Local regulation of gene expression by lncRNA promoters, transcription and splicing

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Mammalian genomes are pervasively transcribed1,2 to produce thousands of long non-coding RNAs (lncRNAs)3,4. A few of these lncRNAs have been shown to recruit regulatory complexes through RNA–protein interactions to influence the expression of nearby genes5–7, and it has been suggested that many other lncRNAs can also act as local regulators8,9. Such local functions could explain the observation that lncRNA expression is often correlated with the expression of nearby genes5–7, 10,11. However, these correlations have been challenging to dissect12 and could alternatively result from processes that are not mediated by the lncRNA transcripts themselves. For example, some gene promoters have been proposed to have dual functions as enhancers13–16, and the process of transcription itself may contribute to gene regulation by recruiting activating factors or remodelling nucleosomes10,17,18. Here we use genetic manipulation in mouse cell lines to dissect 12 genomic loci that produce lncRNAs and find that 5 of these loci influence the expression of a neighbouring gene in cis. Notably, none of these effects requires the specific lncRNA transcripts themselves and instead involves general processes associated with their production, including enhancer-like activity of gene promoters, the process of transcription, and the splicing of the transcript. Furthermore, such effects are not limited to lncRNA loci: we find that four out of six protein-coding loci also influence the expression of a neighbour in cis. These results demonstrate that cross-talk among neighbouring genes is a prevalent phenomenon that can involve multiple mechanisms and cis-regulatory signals, including a role for RNA splice sites. These mechanisms may explain the function and evolution of some genomic loci that produce lncRNAs and broadly contribute to the regulation of both coding and non-coding genes.

We analysed 12 lncRNA loci whose RNA transcripts in mouse embryonic stem cells (mES cells) show preferential localization to the nucleus and span a range of abundance levels (Methods and Extended Data Fig. 1). For each locus, we looked for direct regulatory effects on local gene expression by using a genetic approach based on classical cis–trans tests (Fig. 1a and Supplementary Note 1). Specifically, we generated clonal cell lines carrying heterozygous knockouts of the promoter (~600–1,000-bp deletions) (Fig. 1b) and compared the expression of nearby genes within 1 Mb on the cis and trans alleles (that is, on the modified and unmodified homologous chromosomes in the same cells) (Supplementary Note 2). Changes in neighbouring gene expression that involve only the cis allele very probably result from direct, local functions of the lncRNA locus, while changes that involve both the cis and trans alleles probably result as indirect, downstream consequences of the lncRNA acting elsewhere (Supplementary Note 1). We performed genetic modifications in 129/castaneus F1 hybrid mES cells that contain a polymorphic site every ~140bp, enabling us to distinguish the two alleles using RNA sequencing (Fig. 1b, Extended Data Fig. 2 and Supplementary Note 3).

At 5 of these 12 lncRNA loci, promoter knockouts significantly affected the expression of a nearby gene in an allele-specific manner (false discovery rate <10%), including both activating and repressive effects (Fig. 1c, d, Supplementary Note 4 and Extended Data Fig. 3). For each locus, the affected gene was located immediately adjacent to, and within 5–71 kb of, the knocked-out promoter (Fig. 1c and Extended Data Fig. 4). This indicates that a substantial fraction of lncRNA loci influence the expression of a neighbouring gene.

To test whether such effects were specific to lncRNA loci, we deleted the promoters of six protein-coding genes (Extended Data Fig. 1). Surprisingly, knockouts at four of these loci also affected the expression of a neighbour in cis (Fig. 1c, d and Extended Data Fig. 5). Thus, both non-coding and coding loci can directly influence local gene expression. These regulatory connections may contribute to the observed correlations in the expression of neighbouring genes, which have been reported both for lncRNAs and for mRNAs10,11,19,20.

Because in these experiments we deleted gene promoters, the mechanisms underlying such cis effects could in principle involve (i) DNA regulatory elements in gene promoters13–16, (ii) the process...
of transcription; or (iii) the RNA transcripts themselves, (Extended Data Fig. 6a). To begin to distinguish among these possible mechanisms, we inserted early polyadenylation signals (pAS) 0.5–3 kb downstream of each transcription start site (TSS) that eliminated the production of most of the RNA while retaining the promoter sequence intact (Fig. 2 and Extended Data Fig. 6b, c, see Methods). We examined four lncRNA loci and two mRNA loci where promoter deletion affected the expression of a neighbouring gene (see Supplementary Note 5).

As one example, we describe the linc1536 locus, hereafter called Bendr (Bendr-regulating effects not dependent on the RNA, Fig. 2a). Whereas deleting the Bendr promoter reduced the expression of the adjacent Bendr gene by 57%, inserting a pAS into the first intron of Bendr (~570 bp downstream of the TSS in this ~13 kb locus) had no effect on Bendr expression despite eliminating the spliced Bendr RNA (Fig. 2b, c). Furthermore, global run-on sequencing (GRO-seq) did not detect any transcriptionally engaged polymerase upstream of the pAS insertion (Fig. 2c and Extended Data Fig. 7a), perhaps because the pAS prevents RNA splicing, which may substantially reduce transcriptional activity in the modified locus. Therefore, cis activation of Bend4 requires neither the mature Bendr RNA transcript nor significant Bendr transcription. Instead, this effect appears to be mediated by DNA regulatory elements in the ~750 bp knocked-out promoter-proximal region.

In total, at five of the six loci examined with pAS insertions (including three lncRNAs and two mRNAs), DNA regulatory elements in the promoter-proximal sequences appear to be responsible for activating a neighbouring gene (Extended Data Fig. 7b). Although the promoters in these loci would not be classified as ‘enhancers’ based on H3K4me3/H3K4me1 ratios, they are bound by mES cell transcription factors (Extended Data Fig. 7c) and are located in close proximity to their neighbouring target genes (Fig. 1c and Extended Data Fig. 7d, e), suggesting that these promoters may affect local gene expression through mechanisms similar to enhancers.

We also identified one locus, linc1319 (renamed Blastr: bivalent locus of Sfmbt2) is upregulated by the splicing and transcription of an RNA, where both promoter deletions and pAS insertions substantially reduced the expression of a neighbouring gene, Sfmbt2, located 5 kb upstream (Fig. 3a). To dissect the regulatory mechanism, we tested whether the activation of Sfmbt2 is mediated by a sequence-specific function of the Blastr transcript or the process of transcription (by which we mean one or more sequence-independent functions associated with transcription, such as changes in chromatin state or recruitment of co-factors). To test the first possibility, we knocked out each of the three downstream exons and three introns. None of these deletions impaired Sfmbt2 activation (Fig. 3b, Supplementary Note 6), suggesting that the activation of Sfmbt2 does not require unique sequences or structures in the Blastr transcript itself. To test the second possibility, we engineered pAS insertions at five different locations in the first exon or intron (+40 bp to +15 kb downstream of the TSS) and found that increasing the length of the Blastr transcribed region led to increased activation of Sfmbt2 (Fig. 3b and Extended Data Fig. 8a, b). We note that changing the length of the transcribed region affected the total amount of engaged polymerase in the Blastr locus (Fig. 3c). Thus, Sfmbt2 activation responds to changes in the length/amount of transcriptional activity in the Blastr locus but does not appear to require specific sequence elements in the mature Blastr transcript (Supplementary Note 7).

