A genome-wide library of MADM mice for single-cell genetic mosaic analysis

Graphical abstract

Mosaic Analysis with Double Markers

Cell-autonomous gene function

Lineage Tracing

Genomic Imprinting

Chromosome Segregation

uniparental disomy

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In brief
Contreras et al. generate a resource and suite of transgenic MADM mice for genetic mosaic analysis with double markers of >96% of the entire mouse genome. In addition to providing a proof of principle, they find non-random mitotic sister chromatid segregation in distinct somatic cell lineages in vivo.

Highlights
- Genome-wide resource for genetic mosaic analysis with double markers in mice
- Resource for dissection of cell-autonomous gene function of >96% of the mouse genome
- Resource for genome-wide analysis of genomic imprinting phenotypes
- MADM chromosomes reveal non-random mitotic sister chromatid segregation in vivo

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A genome-wide library of MADM mice for single-cell genetic mosaic analysis

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SUMMARY

Mosaic analysis with double markers (MADM) offers one approach to visualize and concomitantly manipulate genetically defined cells in mice with single-cell resolution. MADM applications include the analysis of lineage, single-cell morphology and physiology, genomic imprinting phenotypes, and dissection of cell-autonomous gene functions in vivo in health and disease. Yet, MADM can only be applied to <25% of all mouse genes on select chromosomes to date. To overcome this limitation, we generate transgenic mice with knocked-in MADM cassettes near the centromeres of all 19 autosomes and validate their use across organs. With this resource, >96% of the entire mouse genome can now be subjected to single-cell genetic mosaic analysis. Beyond a proof of principle, we apply our MADM library to systematically trace sister chromatid segregation in distinct mitotic cell lineages. We find striking chromosome-specific biases in segregation patterns, reflecting a putative mechanism for the asymmetric segregation of genetic determinants in somatic stem cell division.

INTRODUCTION

Genetic mosaic individuals contain cells of distinct genotypes. The phenomenon of genetic mosaicism occurs naturally and is widespread across multicellular organisms. Mosaicism may progressively emerge during life but remain silent with no obvious phenotypes (Yizhak et al., 2019). However, mosaicism is also associated with pathologies in humans including cancer or many neurological disorders (Biesecker and Spinner 2013; D’Gama and Walsh 2018). Genetic mosaic animals have been experimentally created in a number of species, and mosaic analyses provided fundamental insights in a variety of biological systems (Xu and Rubin 1993; Rossant and Spence 1998; Lee and Luo 1999, 2001; Yochem and Herman 2003; Zugates and Lee 2004; Zong et al., 2005; Lozano and Behringer 2007; Luo 2007; Germani et al., 2018; Kim et al., 2019).

One powerful application inherent to induced genetic mosaics is the ability to alter gene function at a high spatiotemporal resolution. A certain tissue can contain homozygous mutant cells for a gene of interest and wild-type cells whose phenotypes can be compared with each other directly. If the genetic mosaic is sparse, even essential genes can be manipulated without affecting the overall health or viability of the animal. Furthermore, sparse genetic mosaics provide a highly effective means with which to study the causal relationship of genetic alteration and phenotypic manifestation at the individual cell level. Genetic mosaics also facilitate the analysis of cell competition and provide an assay to create models of disease. Genetic mosaics have been most extensively generated in the fruit fly by capitalizing upon mitotic recombination between homologous chromosomes (Morgan and Bridges 1919; Stern 1936; Hotta and Benzer 1970; Xu and Rubin 1993; Lee and Luo 1999, 2001; Zugates and Lee 2004). Although technically slightly more challenging, the generation of genetic mosaics in mice is becoming routine. A number of experimental approaches have been established including mosaic analysis with double markers (MADM) that is also based on mitotic recombination (Zong et al., 2005; Luo 2007; Tasic et al., 2012; Hippenmeyer 2013).

MADM relies on Cre/loxP-mediated interchromosomal recombination to simultaneously generate homozygous mutant cells for a candidate gene of interest and wild-type cells in an otherwise heterozygous background. The induction of genetic mosaicism can be spatiotemporally controlled by the use of cell-type-specific Cre/ER driver lines (Zong et al., 2005; Hippenmeyer et al., 2013; Ali et al., 2014; Beattie et al., 2020). Concurrent to the generation of genetic mosaicism, two split genes, encoding green fluorescent protein (GFP) and tdTomato (tdT) fluorescent markers, are reconstituted that permit unequivocal tracing of individual cellular phenotypes in the homozygous mutant and heterozygous and wild-type cells, with each labeled...
Figure 1. Extension of MADM to all 19 mouse autosomes

(A) Summary of the MADM principle. For MADM, two chimeric split marker genes containing partial coding sequences of EGFP and tdT are inserted into identical genomic loci of homologous chromosomes. Following Cre-recombinase-mediated interchromosomal (trans) recombination during mitosis, the split marker genes are reconstituted and functional green and red fluorescent proteins expressed. As a result, green GFP+, red tdT+, and yellow GFP+/tdT+ cells appear sparsely, due to an inherently low stochastic interchromosomal recombination rate, within the genetically defined cell population expressing Cre recombinase. Introduction of a mutant allele distal to the MADM cassette results in a genetic mosaic with homozygous mutant cells labeled in one color (e.g., green GFP+) and homozygous wild-type sibling cells in the other (e.g., red tdT+). Heterozygous cells appear in yellow (GFP+/tdT+).

