1% formaldehyde for 10 min at room temperature and quenched with 0.33M Glycine. Sequences were mapped to the mouse reference genome (NCBI37/mm9) and human reference genome (hg19) using Burrows-Wheeler Alignment tool (bwa) (version 0.6.2) (Li and Durbin, 2009) with default parameters. All antibodies used in ChIP-seq experiments were ChIP-grade. In each experiment a non-specific IgG was used as control. Following antibodies were used for ChIP-seq experiments: rabbit anti-H3 lysine 27 acetylation (H3K27ac) (abcam, ab4729; RRID:AB_2118291), mouse anti-H3 lysine four trimethylation (H3K4me3) (abcam, ab1012; RRID:AB_442796), rabbit anti-Rad21 (Santa Cruz, sc-98784; RRID:AB_2238151), goat anti-CTCF (Santa Cruz, sc-15914X; RRID:AB_2086899), rabbit anti-SMC1A (Bethyl Laboratories, A300-055A; RRID:AB_2192467), rabbit IgG (Santa Cruz, sc-2027; RRID:AB_737197), mouse IgG (Santa Cruz, sc-2025; RRID:AB_737182), goat IgG (Santa Cruz, sc-2028; RRID:AB_737167). ChIPseq data for Tcf7l2 was used from ENA accession number PRJEB3354 (Sur et al., 2012a) and for GP5d cells from ENA accession number PRJEB1429 (Yan et al., 2013a). For visualization, ChIP-seq read depth data were average smoothed across windows of 10 pixels (H3K27ac and H3K4me3) or five pixels (Tcf7l2) in UCSC Genome Browser; RRID: SCR_005780 or alternatively visualized in Integrative Genomics Viewer (IGV, version 2.3; RRID:SCR_011793).

**Bisulfite sequencing**

Genomic DNA was isolated using Qiagen’s Blood & Tissue Genomic DNA extraction kit. Around 1 µg of wild-type and 250 ng of Myc<sup>Δ2-540/Δ2-540</sup> null fibroblast genomic DNA was sonicated to 300 bp fragments using Covaris S220 sonicator. Subsequent to end polishing and A base addition, cytosine methylated paired end adapters (Integrated DNA technologies) were ligated to the DNA fragments. The adapter sequences are as follows

5'-P-GATCGGAAGAGCGGTTCAGCAGGAATGCCGAG
5'-ACACTCTTTCCCTACACGACGCTCTTCCGATCT

After adapter ligation 300–600 bp fragments were size-selected on a 2% agarose gel. Bisulfite-conversion was carried out using ZYMO EZ DNA Methylation-Gold kit (cat. no. D5005). PCR amplification with 12 and 18 cycles was carried out to prepare libraries from the wild-type and Myc<sup>Δ2-540/Δ2-540</sup> null mouse fibroblasts, respectively. The primer pair used for PCR amplification was as follows

**PE PCR Primer P1:**

5'-AATGATACGGCGACCACCGAGATGTCTACTCTTCTTCCCTACACGACGCTTCCGATCT
PE PCR Primer P2:

5'-CAAGCAGAAGACGGCATACGAGATCGGTCTCGGCATTCCTGCTGAACC
GCTCTTCCGATCT

The final library was size-selected for 250–300 bp fragments on a 2% agarose gel and 150 bp sequenced from both ends on two lanes of a HiSeq 4000 (Illumina). Raw sequencing reads were quality and adapter trimmed with cutadapt version 1.8.1 (RRID:SCR_011841) in Trim Galore version 0.4.0 (RRID:SCR_011847). Trimming of low-quality ends was done using Phred score cutoff 30. In addition, all reads were trimmed by 2 bp from their 3’ end. Adapter trimming was performed using the first 13 bp of the standard Illumina paired-end adapters with stringency overlap two and error rate 0.1. Read alignment was performed against mouse genome mm9 with Bismark (version v0.14.3; RRID:SCR_005604) (Krueger and Andrews, 2011) and Bowtie 2 (version 2.2.4; RRID:SCR_005476) (Langmead and Salzberg, 2012). Duplicates were removed using the Bismark deduplicate function. Extraction of methylation calls was done with Bismark methylation extractor discarding first 10 bp of both reads and reading methylation calls of overlapping parts of the paired reads from the first read (–no_overlap parameter). Genomic sites with the coverage of at least 10 reads were considered and methylation ratios smoothed with loess method across 49 bp windows.