Because promoter-proximal splice sites and the process of splicing can enhance transcription—in some cases by as much as 100-fold— we tested whether the splicing of Blastr is involved in Sfmbt2 activation. Upon deleting the 5′ splice site of the first intron of Blastr (Extended Data Fig. 8c), we observed a 94% reduction in Blastr transcription (as assayed by GRO-seq), a 92% reduction in the levels of the mature Blastr transcript, and an 85% reduction in Sfmbt2 expression (Fig. 3b, c and Extended Data Fig. 8a, b), demonstrating that the first 5′ splice site of Blastr has a critical role in activating Blastr and Sfmbt2 transcription. By contrast, downstream splice sites were dispensable: upon deleting downstream Blastr exons, splicing skipped over the removed exon to the next available 3′ splice site (Extended Data Fig. 8d) and Sfmbt2 expression was unaffected (Fig. 3b).

Together, these data demonstrate that the 5′ splice site and the process of transcription in the Blastr locus are important for its ability to regulate Sfmbt2. This indicates that the Blastr RNA is in fact required for Sfmbt2 activation (splicing involves direct interactions between the spliceosome and the nascent transcript), although this mechanism does not appear to depend on the precise sequence of the RNA beyond the presence of initial splice signals. One possibility is that the 5′ splice site promotes transcriptional activity in the Blastr locus, which in turn recruits components of the transcriptional machinery that act on the nearby Sfmbt2 promoter (Fig. 3d, Supplementary Note 7). Consistent with this model, altering transcription or splicing in the Blastr locus led to changes in chromatin state at the Sfmbt2 promoter (including reductions in H3K4me3 and spreading of H3K27me3) and reduced occupancy of engaged RNA polymerase in the paused position just downstream of the Sfmbt2 TSS (Extended Data Fig. 8b, e, f). Thus, changes in Blastr transcription and splicing may affect Sfmbt2 expression in part by altering chromatin state and RNA polymerase occupancy at the Sfmbt2 promoter (Fig. 3d and Supplementary Note 7).

In summary, genetic dissection of 12 lncRNA loci and 6 mRNA loci found that 9 loci (50%) regulate the expression of a neighbouring gene (Extended Data Fig. 9). In most of these loci, including Bendr, local effects are mediated by enhancer-like functions of DNA elements in promoters. In one locus, Blastr, the processes of transcription and splicing also contribute to cis-regulatory functions, perhaps by increasing the local concentration of transcription-associated factors. We did not identify any lncRNA loci in which local effects are mediated by sequence-specific functions of the lncRNA transcript. Because there exist thousands of other loci that fit our selection criteria, we expect that similar mechanisms broadly contribute to gene regulation in many loci (Supplementary Note 8).

The frequent cross-talk between neighbouring genes observed in our study indicates that gene loci can encode multiple independent categories of functions. Category I involves functions of the RNA product: mRNAs provide a template for protein synthesis, and some non-coding transcripts (for example, XIST) act as functional lncRNAs. Category II involves the effects of transcription-related functions of DNA regulatory elements.
processes—including mechanisms mediated by promoters, transcription, and splicing—on the regulation of other nearby genes.

The fact that many lncRNA loci have category II functions does not necessarily mean that they do not also have category I functions, and we note that our experiments do not rule out the possibility that the lncRNAs dissected in this study have RNA-mediated functions other than on local gene regulation. However, the prevalence of category II functions suggests a model for the evolutionary origins of some lncRNAs. In loci where a promoter acts as an enhancer, RNA transcripts may arise as non-functional by-products. In loci where co-transcriptional processes have cis-regulatory functions, the nascent transcripts might arise as non-functional by-products. In loci where co-transcriptional processes have cis-regulatory functions, the nascent transcripts might arise as non-functional by-products. In loci where co-transcriptional processes have cis-regulatory functions, the nascent transcripts might arise as non-functional by-products. In loci where co-transcriptional processes have cis-regulatory functions, the nascent transcripts might arise as non-functional by-products. In loci where co-transcriptional processes have cis-regulatory functions, the nascent transcripts might arise as non-functional by-products. In loci where co-transcriptional processes have cis-regulatory functions, the nascent transcripts might arise as non-functional by-products. In loci where co-transcriptional processes have cis-regulatory functions, the nascent transcripts might arise as non-functional by-products. In loci where co-transcriptional processes have cis-regulatory functions, the nascent transcripts might arise as non-functional by-products. In loci where co-transcriptional processes have cis-regulatory functions, the nascent transcripts might arise as non-functional by-products.

Figure 3 | Transcription and splicing of Blustr activates Sfmbt2 expression. a, Poly(A)+ RNA-seq, GRO-seq, and H3K4me3 ChIP–seq in the Blustr locus. Sfmbt2 has two alternative TSSs. b, Poly(A)+ RNA expression on modified alleles compared to controls (arrows). Error bars, 95% confidence interval for the mean (n = 2 alleles, except for pAS +15 kb where n = 1, see Supplementary Table 1). Sfmbt2 pAS comparisons: two-sided t-test; *P < 0.05; **P < 0.01. c, Allele-specific GRO-seq signal for clones carrying indicated modifications. Only reads mapping to the modified allele are shown (castaneus for pAS +2 kb; 129 for others). d, Model for how transcription in the Blustr locus activates Sfmbt2.

Figure 4 | Evolutionary conservation of mES cell lncRNAs and their promoters. a, Classification of a subset of lncRNAs expressed in mES cells (see Supplementary Note 9, Methods). b, Eleven lncRNAs have promoters whose syntenic sequence corresponds to putative DNA regulatory elements (REs) marked by DNase I hypersensitivity (HS) in human ES cells. c, Example: linc1494. d, Enhancers and lncRNA promoters are significantly enriched for corresponding to human regulatory elements (pie chart, ***P < 1 × 10−10; χ² test versus GC-matched random regions) and show elevated sequence conservation compared to GC-matched regions (bar plot, **P < 0.01; ***P < 0.001, Mann–Whitney U-test versus ii + iii).
cis-regulatory connections—including mechanisms for specificity and the potential for cooperative dynamics of gene activation—represent key areas for future investigation.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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METHODS

Cells and line culture. F1 hybrid 129/castaneus female mouse embryonic stem cells (gift from K. Plath) or V6.5 male mouse embryonic stem cells (gift from A. Meissner) were cultured in serum-free N2B27-based medium (250 μl neurobasal media (Gibco), 250 μl DMEM/F12 (Gibco), 5 ml 100 mM l-glutamine (Gibco), 3 μl 2-mercaptoethanol, 50 μg human leukemia initiation factor (5 × 10^5 units, EMD Millipore), 7.4 μg progesterone, 10 μg bovine insulin (Sigma), 350 μl 7.5% BSA, 0.2 mM dNTPs (Gibco), supplemented with MEK inhibitor PD9081, 50 μl 10 mM, SelleckChem), and GSK3b inhibitor CHIR99021 (150 μl 10 mM, SelleckChem)). Prior to plating cells, tissue culture dishes were pre-treated with PBS + 0.2% gelatin (Sigma) and 1.75 μg/ml 1-laminin (Sigma) for 2–10 h at 37°C.