(B) Expansion of MADM to all mouse autosomes. Transgenic mice with MADM cassettes inserted close to the centromere have been generated for all 19 mouse autosomes. The directionality (forward, centromere-telomere; reverse, telomere-centromere) of marker gene transcription is indicated.

(C) MADM labeling scheme for cassettes inserted in forward direction. MADM experiments involving forward cassettes require that the mutant allele of a candidate gene must be linked to the T-G MADM cassette in order for mutant cells to be labeled in green upon a G2-X MADM event.

(D) MADM labeling scheme for cassettes inserted in reverse direction. MADM experiments involving reverse cassettes require that the mutant allele of a candidate gene must be linked to the G-T MADM cassette in order for mutant cells to be labeled in green upon a G2-X MADM event.

(E) Generation of recombinant MADM chromosomes. To genetically link a mutant allele of a candidate gene of interest to the corresponding chromosome containing the T-G MADM cassette (i.e., forward orientation), it is necessary to first cross mice bearing the T-G MADM cassette with mice bearing the mutant allele. Resulting F1 transheterozygous offspring are then backcrossed to mice homozygous for the T-G MADM cassette. In F2, recombinant offspring emerge from meiotic recombination events in the germline. These F2 recombinants now contain both the MADM cassette (in homozygous configuration) and the mutant allele.

(F) Calculation of meiotic recombination probability

$$\text{meiotic recombination} = \frac{B - A}{2}$$

(legend continued on next page)
in distinct colors with 100% accuracy (Zong et al., 2005; Hippenmeyer et al., 2010; Figure 1A; Figure S1).

The MADM approach enables unparalleled lineage tracing, and MADM-labeled cells can be assessed by histological means, physiological analysis, and optical imaging in vivo (Espinosa et al., 2009; Hippenmeyer et al., 2010; Liang et al., 2013; Gao et al., 2014; Joo et al., 2014; Riccio et al., 2016; Beattie et al., 2017; Henderson et al., 2019; Lv et al., 2019; Ortiz-Alvarez et al., 2019).

MADM technology represents one approach to probe genomic imprinting and the function of imprinted genes (Hippenmeyer et al., 2013; Laukoter et al., 2020b). MADM can be applied to create uniparental chromosome disomy (UPD; somatic cells with two copies of either the maternal or paternal chromosome) and visualize imprinting effects at morphological and transcriptional levels with single-cell resolution (Hippenmeyer et al., 2013; Laukoter et al., 2020a, 2020c).

One clinically relevant application of MADM is the tracing of tumor growth upon sparse or clonal ablation of tumor suppressor genes and/or to assay for the effects of therapeutic agents. As such, MADM has been used for the analysis of tumor formation and the delineation of cancer cell origin at the single-cell level in the brain and distinct organs (Muzumdar et al., 2007, 2016; Liu et al., 2011; Gonzalez et al., 2018; Tian et al., 2020; Yao et al., 2020).

A current limitation of the MADM technology is that it can only be applied to study candidate genes located on chromosome 7 (chr7), chr11, and chr12 and distal to the Rosa26 locus on chr6, where MADM cassettes have been introduced (Zong et al., 2005; Hippenmeyer et al., 2010, 2013). Thus, less than 25% of all genes in the mouse genome can be subjected to MADM analysis as described above. Here, we overcome this constraint and expand MADM technology to all mouse autosomes. We provide validation of all MADM reporters and quantitative assessment of the efficacy of MADM labeling in a variety of organs and tissues and a number of clinically relevant stem cell niches across the entire mouse. Furthermore, we use engineered MADM chromosomes to systematically determine sister chromatid segregation patterns in several somatic cell lineages. Our analysis revealed that sister chromatid segregation patterns in mitotic progenitor cell divisions are highly biased in a chromosome-specific manner and are further affected by cell type in vivo.

**RESULTS**

**Expansion of MADM to all mouse autosomes**

For MADM, two reciprocally chimeric marker genes need to be targeted to identical loci on homologous chromosomes (Zong et al., 2005). The chimeric marker genes (GT and TG alleles) consist of N- or C-terminal halves of the coding sequences for GFP (enhanced GFP) and red fluorescent protein (tdT) interspersed by an intron with the loxP site (Hippenmeyer et al., 2010; Figure 1A; Figure S1). Here, we expanded MADM to all 19 mouse autosomes with the goal to enable MADM for the vast majority, nearly genome-wide, of autosomal genes in the mouse genome. Mouse autosomes consist of only one chromosome arm (i.e., telocentric conformation). We thus rationalized that inserting the MADM cassettes as close as possible to the centromere would maximize the number of genes located distally to the MADM cassette insertion site for prospective MADM experiments (Hippenmeyer et al., 2010, 2013; Figures 1A and 1B).