All sequencing data is uploaded to European Nucleotide Archive (ENA, EMBL-EBI; RRID:SCR_006515) under accession number PRJEB11397 (Dave et al., 2016; http://www.ebi.ac.uk/ena/data/view/PRJEB11397).

**Immunohistochemistry and flow cytometry**

Five micron paraffin embedded tissue sections were processed for immuno-histochemistry as previously described (Sur et al., 2012b). Rabbit polyclonal anti-Myc (Santa Cruz, sc-764; RRID:AB_631276) (1:500), Rabbit monoclonal anti Ki-67 (abcam, ab16667; RRID:AB_302459) (1:200), Goat polyclonal anti-Vimentin (Santa Cruz, sc-7557; RRID:AB_793998) (1:500), biotinylated goat anti-Rabbit IgG (Vector Laboratories, BA1000; RRID:AB_2313606) and biotinylated rabbit anti-Goat IgG (Vector Laboratories, BA5000; RRID:AB_2336126) (1:350) antibodies were used. For flow cytometry, single cell suspensions of spleen and bone-marrow and cells from peripheral blood were stained with Fc-block (CD16/CD32 clone 93, Biolegend, 101302, RRID:AB_312801) and subsequently with CD19 (clone 1D3, BD Biosciences, RRID:AB_11154223), TER119 (clone TER119, Biolegend 100308, RRID:AB_312673), NK1.1(clone PK136, Biolegend, 108716, RRID:AB_493590), GR1/LY6G (clone RB6-8C5, Biolegend, 108410, RRID:AB_313375), CD4 (clone RM4-5, BD Biosciences, 563747) and CD8a (clone 53–6.7, BD Biosciences, 563332). Dead cells were visualized using Propidium iodide. Samples were analyzed using a BD LSRFortessa instrument.

**Isolation and culture of mouse primary fibroblasts**

Fibroblasts were isolated from adult mice by dissecting the skin to ~1 mm³ pieces, and allowing the pieces to adhere to cell culture plates, followed by addition of DMEM medium supplemented with 10% FCS and antibiotics. The fibroblasts were allowed to migrate out from the explants, after which the cells were collected by trypsinization and passaged in the same media for 1–3 passages. For growth assays, 2 × 10⁶ cells were plated per well in 96 well plates. Cells were trypsinized and counted using hemocytometer at respective time points.

**Tumor induction**

Mammary tumors

Six week-old female mice were implanted s.c. with medroxyprogesterone acetate (MPA) pellets (50 mg with a 90 days release period from Innovative Research of America). Subsequently 100 µl of 10 mg/ml dimethylbenz[a]anthracene (DMBA)/oil solution (Sigma) was administered via gavage at 7, 8, 10, 11, 13 and 14 weeks of age. Mice were checked twice a week for development of palpable tumors. Detection of palpable mass in the mammary gland was taken as the end point for tumor-free survival analysis.
Bladder tumors
Ten week-old male mice were administered 0.1% N-Butyl-N-(4-hydroxybutyl) nitrosamine (BBN) (Sigma) in drinking water for five months. At the end of the treatment the mice were sacrificed and the bladders scored for tumor development.

Intestinal tumors
Apc\textsuperscript{min} mouse strain (Jackson Laboratory RRID:MGI:5438590) was used as a model for spontaneous development of intestinal tumors.