At each passage, cells were trypsinized for 3–5 min in TVP solution (0.025% trypsin, 1% chicken serum (Sigma), and 1 mM EDTA in PBS pH 7.4) at room temperature.

Cells tested negative for mycoplasma contamination and were authenticated by comparing polymorphisms to 129S1 and castaneus genomes.

Cellular fractionation. To estimate the relative abundance of lncRNAs in different cellular compartments, we performed cellular fractionation to isolate chromatin-associated, soluble nuclear, and cytoplasmic fractions essentially as described [25]. In brief, we first lysed 5 million cells in 200 μl cold cell lysis buffer (10 mM Tris-HCl pH 7.5, 0.05% IGEPAL CA-630, 150 mM NaCl), incubating on ice for 5 min. We layered the cell lysate over 2.5 volumes of chilled sucrose (Gibco), 5 ml 50 mM neurobasal media (Gibco), 250 ml DMEM/F12 (Gibco), 5 ml 100 μM l-glutamine (Gibco), and supplemented with 1 mM EDTA in PBS pH 7.4 at room temperature. Cell lysates were microcentrifuged at 15,000 g for 10 min. The supernatant from this spun became the cytoplasmic fraction. After washing the pellet of nuclei with PBS (pH 7.5) + 1 mM EDTA, we resuspended the pellet in 100 μl of cold glycerol buffer (20 mM Tris-HCl pH 7.5, 75 mM NaCl, 0.5 mM EDTA, 0.85 mM DTT, 0.125 mM PMSF, 50% glycerol) by gently flicking the tube. We added 100 μl of cold nuclei lysis buffer (10 mM HEPES pH 7.5, 1 mM DTT, 7.5 MgCl2, 0.2 mM dNTPs (Gibco), 2 μM EDTA, 0.3 M NaCl, and 1 μM Igepal CA 630), then vortexed for four seconds. After 2 min on ice, we spun the nuclear lysate at 15,000 g for 2 min. This supernatant was collected as the soluble nuclear (nucleoplasm) fraction. We rinsed the remaining pellet (chromatin fraction) in PBS + 1 mM EDTA, then resuspended the chromatin in 300 μl chromatin DNase buffer (20 mM Tris-HCl pH 7.5, 50 mM KCl, 4 mM MgCl2, 0.5 mM CaCl2, 2 mM TCEP, 0.5 mM PMSF, 0.4% sodium deoxycholate, 1% Igepal CA 630, 0.1% N-lauroylsarcosine) plus 15 μl murine RNase inhibitor (NEB) and 30 μl TURBO DNase (Ambion). The DNase digestion proceeded for 20 min at 37°C and was halted by adding 10 mM EDTA and 5 mM EGTA. Protein was digested with proteinase K for 1 h at 37°C. LncRNA was isolated using Zymo RNA Concentrator-25 columns (two columns for each lncRNA sample).

To select for transfected cells, we replaced the media 24 h after transfection with 8 ml of 0.5% puromycin. We allowed cells to grow for 7–8 days, replacing the media every 2–3 days. We hand-picked 88 individual colonies and 8 control colonies for each transfection in 5 μl media, added 20 μl of TVP for ~10–20 min at 37°C to dissociate the colonies, and then split the colonies into two identical plates. We verified efficient transfection by examining GFP expression after 24 h.

To design single-guide RNAs (sgRNAs), we built custom software to calculate a specificity score (based on potential off-target sites using the algorithm described at http://crispr.mit.edu (see ref. 31)) and an efficacy score (based on a sequence model for sgRNA efficiency as previously described [22]) for each 20-nt targeting sequence. We removed guides with specificity scores <20 or efficacy scores >0.7. To avoid T-rich sequences that result in premature termination of Pol III-mediated sgRNA transcription, we removed guides with more than one T in the four bases closest to the seed region, guides with more than three consecutive Ts, and guides with more than eight Ts total. We removed guides with homopolymer stretches of five or more bases and guides with GC content <20% or >90%. We removed guides that overlapped a known 129/castaneus SNP [33]. Within a given region, we typically chose the three remaining guides with the highest specificity scores. The sequences of all sgRNAs used in this study are listed in Supplementary Table 2.

Promoter deletion guide placement. To knock out an lncRNA or mRNA promoter, we chose 2–3 sgRNAs located in windows 600–1,000 bp upstream and downstream of the TSS, leading to deletions of approximately 600–1,000 bp surrounding the TSS. We adjusted the precise deletion boundaries outward if we could not successfully design guides in these regions (for example, because they were located in repetitive sequences). We note that we often found that the wild-type alleles in heterozygous knockouts were affected by scars from repair of sgRNA double-stranded breaks. Accordingly, we adjusted the bounds if necessary to cut outside of the exons of the mRNA or lncRNA and thus avoid damaging the exonic sequences on the wild-type alleles in heterozygous knockouts. We note that the presence of these scars (and their lack of allele-specific effects on the expression of neighboring genes) indicate that the cis effects observed upon deleting promoters were not merely a result of CRISPR-mediated cutting and subsequent DNA repair.

Genetic deletions with CRISPR/Cas9. To delete specific sequences, we co-transfected 100 ng of Cas9-expressing plasmids (PX330-NonGuide), 300 ng of a pool of sgRNA-expressing plasmids (pZB-Sg3), and 100 ng of a plasmid expressing EGFP and a puromycin selectable marker from a CAG promoter (ps-pGFPFP). To create PX330-NonGuide, we modified PX330 (gift from F. Zhang, Addgene plasmid #44230 (ref. 34)) to remove the sgRNA expression cassette. To generate pZB-Sg3, we cloned a human U6 promoter and optimized sgRNA scaffold sequence [35] into a minimal vector with an ampicillin-selectable marker and a CoEl1 replication origin. We transfected batches of 250,000 mouse embryonic stem cells using the Neon Transfection System (Invitrogen), using one pulse of 40 ms at 1,200 V and plated two batches of cells (500,000 total) into a 96-well plate in 200 μl media. As an internal control for each set of transfections, we performed a transfection using four guides with no predicted target sites in the mouse genome.

Sanger sequencing with CRISPR/Cas9. To genotype each promoter knockout, we extracted genomic DNA and performed PCR using primers spanning the deleted sequence. We sequenced each clone by running the PCR products on agarose gels and comparing PCR amplicon sizes to predicted wild-type and deletion band sizes. We confirmed the sequences of wild-type and deletion bands by Sanger sequencing or high-throughput sequencing through barcoded amplicon sequencing on an Illumina MiSeq (see Supplementary Table 2). Where possible, we used known polymorphic sites from 129S1 and castaneus genomes [33] to determine the haplotype-resolved genotype of each clone. Based on the genotyping data, we nominated clones for RNA sequencing. We eliminated clones showing evidence of polyclonal or subclonal mutations, or complex mutations such as inversion or duplication of the genomic sequence between the sgRNAs. The sequences of all RNA sequencing libraries are listed in Supplementary Table 2.
We fragmented RNA to an average of ~150 nt by adding 2 μl Ambion fragmentation buffer and incubating at 70 °C for exactly 2.5 min. After transferring quickly to ice, we added 40 μl of a master mix containing 12 μl 5′ FNL buffer (50 mM Tris-HCl pH 7.5, 5 mM MgCl₂, 0.6 mM CaCl₂, 50 mM KCl, 10 mM DTT, 0.01% Triton X-100), 1 μl Murine RNase Inhibitor (NEB), 3 μl FastAP Thermosensitive Alkaline Phosphatase (Thermo Scientific), 3 μl T4 Polynucleotide Kinase (NEB), and 1 μl TURBO DNase (Life Technologies). We incubated this reaction at 37 °C for 30 min, then cleaned the reaction with MyOne SILANE magnetic beads and eluted in 20 μl H₂O.