To identify suitable sites for MADM cassette targeting, we applied a number of key criteria. The loci should (1) locate to intergenic regions to minimize the probability of disrupting endogenous gene function and (2) permit spatially and temporally ubiquitous and biallelic expression of the reconstituted GFP and tdT markers. To fulfill the first criteria, we mapped gene by gene the genetic landscape of the centromeric-most 20 Mbp of all autosomes using the UCSC Genome Browser (https://www.genome.ucsc.edu; GRCm38/mm10). Next, we assessed EST (expression sequence tag) expression patterns of the neighboring genes flanking the putative targeting sites and serving as proxy for the spatiotemporal extent of transgene expression. The final choice of the prime targeting loci (Figure 1B; Figures S2 and S3; Table S1) was based upon the most ideal combination of the above key criteria. In total, more than 20,000 protein-coding genes, corresponding to >96% of the entire annotated mouse genome (GRCm38/mm10), are located distally to the MADM targeting loci across all 19 autosomes (Table S1).

Next, we cloned the selected genomic targeting loci and inserted the MADM cassettes (Hippenmeyer et al., 2010) by homologous recombination in mouse embryonic stem cells (ESCs) (Figure S4; see STAR Methods for details). MADM cassettes were inserted in a centromere-to-telomere transcriptional direction (Figure 1B, forward) except for chr3, chr5, chr6, and chr15, which required opposite directionality (Figure 1B, reverse) in order to best fulfill our locus choice criteria. The directionality of reconstituted MADM marker gene transcription, upon interchromosomal recombination, has consequences for the coupling of mutant and wild-type genotypes with fluorescent labeling upon mitosis (Figures 1C and 1D). In order to genetically link a mutant allele of a candidate gene to the corresponding chromosome containing the MADM cassette, meiotic recombination in the germline can be used (e.g., Hippenmeyer et al., 2010; Laukoter et al., 2020b; Figures 1E and 1F). The probability for meiotic recombination that results in the linkage of the mutant allele with the MADM cassette can be estimated (Figure 1F) once the location (cM) of the mutant allele (genomic locus) has been determined by using, for example, the Mouse Genome Informatics (MGI) database (https://www.informatics.jax.org).

Homologous recombination frequencies in ESCs were relatively high for all selected loci (for some, >50%), hinting at an open chromatin structure that should be an advantage for...
Figure 2. MADM labeling pattern in different organs and stem cell niches
(A) Overview of MADM labeling (green, GFP; red, tdT; yellow, GFP/tdT) in MADM-19GT/TG in combination with Hprt-Cre at P21. Diverse tissues/organs including eye, brain, lung, spinal cord, kidney, spleen, liver, heart, and thymus are illustrated.

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prospective mitotic Cre-mediated interchromosomal recombination. Next, chimeric founder mice were generated by blastocyst injection. Homozygous MADM$^{GT/GT}$ and MADM$^{CAG/CAG}$ stock lines were established upon successful germline transmission of the respective MADM cassettes (Figure S4) by using specific genotyping primers (Table S2).

Ubiquitous labeling in all MADM reporter lines across different organs
We systematically analyzed the MADM labeling pattern upon Cre-mediated interchromosomal recombination in all MADM lines (Figure S4E). First, we crossed all MADM$^{GT/GT}$ lines to mice that carry the Cre transgene within the X-linked Hprt (encoding hypoxanthine guanine phosphoribosyl transferase) genomic locus. The Hprt-Cre driver is spatiotemporally ubiquitously and constitutively expressed (Tang et al., 2002). In female mice, inactivation of the X chromosome results in mosaic Cre expression from the Hprt locus and thus highly variable MADM labeling patterns (Zong et al., 2005; Hippenmeyer et al., 2013). We therefore analyzed male experimental MADM (MADM$^{GT/GT}$;HprtCre$^{+/+}$) animals for a first pass comparative assessment. We detected MADM labeling in all organs analyzed—including brain, spinal cord, eye, heart, lung, liver, kidney, thymus, and spleen (Figure 2A)—and in all MADM lines. The relative recombination frequency, at least at this superficial qualitative level, appeared to correlate in distinct selected organs across all 19 MADM lines (Figures 2 and S5–S9).

MADM labeling in clinically relevant adult stem cell niches
We next evaluated a number of stem cell niches with high clinical relevance. Because it is important to know the approximate scale of labeling for determining sample size in a MADM experiment, we chose two different MADM models in combination with Hprt-Cre driver for these analyses, as follows: MADM-19 that shows relatively dense MADM labeling and MADM-4 that represents one of the sparsest MADM.

First, we focused on the mammary gland (Figure 2B), the site where breast cancer initiates. The mammary gland harbors two types of unipotent stem cell lineages, namely, the K14+ myoepithelial (or basal) cells and the K8+ luminal cells (Van Keymeulen et al., 2011). Myoepithelial and luminal stem cell populations are derived from a multipotent progenitor during embryonic development (Wuidart et al., 2018), become activated, start to proliferate, and expand the hair follicle deep down into the dermis. Progenitors located at the bottom of the activated follicle (anagen follicle) form the matrix, from which epithelial hair lineages are specified (Hsu et al., 2014). Such differentiated hair lineages comprise the companion layer (CP), distinct layers of inner root sheath (IRS), and cuticle and cortex of the hair shaft (HS), as well as the innermost hair layer the medulla (Me). Once hair regeneration is completed, the follicles undergo a destructive phase (catagen) and enter the quiescent resting phase again. In the skin of MADM-19$^{GT/GT}$; HprtCre$^{+/+}$ and MADM-4$^{GT/GT}$;HprtCre$^{+/+}$ mice, we observed prominent MADM labeling in all compartments of the hair follicle and importantly in the hair follicle stem cells (Figures 2D and S7C).