CRISPR-Cas9 mediated deletion of super-enhancer region in GP5d cell line
CRISPR-Cas9 mediated deletion of MYC super-enhancer region on chromosome 8q24 (GRCh37/hg19 chr8: 128226403–128746490) and Immunoglobulin Heavy (IGH) gene locus on chromosome 14q32.33 (GRCh37/hg19 chr14: 106527004–107035452) were carried out in GP5d (Sigma, 95090715; RRID:CVCL_1235, confirmed by STR profiling at ECACC) colon cancer cell line stably expressing Cas9 protein. A lentiviral plasmid containing Cas9 fused via a self-cleaving 2A peptide to a blasticidin resistance gene, was packaged into lentiviral particles using the packaging plasmids psPAX2 (a gift from Didier Trono, Addgene plasmid # 12260, RRID:SCR_000237) and pCMV-VSV-G (a gift from Robert Weinberg (Addgene plasmid # 8454, RRID:SCR_000237). The virus was used to transduce GP5d colon cancer cells. 48 hr after transduction, GP5d cells expressing Cas9 (GP5d-Cas9) were selected in 5 \( \mu \)g/ml Blasticidin (Thermo Fisher Scientific Inc., Cat. no. A1113903). The single guide RNA (sgRNA’s) were designed (http://www.broadinstitute.org/rnai/public/analysis-tools/sgrna-design) to span the entire MYC super-enhancer region and IGH locus (Figure 6), respectively (Eurofins MWG Operon). sgRNAs were cloned into an sgRNA Cloning Vector (Addgene Plasmid #41824, RRID:SCR_000237) using Gibson assembly master mix (NEBuilder HiFi DNA assembly Master Mix, Cat no. E2621S). GP5d-Cas9 (2 \( \times \) 10\(^6\)) cells were transfected (using FuGENE HD Transfection Reagent, Cat.no E2312) with 10 \( \mu \)g of eight pooled equimolar sgRNA constructs. Post transfection half of the cultured cells were collected for PCR genotyping, while the other half was re-plated for culturing. Cells were collected at day 2, 4 and subsequently every fourth day till day 32. DNA from cells was extracted (using DNeasy Blood & Tissue Kit, Qiagen Cat. no. 69506) and genotyped with 300 ng of DNA at following conditions - Initial denaturation of 95°C for 5 min; denaturation of 98°C for 15 s, annealing at 60°C for 30 s, extension at 72°C for 30 s (30 cycles for MYC super-enhancer region and 35 cycles for IGH gene locus deletion genotyping); final extension at 72°C, 5 min. Each experiment was done in triplicate. The sequences of the different guide RNAs and primer pairs used for PCR genotyping of the deletions are given in Supplementary file 2. GP5d cells were cultured in DMEM supplemented with 10% FBS and antibiotics. The cell line was mycoplasma free.

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Additional information

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Author contributions

KD, Data curation, Software, Formal analysis, Validation, Investigation, Visualization, Methodology, Writing—original draft, Project administration, Writing—review and editing; IS, Data curation, Software, Formal analysis, Supervision, Validation, Investigation, Visualization, Methodology, Writing—original draft, Writing—review and editing; JY, LB, Investigation, Methodology, Writing—review and editing; JZ, EK, FZ, Software, Formal analysis, Visualization, Writing—review and editing; XL, SK, CG, ADP, Methodology, Writing—review and editing; RM, Formal analysis, Investigation, Methodology, Writing—review and editing; JT, Conceptualization, Resources, Formal analysis, Supervision, Funding acquisition, Writing—original draft, Project administration, Writing—review and editing

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Ethics

Animal experimentation: All mouse experiments were conducted in accordance with the local ethical guidelines after approval of the protocols by the ethics committee of the The Board of Agriculture, Experimental Animal Authority, Stockholm South, Sweden (Dnr. S50/13, S11/15 and S16/15).

Additional files

Supplementary files

- Supplementary file 1. List of genes that show significant differential expression in the wild-type and the Myc^{2-540/2-540} colon (q-value <0.05).
  DOI: 10.7554/eLife.23382.018
- Supplementary file 2. List of primers and guide RNA sequences used in this study.
  DOI: 10.7554/eLife.23382.019

Major datasets

The following datasets were generated:

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References


Appendix 1

Purification of primary mouse cells

Bone marrow (BM) cells harvested from 8- to 12 weeks old C57BL/6 mice. For isolation of LSK Flt3<sup>−</sup> cells, BM cells were subjected to depletion of mature cells using a cocktail of purified antibodies containing TER119 (TER-119; Biolegend 116202; RRID:AB_313703), CD19 (1D3; BD, 553783; RRID:AB_395047), CD3 (17A2; BD, 555273; RRID:AB_395697), GR1 (RB6-8C5; Biolegend, 108402; RRID:AB_313367) and CD11b (M1/70; Biolegend, 101202; RRID:AB_312785) in combination with sheep anti-rat IgG Dynabeads (Invitrogen; 11035). Non-depleted cells were stained with Ter119 PECy5 (TER-119), NK1.1 PECy5 (PK136), CD3 PECy5 (145–2C11), CD19 PECF594 (1D3), CD11b PECy5 (M1/70), CD11c PECy7 (N418), CD19 PECF594 (1D3), CD24 PECy7 (H1/80), CD45 PECy7 (30-F11), CD49b PECy7 (2B11), CD49d PECy7 (2A3), CD54 PECy7 (1B8), CD71 PECy7 (8E5), CD80 PECy7 (16-10A1), CD86 PECy7 (37B5), CD95 PECy7 (21–512), CD102 PECy7 (1A1), CD117 PECy7 (2B8), KIT (CD117) APCeFlour780 (2B8; eBioscience, 47–1171–82; RRID:AB_1272177), SCA1 PB (D7; Biolegend, 108120; RRID:AB_493273), FLT3 (CD135) PE (A2F10; Biolegend, 135306; RRID:AB_1877217), CD11C PECy7 (N418; Biolegend, 117318; RRID:AB_493568), LY6C APC (HK1.4; Biolegend, 128016; RRID:AB_1732076) and IL7R (CD127) biotin (A7R34 Biolegend 135006; RRID:AB_2126118; visualized using Streptavidin-QD655; Invitrogen Q10121MP). LSK Flt3<sup>−</sup> (HSCs) cells were subsequently FACS sorted as TER119/CD3/GR1/NK1.1/CD102<sup>low</sup>/CD19<sup>−</sup>CD11c<sup>−</sup>IL7R<sup>−</sup>SCA1<sup>++</sup>KIT<sup>−</sup>FLT3. For isolation of mature B cells, BM cells were subjected to MACS column enrichment of CD45R (B220)+ cells using anti-CD45R beads (Miltenyi Biotec, 130-049-501). B220 enriched cells were stained with TER119 PECy5 (TER-119), GR1 PECy5 (RB6-8C5), CD11b PECy5 (M1/70), IgD PB (11–26c.2a), IgM PECy7 (11/14), CD19 PECF594 (1D3). Mature B cells subsequently FACS sorted as TER119/GR1/CD102<sup>−</sup>CD19<sup>−</sup>IgM<sup>+</sup>IgD<sup>+</sup>. Propidium iodide (Life technologies, p3566) was used as a dead cell discriminator when sorting live cells (for RNAseq experiments) and Aqua fluorescent reactive dye (Life technologies; L34957) when sorting fixed cells (for ChIPseq experiments).

For ChIPseq experiments, fully antibody/viability dye-stained cells (5 × 10<sup>6</sup> cells/ml) were fixed by incubation with 1% formaldehyde (ThermoFisher Scientific; 28908) for 10 min at room temperature (RT). Formaldehyde was quenched using 0.1 vol 1 M glycine and incubated for 10 min at RT. Cells were additionally washed with 0.1 M glycine before being resuspended in PBS with 2% FCS prior to FACS sorting. Cell sorting was done on a BD FACSAriaIII cell sorter (BD Biosciences).