Allele-specific gene expression measurements from RNA sequencing. We sequenced RNA libraries on an Illumina HiSeq 2500 (Read 1: 38 cycles; Read 2: 30 cycles; Index: 8 cycles). The first read includes the 8–nt barcode added during the first adaptor ligation (see above). Following processing to separate samples based on the inline barcodes, we filtered out sequencing reads that aligned to highly abundant RNA transcripts, including ribosomal RNAs, snRNAs, and repetitive sequences, as defined by RefSeq and RepeatMasker. A FASTA file containing these sequences is available at the Gene Expression Omnibus (GSE55914).

We developed a computational pipeline to estimate allele-specific expression from RNA-sequencing data. We created two separate reference files for the 129S1 and castaneus haplotypes, starting with the mm9 genome build and layering on SNPs based on whole-genome sequencing of each of the two mouse strains. We aligned RNA-sequencing data separately to each of the two haplotypes using TopHat (version 2.0.8). We combined the results of the two alignments using PySuspenders, which identifies reads that map specifically to one or the other allele and splits them into separate BAM files. We discarded duplicate reads and reads with MAPQ < 30. After generating separate BAM files containing the reads mapping to each allele, we counted reads that mapped to each RefSeq transcript (including both spliced and unspliced isoforms) using Scripture and calculated ‘allelic expression ratios’ for each gene (counts from 129 allele divided by total counts from both 129 and castaneus alleles). The distribution of allelic expression ratios for all active genes in mES cells was centred on 0.5, indicating that on average each gene is expressed equally from the 129 and castaneus alleles (Extended Data Fig. 2b). This indicates that there is no systematic bias in our mapping procedure towards one allele or the other.

RNA-seq data analysis. We processed RNA-sequencing data sets in batches corresponding to sets of libraries made on the same day with the same hybrid selection probe pool. We removed samples with fewer than 100,000 non-repetitive, unique, allele-informative reads. For within-batch quality control, we performed hierarchical clustering on all samples by their allelic expression ratios and removed the 2–5% of outlier samples, which were largely comprised of clones that showed monoallelic expression from the X chromosome.

Assessment of gene knockout by expression analysis. The PCR genotyping procedure described above provided putative genotypes for the cell clones. We confirmed the genotype of cells by analysing the allele-specific expression of the knocked-out gene in each clone. We required that clones show >80% reduction of expression of the knocked out gene on the appropriate allele in order to include the clone in downstream analysis. Incomplete reduction of expression in some cases appeared to result from use of alternative TSSs that were not included in the deleted sequence. In other cases, incomplete reduction of expression appeared to result from subclonal genetic mosaicism within the cell line, which probably resulted from deletions that occurred after several cell divisions, leading to genetic differences between individual cells in a colony. For further analysis, we focused on gene loci where we obtained at least two heterozygous knockout clones.

Barplots for allele-specific expression data. Barplots that depict allele-specific expression or GRO-seq transcription on modified alleles compared to controls (for example, Fig. 1d) were calculated as follows. For each modified allele, allele-specific measurements were normalized to the corresponding alleles in wild-type clones (for example, values for castaneus knockout alleles were divided by the mean of unmodified castaneus alleles in wild-type clones). We performed the same calculation for unmodified alleles in wild-type clones to create a null distribution. For modified alleles, we further scaled these values by dividing by the mean of the wild-type alleles in heterozygous knockout clones. The value of each bar represents a mean of these normalized measurements.

Identifying significant changes in allele-specific expression. In developing a statistical pipeline to detect allele-specific effects of these genetic manipulations, we sought to distinguish local effects of the genetic deletion from downstream effects that result as a consequence of either IncRNA/mRNA functions elsewhere in the cell, off-target effects, or biological/technical variation between clonal cell lines (Supplementary Note 1). Our power to detect these effects varies between different measured genes (owing to their level of expression and availability of SNPs) and between different knockout targets (owing to differences in the numbers of knockout clones analysed).

To account for these two variables, we developed a statistical approach to empirically estimate the false discovery rate of allele-specific changes in the RNA expression or GRO-seq transcription on modified alleles compared to controls. For each gene in the neighbourhood of one of our promoter deletions, we calculated three statistics: (i) a t-test statistic comparing the average change in expression for each of the knockout alleles (including both heterozygous and homozygous knockout clones), normalized to the expression of the gene on the wild-type allele of the heterozygous clones; (ii) a z-score statistic comparing the
expression of the knockout allele in heterozygous clones to the expression of the wild-type allele in the same clone; and (iii) a t-test statistic comparing the heterozygotes to the wild-type control clones using the allelic expression ratio after applying a variance-stabilizing transformation (arcsin of the square root of the allelic expression ratio). For a given gene, only samples with at least 20 allele-informative reads were considered, in order to enable accurate estimates of allele-specific expression. These three tests differ in whether they incorporate information from homozygous clones and how they normalize between knockout- and non-knockout samples. We followed the transcription of three of the three tests in order to regard the gene as significant, as described below.

We note that each underlying measure was approximately normally distributed, with some apparent outliers across hundreds of control clones; we conservatively included these outliers in calculating each test statistic. We examined differences in variation between knockout and control alleles with Levene’s test. For estimates of the variance of distributions presented in figures, see Supplementary Table 1. Because the distributions are only approximately normal, we assessed the significance of each of these gene-level statistics by permutation, sampling other cell lines from the same experimental batch and randomly assigning them as heterozygous or homozygous knockout clones to match the distribution of genotypes of the real samples. We calculated an empirical false discovery rate for the sum of these permutation ranks, testing each of the neighbouring genes for other genes, transcription was unaffected despite pAS knock-in, consistent with the location-dependent efficiency previously observed for this pAS sequence42.

This short pAS sequence (spA) succeeded in halting the transcription of three RNAs: Blustr

5’site splice junction.

To knock out the 5’splice site of Blustr, we co-transfected mES cells as described above, using a single sgRNA pZB plasmid and 200 ng of dsDNA oligonucleotide donor for homologous recombination (Extended Data Fig. 8c). The oligo was used as an ultrasome from Integrated DNA Technologies (Supplementary Table 2). We genotyped these insertions through amplicon sequencing using an Illumina MiSeq (primers in Supplementary Table 2). Transcriptional activity with GRO-Seq. We used precision run-on sequencing (PRO-seq)43, a variant of global run-on sequencing44, to map transcriptionally engaged RNA polymerase for a subset of clones. Clones for PRO-seq (as well as ChIP-seq and assays for transposase-accessible chromatin with high-throughput sequencing (ATAC-seq)) were chosen from among the recoverable knockout cell lines with a preference for clones with homologous knockouts or knockouts on the 129 allele only. We performed PRO-seq as previously described45, with modifications. We harvested 10 million mES cells by scraping, washing in cold PBS, and spinning at 330g for 3 min. The cell pellet was resuspended in 1 ml cold dousing buffer (10 mM Tris-HCl pH 7.4, 300 mM sucrose, 3 mM MgCl2, 2 mM MgCl2, 0.1% (v/v) Triton X-100, and 0.5 mM DTT) immediately frozen, and stored at −80°C until use.