Next, we analyzed MADM labeling in the small intestine that represents another critical model for the study of stem-cell-mediated regeneration but also intestinal cancer (Barker et al., 2009). Intestinal stem cells replenishing the epithelium are LGR5+ and located in the crypt base (Barker et al., 2007). They are intermingled with secretory Paneth cells and divide constantly in order to rejuvenate the epithelial cell layer on the villus surface. Interestingly, LGR5+ stem cells mostly divide symmetrically and undergo neutral competition within the crypt, thus driving the crypt toward monoclonality (Snippert et al., 2010). In order to evaluate the potential for MADM-based lineage tracing, the study of loss of gene function, and analysis of stem cell compartmental and functional behavior in vivo, we used adult lactating 4-month-old female MADM-19$^{GT/GT}$;HprtCre$^{+/+}$ (Figure 2B) and MADM-4$^{GT/GT}$;HprtCre$^{+/+}$ (Figure S7A) mice and could readily detect GFP+ (green), tdT− (red), and GFP+/tdT+ (yellow) cells in both K14+ basal and K8+ luminal cells. Next, we analyzed pancreatic epithelial cells that can be divided into secretory acinar cells and ductal epithelial cells. Although the tumor cell of origin for pancreatic cancer remains controversial, oncogenic drivers can trigger pancreatic ductal adenocarcinoma (PDAC) from both ductal and acinar cells (Ferreira et al., 2017; Lee et al., 2019). In both, MADM-19$^{GT/GT}$;HprtCre$^{+/+}$ and MADM-4$^{GT/GT}$;HprtCre$^{+/+}$ mice at postnatal day 21 (P21), we noticed MADM-labeled cells in the acinus and duct within the pancreas (Figure 2C and S7B).

Hair follicles are a prime stem cell model for the study of tissue regeneration but also for skin cancer including melanoma (Sun et al., 2019). Hair follicles are appendages of the epidermal lineage and undergo cycling rounds of stem cell activation in order to generate new hair (Fuchs and Nowak 2008). The stem cells are located in the secondary hair germ (2° HG) and lower part of the bulge (Bu) of a resting follicle (telogen follicle) (Figure 2D). They become activated, start to proliferate, and expand the hair follicle deep down into the dermis. Progenitors located at the bottom of the activated follicle (anagen follicle) form the matrix, from which epithelial hair lineages are specified (Hsu et al., 2014). Such differentiated hair lineages comprise the companion layer (CP), distinct layers of inner root sheath (IRS), and cuticle and cortex of the hair shaft (HS), as well as the innermost hair layer the medulla (Me). Once hair regeneration is completed, the follicles undergo a destructive phase (catagen) and enter the quiescent resting phase again. In the skin of MADM-19$^{GT/GT}$; HprtCre$^{+/+}$ and MADM-4$^{GT/GT}$;HprtCre$^{+/+}$ mice, we observed prominent MADM labeling in all compartments of the hair follicle and importantly in the hair follicle stem cells (Figures 2D and S7C).

(B) Schematic (left) and MADM labeling (middle/right; green, GFP; red, tdT; yellow, GFP/tdT) in mammary gland of lactating MADM-19$^{GT/GT}$;HprtCre$^{+/+}$ female at 4 months of age. Basal/myoepithelial (middle) and luminal (right) cells are stained with antibodies against K14 and K8 (white), respectively.

(C) Schematic (left) MADM labeling (right; green, GFP; red, tdT; yellow, GFP/tdT) in MADM-19$^{GT/GT}$;HprtCre$^{+/+}$ pancreas, acinus, and duct, at P21. Epithelial cells are visualized by antibody staining against β-catenin (white; β-Cat). Acinar cells are identified by the presence of intracellular secretory granules.

(D) Schematic (left) and MADM labeling (middle/right; green, GFP; red, tdT; yellow, GFP/tdT) in telogen (middle) and anagen (right) hair follicles in MADM-19$^{GT/GT}$;HprtCre$^{+/+}$ at P21 (telogen) and P28 (anagen), Bu, bulge; 2° HG, secondary hair germ; SG, sebaceous gland; IRS, inner root sheath; CP, companion layer; ORS, outer root sheath; Mx, matrix.

(E) Schematic (left) and MADM labeling (right; green, GFP; red, tdT; yellow, GFP/tdT) in small intestine in MADM-19$^{GT/GT}$;HprtCre$^{+/+}$ at P21. Epithelial cells are visualized by antibody staining against β-catenin (white; β-Cat). Asterisk marks a Paneth cell, identified by the presence of intracellular granules. TAC, transit-amplifying cell; LGR5, leucine-rich repeat-containing G-protein coupled receptor 5. Nuclei were stained using DAPI. Scale bar: 50 μm (A) and 20 μm (B–E).