ChIP-sequencing

3 μg of polyclonal anti-H3K27Ac (Diagenode, cat# C15410196, lot# A1723-0041D, RRID:AB_2637097) or H3K4me2 (Millipore, cat#07–030, lot#2089140 and lot#2309072, RRID:AB_11213050) antibody was bound to 10 μl Protein G-coupled Dynabeads (ThermoFisher) per ChIP and incubated with rotation for 4 hr at 4°C. Pellets of 0.5 × 10<sup>6</sup> PFA-fixed cells were resuspended in 100 μl SDS lysis buffer (50 mM Tris/HCl, 0.5% SDS, and 10 mM EDTA), placed cold for 15 min and sonicated for 12 cycles of 30 s on/30 s off on high power using a Bioruptor Plus (Diagenode). Samples were centrifuged, and supernatants transferred to new tubes. After addition of 200 μl of ChIP dilution buffer (50 mM Tris/HCl, 225 mM NaCl, 0.15% NaDod, and 1.5% Triton-X) and 4 μl of 50X protease inhibitors (Roche), samples were incubated at room temperature for 10 min. 10% of each sample was saved for input controls. Antibody-coated dynabeads were washed, resuspended with cell lysate and rotated overnight at 4°C.
Immunoprecipitated chromatin was washed once with low salt buffer (50 mM Tris/HCl, 150 mM NaCl, 0.1% SDS, 0.1% NaDOC, 1% Triton X-100, 1 mM EDTA), high salt buffer (50 mM Tris/HCl, 500 mM NaCl, 0.1% SDS, 0.1% NaDoc, 1% Triton X-100, and 1 mM EDTA) and LiCl buffer (10 mM Tris/HCl, 250 mM LiCl, 0.5% IGEPAL CA-630, 0.5% NaDoc, 1 mM EDTA) followed by two washings with TE buffer. For reversal of crosslinking, chromatin complexes and input control samples were diluted in 200 μl ChIP elution buffer (10 mM Tris/HCl, 0.5% SDS, 300 mM NaCl, and 5 mM EDTA) and 2 μl of 20 μg/ml proteinase K (Thermo Scientific). Samples were vortexed and incubated shaking overnight at 65°C. After reverse crosslinking, 1 μl 20 μg/ml RNAse (Sigma) was added and incubated at 37°C for 30 min. After another 2 hr of incubation with 2 μl of proteinase K at 55°C, samples were placed in a magnet to trap magnetic beads and supernatant collected. DNA purification was carried out using Qiagen MinElute PCR Purification Kit.

DNA concentrations in purified samples were measured using Qubit dsDNA HS Kit (Invitrogen). Libraries were prepared using Rubicon ThruPLEX DNA-seq 12S Kit, according to manufacturer’s instructions. 2 ng of chromatin was used when available but samples below Qubit detection levels (<0.5–1.5 ng) were frequently used. After 11 cycles of PCR amplification, adapter cleanup was done using Agencourt AmPureXP beads (Beckman Coulter) at a ratio of 1:0.88. Libraries with an average size of 400–500 bp were pooled and single-end sequenced (50 cycles) using the Illumina sequencing platform (HiSeq2000).

RNA-sequencing
For RNA extraction 5,000–10,000 cells were sorted into buffer RLT (Qiagen) with β-mercaptoethanol and total RNA was extracted using Rneasy Micro Kit (Qiagen) according to manufacturer’s instructions. On-column DNase I treatment was performed to minimize DNA contamination. Strand specific RNAseq libraries were prepared using TotalScript RNA-seq kit (Epicentre) according to the manufacturer’s instructions. Barcoded libraries were pooled and pair-end sequenced (2 × 50 cycles) using the Illumina platform (mainly HighSeq 2500).

ChIP-seq data analysis
Quality of sequencing samples was assessed with FastQC (v0.11.2). Samples were mapped to the mm10 genome using Bowtie2 (v2.2.3) with default parameters (Langmead and Salzberg, 2012). Mapped reads were filtered with HOMER (v4.6) (Heinz et al., 2010) using the makeTagDirectory command, only keeping uniquely mapped reads and removing possible PCR duplicates by restricting the tags per base pair to 1 (-tbp 1). Resulting filtered reads were visualized by generating bigWig files from tag directories, using HOMER’s makeBigWig.pl with a set fragment length (fragLength 130) and normalization to 10 million reads (-norm 1e7).

RNA-seq data analysis
Quality of sequencing samples was assessed with FastQC (v0.11.2) (Andrews, 2010). Samples were mapped to the mm10 genome using STAR (v2.4) with default parameters for paired-end reads (Dobin et al., 2013). Mapped reads were filtered with HOMER (v4.6) using the makeTagDirectory command with strand specific pair-end read settings (-sspe), and removing excessive possible PCR duplicates by restricting the tags per base pair to 3 (-tbp 3). Resulting filtered reads were visualized by generating bigWig files from tag directories, using HOMER’s makeBigWig.pl with a set fragment length (-fragLength 75), normalization to 10 million reads (-norm 1e7) and stranded data setting (-strand).
The above sequencing data is accessible via ENA accession number PRJEB20316 (Dave et al., 2017; http://www.ebi.ac.uk/ena/data/view/PRJEB20316).