A 28S 1→ Nuclear Run-On (NRO) mix was prepared as follows: 1 M Tris-HCl, pH 8.0, 1 M MgCl2, 2 M KCl, and 0.1 M DTT. 5 μl of 1 mM Biotin-11-CTP (Perkin Elmer), 1 μl of 0.05 mM CTP, 2.5 μl of 2 M ATP, 2.5 μl of 2 M GTP, 2.5 μl of 2 mM UTP (Sigma Aldrich), 6.5 μl of nuclelease free water, and 2 μl of SUPERaseIn (Ambion) were added to the 2× NRO mix and mixed well before the addition of 50 μl of 2% NLS. The NRO reaction mix was mixed well and preheated to 37°C. 100 μl of NRO mix was added to 100 μl of nuclei in storage buffer. The reaction was mixed gently by pipetting and incubated at 37°C for 3 min, mixing halfway through. To halt the reaction 500 μl of TRIzol LS (Thermo Fisher) was added. The sample was mixed well, and incubated at room temperature for 5 min. RNA was isolated through a chloroform extraction and ethanol precipitation, and resuspended in 20 μl of H2O. The RNA was heat denatured at 65°C for 40s and fragmented on ice for 10 min with 5 μl of 1N NaOH. To stop the reaction, 5 μl of 1 M acetic acid and 20 μl of 1 M Tris-HCl, pH 7.4 were added. To remove unincorporated biotinylated nucleotides, the sample was passed through a P-30 exchange column (BioRad). 1 μl of RNase inhibitor was added to the ~50 μl of RNA and the first biotin enrichment was then performed.

Each biotin enrichment was performed as follows. To prepare the Streptavidin M280 beads, 1 μl of 5 μg/ml Streptavidin-Agarose (Novagen) was incubated, 100 μl of beads were taken per sample and washed once in 0.1N NaOH with 50 mM NaCl and twice in 100 mM NaCl. Beads were resuspended in 160 μl of binding buffer (10 mM Tris-HCl, pH 7.4, 300 mM NaCl, and 0.1% (v/v) Triton X-100). To each sample an equal volume of Streptavidin M280 beads was added, mixed, and incubated on a rotator for 20 min at room temperature. The beads were magnetically separated and washed twice in 500 μl of ice cold high salt wash buffer (50 mM Tris-HCl, pH 7.4, 2.5 mM NaCl, and 0.5% (v/v) Triton X-100), twice in 500 μl of binding buffer, and once in 500 μl of low salt wash buffer (50 mM Tris-HCl, pH 7.4 and 0.1% (v/v) Triton X-100). To harvest the RNA, 300 μl of TRIzol (Thermo Fisher) was added to the mixture and incubated at room temperature for 5 min. 40 μl of chloroform was added and mixture was incubated at room temperature for 3 min. The samples were centrifuged at 14,000g for 5 min at 4°C. The aqueous phase was collected and transferred to a new tube; the remaining organic phase was removed from the beads. The TRIzol extraction was then repeated as above and the two aqueous phases were combined. RNA was purified with a chloroform
extraction and ethanol precipitation, and resuspended in nuclease-free water. RNA sequencing libraries were then prepared as described above, except that SILANE clean-ups were replaced with streptavidin-biotin capture enrichments until after reverse transcription (a total of three enrichments).

We sequenced PRO-seq libraries to a depth of ~10 million 30-bp paired-end reads. To analyse the data, we mapped and processed the RNA sequencing data as described above, including aligning individually to the 129 and castaneus genomes. Figures showing Allele-specific GRO-seq’ depict coverage for reads that uniquely map to either allele. Figure S3A depicts the reads distributed in the two allelic replicates. To assess the relative read density in the promoter-proximal region and gene body of Sfmbt2, we counted reads in the 2 kb region downstream of the first Sfmbt2 TSS and in the remainder of the gene body.47 We calculated the pause index as the ratio of these two quantities, normalized to total read count. We noticed that different PRO-seq libraries had subtle biases in the relative fraction of reads aligning to the TSS versus the gene body, leading to slightly offset distributions of pause indices across all genes, and so we corrected for these biases in each library by normalizing TSS and gene body RPKM to the median of the ~5,000 genes with coverage across all samples.

Chromatin accessibility with ATAC-seq. Libraries were generated as previously described35 using 50,000 mES cells. We generated duplicate ATAC-seq libraries for each clonal cell line examined and sequenced each to a depth of ~40 million 30-bp paired end reads. We aligned paired-end DNA sequencing reads using Bowtie2 (ref. 48) to each of the 129 and castaneus genomes with the following parameters: --minimap mismatch 1000, removed duplicate reads using Picard (http://picard.sourceforge.net), and filtered to uniquely aligning reads using samtools (MAQ < 30, https://github.com/samtools/samtools). For plotting normalized read coverage at the first Sfmbt2 promoters, we combined data from the two biological replicates (two independent measures of the same cell line) and connected paired-end reads to generate fragments. Fragment coverage was normalized by the total number of uniquely aligned reads.

Chromatin immunoprecipitation. ChiP–seq for H3K4me3 and H3K27me3 was performed using monoclonal antibodies as previously described48. Sequencing data was analysed as for ATAC-seq described above.

Validation of allele-specific RNA expression with ddPCR. To validate our RNA-seq based measurements of allele-specific expression, we used a quantitative allele-specific PCR assay to verify measurements for Blastr and Sfmbt2. We isolated RNA from harvested mES cells using RNeasy 96 columns and performed a DNAse treatment followed by reverse transcription of 500 ng of RNA (total reaction volume 20 μL). We performed droplet digital PCR (ddPCR) using Bio-Rad Custom ddPCR Assays that involve qPCR primers flanking a polymorphic site and two allele-specific fluorescent probes. For Blastr: left primer sequence: GACACATTCCTCCTTCAACA; right primer sequence: GAACAGTTTGTCCTGCC; probe sequence: TAAAGGTTGGAAGCTCCCAAG (129 allele, FAM) or AGTGGAGGCGAACTTCAAG (castaneus, HEX). ddPCR was done with 2.2 μl of CDNA, 11 μl of Supermix (BioRad), 1.1 μl of each probe, and 7.7 μl of cells (10^6 cells) per reaction followed by droplet generation. PCR was performed as follows: 95°C for 10 min; cycling at 94°C for 30s and 55°C for 1 min for a total of 40 cycles; and 98°C for 10 min. Readout was done using the QX200 Droplet Reader and Quantasoftware (BioRad) to determine the total number of droplets containing each allele. We calculated allelic expression ratios from these values and compared it to values generated through RNA-sequencing and hybrid selection of the same RNA samples (Extended Data Fig. 2d, e).