See also Figures S5–S8.

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cell behavior in the intestinal crypts, we dissected the intestine of MADM-19G/(G:HprtCre/; and MADM-4G/(G:HprtCre/ mice at P21. We observed MADM-labeled cells in all compartments of the intestinal unit, including the villus and the crypt (Figure 2E and S7D).

Lastly, we validated one of the MADM reporters in a disease-relevant setting. We used MADM-18 lines to examine the effect of clonal loss of Apc (adenomatous polyposis coli). Apc functions as a tumor suppressor and mutations in Apc cause hereditary and sporadic human bowel cancers upon loss of heterozygosity (Behrens et al., 1998; Fodde et al., 2001). To mimic the intestinal tumor initiation, we generated genetic mosaic mice harboring green Apc−/− cells by using an Apc-flox allele (Cheung et al., 2010) and red wild-type cells in an otherwise heterozygous environment (Figure 3). At 3 months of age, MADM-18G/(G:HprtCre/; control mice showed several red- or green-labeled normal crypt-villus units (Figures 3A−3D). Note that MADM labeling within crypt-villus units appeared exclusively unicolor, reflecting monoclonality due to stochastic competition between dividing intestinal stem cells (Snippert et al., 2010). In contrast, all MADM-18G/(G:Apc Cre/−;HprtCre/; experimental mice, in which Apc−/− cells are labeled in green and Apc+/+ cells in red by design, displayed one or several green-labeled cancerous lesions (adenomas), derived from Apc−/− stem cells, in their small intestine and colon (Figures 3E−3G; Barker et al., 2009). Yet, we did not detect any tumors derived from red control cells in MADM-18G/(G:Apc Cre/−;HprtCre/; experimental mice (Figure 3D). Antibody staining against phosphohistone H3 (P-H3) confirmed that normal crypt-villus units display proliferation only within the crypt compartment but not within the villus epithelium (Figures 3B, 3C, and 3F). In contrast, adenomas derived from green Apc−/− cells in mosaic mice contained proliferating tumor cells in regions outside the crypt compartment (Figure 3F) as previously reported (Schepers et al., 2012). In summary, we validated one of the MADM lines for functional genetic mosaic analysis in the context of the Apc model (Figures 3H and 3I) for tumor initiation and growth.

Genomic imprinting phenotypes in liver cells with UPD
MADM can create UPD (Figure 4A) to analyze imprinting phenotypes at the single-cell level that result from the imbalanced expression of imprinted genes (Hippenmeyer et al., 2013; Lautkot et al., 2020b, 2020c; Paulet al., 2021). Prominent imprinting phenotypes have been observed in the liver where, for instance cells with MADM-induced paternal UPD of chr7 exhibit overgrowth (Hippenmeyer et al., 2013), in accordance with the kinship hypothesis that stipulates a major growth regulatory function of genomic imprinting (Haig 2004; Tucci et al., 2019). Because imprinted genes are located throughout the genome, we analyzed the liver in all 19 MADM reporters in combination with Hprt-Cre (Figures 4B−4U) for potential imprinting phenotypes. We readily observed the growth advantage of hepatocytes with paternal UPD of chr7 (Figures 4H and 4V) but also noticed that cells with paternal UPD of chr11 (Figures 4L and 4V) and chr17 (Figures 4R and 4V) showed significant overrepresentation in comparison to cells with maternal UPD. The maternally expressed growth inhibitory imprinted genes Grb10 and Igf2r are located on chr11 and chr17, respectively. Thus, although overexpression of growth-promoting Igf2r in UPD of chr7 leads to paternal growth dominance (Hippenmeyer et al., 2013), the absence of growth-antagonizing Grb10 or Igf2r (Smith et al., 2006) may result in the growth advantage of cells with paternal UPD of chr11 or chr17. We did not find significant UPD-mediated phenotypes in the liver of any other MADM (Figures 4B−4U).

Quantification of recombination efficiency of all MADM chromosomes
To systematically determine recombination frequencies comparatively in all MADMs, we quantified the absolute density of MADM-labeled neurons in the neocortex of P21 mice by using the Emx1-Cre driver (Figures 5A, 5B, and S9). We first assessed MADM labeling originating from G2-X events and quantified the numbers of green GFP+ and red tdT+ projection neurons per cubic millimeter (Figures 5A and 5B). The relative number of red tdT+ versus green GFP+ projection neurons was not significantly different across MADM lines (Figures 5B; Table S3). We classified the MADM reporters into three categories, as follows: (1) sparse (<25 cells/mm3), (2) intermediate (25−100 cells/mm3), and (3) dense (>100 cells/mm3). Because all MADM-targeting loci have been selected by using the same criteria, the origin of the variability in recombination frequency across all MADMs is currently not clear. In mice, the pairing of homologous chromosomes in somatic cells is infrequent and under tight regulation, unlike in the fruit fly Drosophila (Apte and Meller 2012). Thus, the individual dynamic organization of different homologous chromosomes within the nucleus may result in distinct probabilities of Cre-mediated interchromosomal recombination. It is also important to mention that insertion of the MADM cassettes at more distal locations in the same chromosome could lead to a distinct recombination probability. In any case, all MADM reporters do work as predicted from the MADM principle (Figures 1 and S1) in all organs analyzed (Figures 2, 3, 4, 5, and S5−S9). Importantly, even the sparsest MADM lines (including MADM-4 [Hansen and Hippenmeyer, unpublished observation] and MADM-5 [Takeo et al., 2021]) reliably permit functional genetic mosaic analysis of candidate genes.

MADM reveals chromosome-specific biases in mitotic sister chromatid segregation patterns
Previous in vitro studies have used mitotic recombination in ESCs to monitor the randomness of sister chromatid segregation patterns upon mitosis (Liu et al., 2002; Armakolas and Klar 2006). Against common belief, initial results indicated that sister chromatids derived from a homologous pair of chromosomes did not segregate randomly to daughter cells. Instead, G2-X segregation (two recombinant chromosomes segregate away from each other), thus reflecting one particular pattern of sister chromatid segregation, prevailed in ESCs for chr7 (Liu et al., 2002). Furthermore, ESC-derived endoderm cell lines exhibited complete bias toward G2-X (Armakolas and Klar 2006). Conversely, ESC-derived neuroectoderm cell lines never showed G2-X (Armakolas and Klar 2006). Although these results indicated that cell type may influence the selective segregation of sister chromatids, such a hypothesis is based on the analysis of only one chromosome and has not been examined in the context of intact
Figure 3. Apc-tumor model at single-cell resolution using the MADM-18 line

(A) Schematic representation of MADM labeling (green, GFP; red, tdT) and respective cellular genotypes in wild-type MADM-18<sup>G1T/G0</sup>:Hprt<sup>Cre</sup> mice.

(B and C) P-H3 staining (white) in small intestine in MADM-18<sup>G1T/G0</sup>:Hprt<sup>Cre</sup> mice at 3 months of age. (B) Overview of unicolor (monoclonal) green wild-type crypt-villus units with insets highlighting non-proliferative villus epithelium (i) and a proliferative cell within the crypt (white arrow) (ii); (C) overview and unicolor (monoclonal) red wild-type crypt-villus units with insets highlighting non-proliferative villus epithelium (iii) and a proliferating cell within the crypt (iv).

(D) Quantification of the percentage of intestinal structures displaying MADM labeling. Data obtained from n = 3 male MADM-18<sup>G1T/G0</sup>:Hprt<sup>Cre</sup> mice at 3 months of age.

(E) Schematic representation of MADM labeling (green, GFP; red, tdT) and respective cellular genotypes in genetic mosaic MADM-18<sup>G1T/G0,Apc</sup>:Hprt<sup>Cre</sup> mice.

(F) P-H3 staining (white) in small intestine in male MADM-18<sup>G1T/G0,Apc</sup>:Hprt<sup>Cre</sup> mice at 3 months of age with insets highlighting a proliferating adenoma cell at the boundary to the non-proliferative villus epithelium (white arrow) (i), proliferating adenoma cells within the tumor (white arrows) (ii), non-proliferative normal villus epithelium (iii), and proliferative cells within a normal crypt compartment (iv).

(G) Quantification of the percentage of intestinal structures displaying MADM labeling. Green Apc<sup>/−/−</sup> cells display 100% transformation and tumor initiation, whereas red wild-type cells solely give rise to normal crypt-villus-units. Data obtained from n = 3 male MADM-18<sup>G1T/G0,Apc</sup>:Hprt<sup>Cre</sup> mice at 3 months of age.

(H and I) Summary of MADM labeling in small intestine of control MADM-18<sup>G1T/G0</sup>:Hprt<sup>Cre</sup> (H) and genetic mosaic MADM-18<sup>G1T/G0,Apc</sup>:Hprt<sup>Cre</sup> (I). Note that in the mosaic, red wild-type cells give rise to normal crypt-villus units and green Apc<sup>/−/−</sup> cells initiate tumor development and subepithelial invasion of adenomas.

Nuclei were stained using DAPI. Scale bar: 100 μm (B, C, and F) and 25 μm (i–iv).
tissue. To this end, we used the entire library of MADM-rendered homologous chromosomes to systematically trace sister chromatid segregation patterns of all 19 mouse autosomes in a number of somatic cell lineages in vivo.

We exploited the inimitable feature provided by the MADM principle (Figures 5C and S1)—the differential fluorescent labeling of pairs of nascent sister cells upon mitosis that is dependent on how recombinant chromosomes segregate during cell division. G2-X segregation of recombinant MADM chromosomes can be unambiguously identified by the presence of red and green cells. However, G2-Z segregation, producing yellow cells, cannot be identified without ambiguity because G1 and/or postmitotic G0 events also result in yellow cells (Zong et al., 2005; Figures 5C and S1). Therefore, we capitalized upon the power of unequivocal G2-X identification—but also taking into consideration the caveat of yellow cells potentially reflecting a mix of G2-Z and G1/G0—and defined "yellow-green-red-index" (YGRI) as a proxy for sister chromatid segregation patterns (Figure 5D).