External ChiP–seq, RNA-seq, and DNase HS data. We used the following data from ENCODE46: H3K4me3, H3K4me1, H3K27ac, and CTCF ChiP–seq in mES cells (ES-Bruc); DNase hypersensitivity sequencing in mES cells (E14); H3K4me3 and CTCF ChIP–seq in mouse embryonic stem cells, and RNA-seq and RNA-sequencing in data in H1-hES cells (nuclear poly(A)^+ total nuclear). To assess transcription factor binding to mRNA and lincRNA promoters (Extended Data Fig. 7c), we examined mES cell ChiP–seq peaks available from Kagey et al. at the Gene Expression Omnibus (GSE22563).31

DNA purification for assessing proximity contacts. To examine the proximity contacts of the linc1405 locus, we used the RAP-DNA protocol, which we initially developed in order to map RNA localization to chromatin, to capture linc1405 DNA.35 In brief, we cross-linked live cells to fix endogenous chromatin complexes, then purified a target RNA region using a pool of oligonucleotides targeting the linc1405 locus (Supplementary Table 3). Here we note that the read start as the linc1405 RNA—In this way, we specifically capture the linc1405 DNA and do not directly capture the linc1405 RNA itself. We mapped the 3D proximity contacts of the linc1405 locus through high-throughput sequencing of co-purified DNA and calculated the normalized enrichment to an input DNA library in 1-kb windows (Extended Data Fig. 7e). Annotations for topologically associated domains (TADs) were downloaded from the Ren Laboratory (http://chromosome.sdsce.edu/mouse/hc-d/download.html)52. LncRNA transcript annotations. For evolutionary conservation analysis, we used lincRNA annotations and isoforms previously defined based on RNA sequencing in mouse embryonic stem cells, combining annotations generated with multiple methods (Scripture41 and slncky28). We filtered the combined list using slncky28 to eliminate transcripts predicted to encode proteins or micropeptides by UCSC, transcripts that partially align to protein-coding genes (for example, pseudogenes or incomplete reconstructions), and transcripts containing coding gene duplications. Subsequently we performed several manual curation steps. We examined each isoform using a combination of long-read RNA-sequencing data, total chromatin-associated RNA sequencing data, cap analysis of gene expression (CAGE) data, and poly(A)^+ 3′-end sequencing data from mES cells38,46,47. We eliminated transcripts that appeared to result from an extended 3′ UTR of an upstream protein-coding transcript. Because the precise 5′ ends of transcripts are imprecisely assigned based on RNA-sequencing data alone, we re-assigned 5′ ends (TSSs) using a sliding-window approach to find the 10–bp window with the highest number of same-strand CAGE reads within 300–bp of the initial calculated TSS. We additionally manually curated the TSS of each lincRNA, some of which were incorrectly assigned by more than 300 bp, based on CAGE and H3K4me3 ChIP–seq data, and eliminated any where we could not identify the TSS (for example, due to an unmappable sequence or very low abundance).

Analysis of lincRNA and promoter conservation. To categorize lincRNAs by their conservation properties and promoter locations, we examined a set of 307 lincRNAs expressed in mES cells as described above. We assessed the conservation of each lincRNA through a two-step approach. We first used slncky to look in syntenic locations for evidence of lincRNA transcripts in deep poly(A)^+ RNA-seq of rat, chimp, and human induced pluripotent stem cells (iPSCs)42. lincRNAs called ‘conserved’ by this first filter have substantial evidence based on RNA-seq data that allows for independent reconstruction of the transcript in one or more of these other organisms. We categorized the remaining lincRNAs by the location of their TSS: 71 lincRNAs originate within 500–bp of an mRNA TSS on the opposite strand (divergent); 59 lincRNAs originate within the long-terminal repeats (LTRs) of endogenous retroelements; and 79 lincRNAs have their promoters in intergenic regions that do not overlap with LTRs and do not emerge from a bidirectional mRNA promoter (herein, ‘intergenic’).

Because some conserved lincRNAs might be expressed at too low a level to assemble a transcript de novo in a given species, we examined more closely the 79 intergenic lincRNAs that were called ‘mouse-specific’ in the initial slncky analysis. We applied a second, more stringent threshold to remove lincRNAs misclassified as mouse-specific due to low abundance. For each intergenic lincRNA locus, we used liftOver48 to map the 10 bp surrounding the mouse TSS (mm9) to the human genome (hg19) (minmatch = 0.1, UCSC chain). 37 of these transcripts did not lift over this step, and thus were considered mouse-specific. For the 42 that did lift over, we examined the syntenic region for evidence of poly(A)^+ RNA-seq data from human iPSCs50 or poly(A)^+ nuclear-fraction RNA-seq from hES cells (~100 to ~900 bp relative to the TSS), or for evidence of poly(A)^+ nuclear-fraction or whole-cell CAGE from hES cells (~250 to ~250 bp relative to the TSS), and removed from consideration any lincRNAs that showed evidence for RNA-seq or CAGE above a certain threshold. We chose this threshold based on a set of random intergenic regions, which were matched to the set of intergenic mouse-specific lincRNAs based on GC content. We eliminated from consideration the ten lincRNAs that showed RNA-seq or CAGE signals greater than the 90th percentile of random regions, corresponding to approximately two CAGE or RNA-seq reads in the windows described above. These ten lincRNAs were added to the ‘conserved’ section of the pie chart in Fig. 4a. Several of these ten lincRNAs correspond to substantially shortened, single-exon poly(A)^+ transcripts that show minimal overlap with the syntenic regions in mm9, although a majority of the exonomic sequence of these transcripts are not in fact conserved between human and mouse, we excluded these from consideration as putative mouse-specific lincRNAs.

For the purposes of examining the conservation properties of these intergenic mouse-specific lincRNAs, we defined a matched set of ‘enhancer’ elements. We first generated a list of regulatory elements in mES cells using the DNase hotspots called by ENCODE-UW in ES-E14 cells. As an estimate of the activity of each element, we calculated the density of H3K27ac reads in the region. From the set of intergenic elements that did not overlap a promoter, lincRNA promoter, or LTR, we selected a random subset matched to the intergenic lincRNA promoters for H3K27ac (804 elements) and of appropriate genomic distance to the TSS of the closest active gene (binned by 5kb). We call these elements ‘enhancers’ because they are marked by DNase hypersensitivity and H3K27ac but do not overlap a known gene promoter.

We compared the sequence conservation and functional conservation of three classes of elements: intergenic mouse-specific lincRNAs, matched intergenic...
enhancer elements, and GC-matched random intergenic elements. First, we computed the rate at which each set maps to human sequence. We centred each element and used liftOver (~minMatch = 0.1) to identify the syntenic region in the human genome. Elements that did not lift over at this step correspond to the white segment of the pie charts in Fig. 4 (ii, did not map). For elements that did lift over to human, we next defined the subset that map to putative regulatory elements in human. We examined a 500-bp window centred on the lifted over region and counted reads in hES cell DNase-seq data from ENCODE. We defined regions showing 50% of hES scores higher than 95% of the mapped to random intergenic regions as putative DNA regulatory elements. We note that these random intergenic regions include some enhancers; they are matched to IncRNA promoters for GC content, and thus frequently correspond to regulatory elements (which are GC-rich) that happen to be active in hES cells. For both intergenic mouse-specific IncRNAs and enhancers, ~33% of elements corresponded to putative DNA regulatory elements in human (Fig. 4d), representing a ~6.6-fold enrichment versus the random intergenic controls. To compare sequence conservation of these classes of elements, we calculated the average SiPhy score across each 500-bp region surrounding the mouse TSS or the centre of the enhancer element, using the 29 mammals alignment from the mouse perspective. We used a two-sided Mann–Whitney U-test to look for changes in the distributions of SiPhy scores to the set of mappable random intergenic regions (Fig. 4d: random ii – iii).