First, we systematically determined the YGRI of pyramidal projection neurons in the P21 neocortex for all 19 MADM reporters in combination with the Emx1-Cre driver (expressed in cortical progenitor cells and thereby limiting G0 events) (Figure 5E; Table S3). Contrary to the prediction and expectation based on cell culture data (no G2-X in neuroectodermal lineage [Armakolas and Klar 2006]), we always observed G2-X events. Interestingly, the YGRI values ranged from ~1 for MADM-2 and MADM-17 to ~10 for MADM-15 (Figure 5E, top). The values of the YGRI did not correlate with the sizes of the respective MADM chromosomes. Next, we compared the values of the YGRI with the absolute recombination frequencies (RFI, recombination frequency index), i.e., density of G2-X MADM labeling as indicated in Figure 5B. In the ranking plot in which the axes indicate YGRI versus RFI, there was no apparent correlation (Figure 5E, bottom) of YGRI with RFI. In summary, we detected highly distinct YGRI for different MADM chromosomes, suggesting distinct sister chromatid segregation patterns in the cortical Emx1+ projection neuron lineage.

Previous studies implicated left-right dynein (LRD)—the only protein thus far in a mammalian cell culture system—in the selective sister chromatid segregation process (Armakolas and Klar 2007). Intriguingly, mutations in the gene (Dnah11) encoding LRD causes randomization of left-right laterality in mice (half of the animals develop with mirror-imaged visceral organs) (Hummel and Chapman 1959; Supp et al., 1997). Based on the above findings from cell culture, we next assessed whether Dnah11 could play a role in biased chromosome-specific sister chromatid in vivo as well. We crossed the MADM system to iv (situs inversus) (a spontaneous mutation in Dnah11 [Hummel and Chapman 1959]) background and analyzed YGRI of chr7, chr12, and chr18 in cortical Emx1+ projection neurons (Figure 6A). However, contrary to the in vitro data (Armakolas and Klar 2007), we could not observe randomization (i.e., a drop of YGRI value to 1) of sister chromatid segregation for the three tested chromosomes (chr7, chr12, and chr18). These data suggest that cultured differentiated ESC lineages and cortical excitatory neurons in vivo differ in mechanisms of biased sister chromatid segregation.

**Chromosome-specific biases of sister chromatid segregation differ in distinct cell types**

To determine the possible influence of cell type on biased, chromosome-specific, sister chromatid segregation patterns, we first analyzed Emx1+ cortical astrocytes and hippocampal CA1 pyramidal cells. The YGRI for cortical astrocytes was markedly different from the YGRI for cortical projection neurons or hippocampal CA1 pyramidal cells for a representative set of 10 MADM chromosomes analyzed (Figures 6B and 6C). Next, we introduced Nestin-Cre to label neural lineages beyond forebrain projection neurons and astrocytes. We focused on cerebellar Purkinje cells and determined the YGRI. Strikingly, the YGRI for Purkinje cells was also markedly different in most MADMs compared to the YGRIs for cortical projection neurons and astrocytes and hippocampal CA1 pyramidal cells (Figures 6B and 6C).

Finally, we assessed sister chromatid segregation patterns for a non-neural somatic cell type. We focused on T cells (CD3+) and B cells (CD19+) within the hematopoietic lineage and determined the YGRI for six different MADM chromosomes by fluorescence-activated cell sorting (FACS) analysis (Figure 6D). Although the distinct MADM chromosomes showed different YGRI values, the YGRI for T cells in comparison to B cells was not significantly different for all six chromosomes analyzed. No significant
A MADM cerebral cortex images

B Classification of MADM lines
- Sparse lines: density < 25 cells/mm²
  - M4, M3, M13, M6, M1, M9
- Intermediate lines: 25 < density < 100 cells/mm²
  - M5, M15, M10, M14, M8, M2, M16, M17
- Dense lines: density > 100 cells/mm²
  - M12, M7, M18, M19, M11

C MADM scheme for Chr segregation

D Index calculation
Yellow Green Red Index (YGRI) = \frac{\# yellow cells}{average (\# green, \# red cells)}

E YGRI in cortical pyramidal neurons

(legend on next page)
correlation could be established when the YGRI of T/B cells was compared to the YGRI of the neural lineages. Altogether, these data indicate that the highly biased and chromosome-specific sister chromatid segregation patterns are further affected by cell type in somatic cell lineages in vivo.

**DISCUSSION**

The analysis of gene function in multicellular systems in vivo requires quantitative and high-resolution experimental tools to analyze the cellular phenotype. Here, we expanded the MADM technology to enable, in principle, the genetic mosaic dissection of cell-autonomous gene function of most genes (>96%) across the entire mouse genome with single-cell resolution. Although functional genetic mosaic analysis clearly represents the most salient utility of MADM, we also extended the application spectrum and used MADM as a proxy to trace the randomness of mitotic sister chromatid segregation patterns upon somatic stem cell division. We first discuss these biological findings in a more general context before we elaborate on the overarching potential of the genome-wide MADM resource for future genetic mosaic analysis.