Impact of expression level on conservation analysis. Although the set of intergenic mES cell IncRNAs examined above does not show any significant evidence for poly(A)+ RNA in the syntenic locus in human, some of these transcripts may not be detected in human and yet still be truly conserved. These transcripts might be misclassified as mouse-specific IncRNAs for several reasons, including: (i) low expression level in hES cells and iPSC cells such that the IncRNA, by chance, is not detected based on the depth of sequencing data available; or (ii) the IncRNA is not expressed in hES cells or iPSC cells, but is expressed in a different human cell type and thus may have a conserved function.

To estimate the false positives resulting from these and other scenarios, we examined the properties of a set of 853 conserved mRNAs matched to the intergenic mouse-specific IncRNAs based on expression in mES cells. We counted the frequency at which these mRNAs would be called ‘not conserved’ by the same procedures described above: we applied the nuclear poly(A)+ CAGE and RNA-seq filters to eliminate transcripts that show detectable transcription in the 1-kb region near the TSS. While 87% of the intergenic IncRNAs described above passed these filters (and thus appeared to be mouse-specific), only 22% of the 853 mouse-expressed mRNAs were not detectably expressed in human. Of the 30 intergenic lncRNAs called mouse-specific by this more conservative analysis, 5 do indeed correspond to putative DNA regulatory elements, including linc1494 (Fig. 4c), representing a >8-fold enrichment versus GC-matched random sequences (chi-squared P < 1 × 10−18). Thus, our conclusions that some IncRNAs appear to evolve from ancestral regulatory elements are robust even with stringent thresholds.

Software for data analysis and graphical plots. We used the following software for data analysis and graphical plots: R Bioconductor (version 3.0)12, Gviz (version 1.10.11), gplots (version 2.17.0), GenomeRanger (version 1.18.4)14, tdracklayer (version 1.26.3)15, BEDTools46, Integrative Genomics Viewer (version 2.3.26)47, and vcftools (version 0.1.12)48.

Data availability. Sequencing data for this study is available at the Gene Expression Omnibus (GSE80262 and GSE85798), and additional visualizations of the data are available at http://pubs.broadinstitute.org/neighboring-genes/.

Code availability. Code for the analyses described in this paper is available from the authors upon request.

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Extended Data Figure 1 | Expression and subcellular localization of knocked-out lncRNAs and mRNAs. a, Expression of lncRNAs and mRNAs in F1 129/castaneus female mES cells, reported in fragments per kilobase per million (FPKM) in whole-cell poly(A)+ RNA-seq. Cumulative fraction is plotted for all mRNAs expressed in mES cells. Large dots represent transcripts whose promoters we deleted in this study. LncRNAs and mRNAs span a >20-fold range of abundance levels. b, Relative subcellular localization of lncRNAs and mRNAs. We sequenced poly(A)+ RNA from chromatin, soluble nuclear, and cytoplasmic fractions (see Methods) and plotted the relative abundance of mature transcripts in each fraction. We selected lncRNAs that showed localization biased towards the nuclear fractions relative to most mRNAs. For comparison, we plotted 1,000 randomly selected mRNAs (light grey).
Extended Data Figure 2 | Generation of knockout clones and measurement of allele-specific RNA expression. a, Overview of knockout and measurement protocol. b, Distribution of allelic expression ratios (number of informative reads mapping to 129S1 allele divided by the number mapping to either the 129S1 or the castaneus allele) across active genes in mES cells. c, Scatter plot of allelic expression ratios for genes with RPKM $\geq 2$ that have more than 100 allele-informative reads across all libraries. Allelic expression ratios are consistent in RNA sequencing data before and after hybrid selection (HS). d, e, Allelic expression ratios as measured by two independent methods for Blastr (d) and Sfmbt2 (e) expression in 15 clonal cell lines containing genetic modifications in the Blastr locus. Each dot represents the mean of two ddPCR technical replicates ($x$ axis) and the value from one RNA-seq technical replicate ($y$ axis). f, Example locus showing hybrid selection strategy and RNA-seq coverage for cell lines with the indicated genotype for deletion of the Bendr promoter. The $y$ axis scales represent normalized read counts and are the same for all hybrid selection tracks. The absolute level of expression for any given gene varies among clonal cell lines; throughout this work, we instead consider the relative level of expression between the two alleles in heterozygous knockout cells. For similar plots of each gene studied, see http://pubs.broadinstitute.org/neighboring-genes/.

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Extended Data Figure 3 | Read-through transcription at Meg3 and Snhg3 loci. a, Snhg3 promoter knockout reduces the levels of Rcc1 mRNA by 23%. However, sequencing of chromatin-associated RNA shows that transcription continues past the annotated 3′ end of Snhg3 into the downstream Rcc1 gene (see Methods). This read-through transcription creates a fusion transcript containing exons of both Snhg3 and Rcc1, as well as intergenic RNA. We note that this fusion transcript is also annotated in the syntenic human locus as an alternative isoform of RCC1.

Bars, relative poly(A)+ RNA expression on modified versus unmodified alleles. Error bars, 95% confidence interval for the mean (n ≥ 2 alleles, see Supplementary Table 1). b, Meg3 promoter knockout eliminates the expression not only of Meg3 but also of two additional IncRNAs encoded downstream in a tandem orientation (Rian and Mrg3). Although these three IncRNAs are annotated as separate genes, they appear to be derived from a single transcript driven by the Meg3 promoter. This is consistent with the presence of continuous chromatin-associated RNA throughout the locus and a lack of CAGE reads at the 5′ ends of Rian and Mrg3.
Extended Data Figure 4 | Promoter knockouts for five intergenic lncRNAs affect the expression of a neighbouring gene. Significance (z-score) of allele-specific expression ratios at all genes within 1 Mb of each of five lncRNA loci. Each dot represents a different heterozygous promoter knockout clone for a given gene. Dots are shown only for genes that are sufficiently highly expressed to assess allele-specific expression (see Methods). The y axis is capped at –10 to +10 standard deviations from the mean. Black, knocked-out lncRNA; blue, gene with significant allele-specific change in gene expression (FDR < 10%). Independent clones are not expected to yield the same significance value (z-score), in part because read depth differs between samples.
Extended Data Figure 5 | Promoter knockouts for four mRNAs affect the expression of a neighbouring gene. Significance (z-score) of allele-specific expression ratios at all genes within 1 Mb of each of four mRNA loci. Each dot represents a different heterozygous promoter knockout clone for a given gene. Dots are shown only for genes that are sufficiently highly expressed to assess allele-specific expression (see Methods). The y axis is capped at −10 to +10 standard deviations from the mean. Black, knocked-out lncRNA; blue, gene with significant allele-specific change in gene expression (FDR < 10%). Independent clones are not expected to yield the same significance value (z-score), in part because read depth differs between samples.
Dissecting mechanisms for how gene loci regulate a neighbour. a, Three categories of possible mechanisms by which a gene locus might regulate the expression of a neighbour. b, We used two strategies to insert pAS downstream of gene promoters. In the first strategy, we inserted a 49-bp synthetic pAS (spA) using a single-stranded DNA oligo with 75-bp homology arms (see Methods). c, In the second pAS insertion strategy, we cloned a donor plasmid containing a selection cassette and three different pAS sequences (see Methods). Homology arms of 300–800 bp were used to integrate the cassette. After isolating clones with successful insertions, we used a second round of transfections to remove the selection cassette, leaving behind three tandem pASs. EFS, elongation factor 1 promoter; Puro, puromycin resistance gene (pac); HSV-tk, herpes simplex virus thymidine kinase.
Extended Data Figure 7 | Promoters of lncRNAs and mRNAs have enhancer-like functions. a, Allele-specific GRO-seq signal for clones with the indicated modifications at the Bendr locus. Only reads specifically mapping to one of the two alleles are shown. The y axis scale represents normalized read count and is the same for all tracks. b, Allele-specific poly(A)⁺ RNA expression for genetic modifications at the linc1405, Snhg17, Gpr19, and Slc30a9 loci. Bars, average RNA expression on modified compared to unmodified (wild-type) alleles. Error bars, 95% confidence intervals for the mean (n ≥ 2 alleles, see Supplementary Table 1). Grey arrows indicate distance from the targeted locus promoter to the affected neighbouring gene. We note that, based on their location, the Snhg17 and Gpr19 pAS insertions probably allow more substantial splicing and transcription; for these loci, it is clear that the majority of the transcript is dispensable but it is possible that transcription close to the promoter may be involved in the cis-regulatory function. c, Presence (grey) or absence (white) of various chromatin marks and transcription factors in mES cells in a 1.5-kb window centred on the TSS of each targeted gene. d, Distance from each knocked-out gene to its neighbouring target gene (x axis) versus the magnitude of the effect on the expression of the neighbouring gene (per cent compared to wild-type, y axis). Blue genes represent those discussed in main text; grey genes are discussed in Supplementary Note 5. e, Proximity-based contacts between the linc1405 and Eomes loci. The y axis shows enrichment in a sequencing-based proximity assay in which we used antisense oligos to capture linc1405 DNA and any interacting, cross-linked proximal DNA (see Methods). TAD annotations are derived from Hi-C experiments in mES cells (see Methods). Blue arrow, focal contact between the linc1405 and Eomes loci.
Extended Data Figure 8 | Characterization of genetic modifications in the *Blustr* locus. a, Allele-specific GRO-seq signal for clones with the indicated modifications at the *Blustr* locus. Only reads specifically mapping to one of the two alleles are shown. The y axis scale represents normalized read count and is the same for all tracks, and is magnified five times at the indicated location to better visualize the reads in the *Sfmbt2* locus. b, Quantification of allele-specific GRO-seq signal in the *Sfmbt2* locus on alleles modified as indicated. TSS, region including the two alternative TSSs of *Sfmbt2* and 2 kb downstream; gene body, region containing the remainder of the *Sfmbt2* gene locus; pause index, ratio of TSS to gene body. Dashed grey lines indicate the 95% confidence intervals for the mean of eight wild-type clones. Bars, *n* = 8 for wild-type and *n* = 1 for others.

c, Schematic of the 5′ end of the *Blustr* locus and genotypes of two knockout clones. The 5′ splice site is located 78 bp downstream of the *Blustr* transcription start site (in this panel, *Blustr* is transcribed from left to right). One of the alleles from the two clones contains insertion of the oligo mediated by homologous recombination; the remaining three alleles contain insertions or deletions resulting from non-homologous end joining repair of sgRNA-mediated double-strand breaks, some of which also disrupt the 5′ splice site. Bar plots show allele-specific RNA expression for knockout clones and control clones (*n* = 18 for +/+ , 1 for others). Error bars, 95% confidence interval for the mean. 
d, Schematic of the observed splice structures of *Blustr* RNA transcripts in poly(A)+ RNA sequencing of the exon deletion clones. Each deletion removes a region including ~50–200 bp on either side of the exon, thereby removing both the exon and its splice sites. The Exon 4 deletion removes the endogenous pAS, leading to new isoforms of the lncRNA transcript that splice into two cryptic splice acceptors downstream. e, GRO-seq, H3K4me3 ChIP–seq, and chromatin accessibility (ATAC-seq FPKM) at the *Blustr* and *Sfmbt2* promoters in cell lines with the indicated genotypes. Deletion of the first 5′ splice site leads to a significant reduction in H3K4me3, RNA polymerase occupancy, and chromatin accessibility at the *Blustr* promoter, as well as H3K4me3 and RNA polymerase occupancy (but not accessibility) at the *Sfmbt2* promoter. f, H3K27me3 ChIP–seq at the *Blustr* and *Sfmbt2* loci in cell lines with the indicated genotypes. Deletion of the *Blustr* promoter or 5′ splice site leads to spreading of the repression-associated H3K27me3 modification across a ~30 kb region.
**Extended Data Figure 9 | Mechanisms for cross-talk between neighbouring lncRNAs and mRNAs.** Proposed mechanisms based on pAS insertion experiments and other genetic manipulations (see text). For proposed mechanisms of lncRNAs marked with daggers see Supplementary Note 5.
Extended Data Figure 10 | Classification of lncRNAs based on conservation and promoter location. 

**a.** Classification of 307 lncRNAs expressed in mES cells. 'Conserved' transcripts are those that show significant evidence of cap analysis of gene expression (CAGE) data and/or poly(A)+ RNA in syntenic loci (see Methods). Divergent, initiating within 500 bp of an mRNA TSS, on the opposite strand; ERV, endogenous retroviral repetitive element (see Supplementary Note 9). Box plot shows sequence-level conservation of the promoters of subsets of lncRNAs expressed in mES cells. Random intergenic regions are matched to lncRNA promoters by GC content. Positive Siphy score indicates evolutionary constraint on functional sequences. Orange category corresponds to mouse-specific lncRNAs that appear to have evolved from ancestral regulatory elements (REs) and correspond to sequences that show evidence for DNase I hypersensitivity in human embryonic stem cells. Significance is calculated compared to random intergenic regions using a Mann–Whitney U-test. ** *** P < 0.001. Box represents first and third quartiles; centre line represents median; whiskers represent data within 1.5 × the interquartile range.

**b.** Chromatin and RNA data for 11 mouse-specific lncRNAs that appear to have evolved from ancestral regulatory elements. In mouse, these elements show evidence for CAGE, H3K4me3, and DNase I hypersensitivity, consistent with their roles as promoters. The syntenic sequences in human do not show evidence for CAGE but nonetheless are DNase I hypersensitive and are frequently marked by H3K4me1 and/or CTCF.

**c.** Model for evolution of lncRNAs from pre-existing enhancers, which often initiate weak bidirectional transcription. Spliced transcripts may neutrally appear through the appearance of splice signals and loss of polyadenylation signals. In some cases, transcription, splicing, or other RNA processing mechanisms may feed back and contribute to the cis-regulatory function of the promoter, producing a lncRNA as a by-product.