**Non-random mitotic sister chromatid segregation in mouse in vivo**

Asymmetric stem cell division requires the non-equivalent distribution of cell-fate determinants including proteins, mRNA, or intracellular organelles (Gönczy 2008; Knoblich 2008; Taverna et al., 2014). Recently, an intriguing model has been postulated whereby asymmetric cell division might also be promoted by differentiation of sister chromatids by epigenetic means, followed by selective segregation of “unequal” sister chromatids to daughter cells (Bell 2005; Armakolas et al., 2010; Yamashita 2013). However, experimental evidence supporting such a model in mice was so far obtained solely from in vitro studies in ESCs and derived lineages and only for one chromosome (chr7; Figure 7, left; Liu et al., 2002; Armakolas and Klar 2006).

In our study we systematically traced sister chromatid segregation patterns of the entire set of mouse autosomes in vivo. We observed that the prevalence of G2-X events, approximated in the value of YGRI, in the same cell type (cortical projection neurons) and by using identical an Emx1-Cre driver vastly differed, up to one order of magnitude for different chromosomes. However, the relative amount of G2-X segregation did not correlate with the absolute recombination frequency. Thus, high absolute recombination frequency does not predict a bias in recombinant sister chromatid segregation toward G2-X nor G2-Z. Previous work has postulated that it is highly unlikely that biased sister chromatid segregation may have evolved for a site-specific recombination system not indigenous to mouse cells (Armakolas et al., 2010). Yet, we note that the dynamic state of chromatin architecture may influence absolute recombination frequency (proximity of loxP sites). We also cannot exclude that the location of the genomic recombination loci, and thus the size of resulting recombinant sister chromatids, may influence segregation bias. Such a hypothesis may be tested in the future by systematic introduction of loxP sites at defined genomic distance intervals.

Our observation that distinct cell types show different YGRIs for the same chromosome could reflect a different recombination activity of respective Cre drivers in a particular cellular lineage. For our MADM-based analysis, we used Emx1- and Nestin-Cre drivers that are mostly active in dividing neural stem cells and turned off in postmitotic cells. The contribution of G0 recombination is thus expected to be minimal. Still, all YGRIs in neural lineages were >1, with some up to an order of magnitude higher indicating increasing rates of G2-Z segregation. However, a certain rate of G1 recombination (also producing yellow cells) that increase the YGRI besides G2-Z segregation may add to the overall YGRI. Although G1 recombination events did not occur in cultured ESCs (Liu et al., 2002; Armakolas and Klar 2006), we cannot currently exclude that interchromosomal recombination efficiency could be distinct in G1 versus G2 phases of the cell cycle for different cell types in vivo. However, for any given cell division cycle, the relative recombination events in G1 versus G2 should be the same. Thus, different YGRIs for different chromosomes must reflect chromosome-specific sister chromatid segregation patterns in genetically identical cells (here, Emx1+ cortical projection neurons). Perhaps most striking was the finding that YGRIs of 10 different chromosomes in astrocytes were rather constant and low, indicating a

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**Figure 5. Mitotic interchromosomal recombination efficiency and sister chromatid segregation patterns for all MADM reporters in cortical projection neurons**

(A) Representative images of MADM-labeling pattern (green, GFP; red, tdT; yellow, GFP:tdT) in cerebral cortex in three exemplary MADM lines in combination with the Emx1-Cre driver at P21. (Top) MADM-9GT/124Emx1Cre−: (middle) MADM-17GT/105Emx1Cre+: (bottom) MADM-19GT/104Emx1Cre+. Scale bar: 100 μm. (B) Classification of MADM lines. (Top) Sparse (<25 cells/mm²); (Middle) Intermediate (25–100 cells/mm²); (Bottom) Dense (>100 cells/mm²). Bars represent mean ± SEM. Data show M7, M11, and M19 (n = 5); M2, M3, M5, M6–M10, and M12–M18 (n = 6); M4 and M6 (n = 8); and M1 (n = 12 mice). (C) MADM principle illustrating G2-X and G2-Z segregation patterns. Upon Cre-mediated interchromosomal recombination at the loxP site in the MADM cassette in G2 phase of the cell cycle, recombinant chromosomes can either segregate together to the same daughter cell (G2-Z segregation; yellow, GFP:tdT and unlabeled cell) or each recombinant chromosome may segregate to distinct daughter cells (G2-X segregation; green, GFP+ and red tdT+ cell, respectively) upon mitosis. (D) Definition of yellow-green-red-index index (YGRI). The YGRI is calculated from the number of yellow cells divided by the average of green and red cells to compensate for G2-Z events that leads to labeling of only one daughter cell (yellow) and an (invisible) unlabelled cell. Note that yellow cells emerging from G1/G0 events contribute to the total number of yellow cells. (E) YGRI index in neuronal lineages. (Top) YGRI for cortical projection neurons in P21 neocortex of all 19 MADM reporter lines in combination with the Emx1-Cre driver. Note that (1) YGRI varies from 1 to 10 but is never below 1 and (2) YGRIs do not correlate with the sizes of the respective MADM chromosomes. Bars represent mean ± SEM. Data show M2, M3, M5, and M7–M19 (n = 6); M4 and M6 (n = 8); and M1 (n = 12). (Bottom) YGRI ranking in correlation (red line) to recombination frequency index (RFI). Note that MADM chromosomes with a high recombination frequency do not necessarily present high YGRI and vice versa. See also Figures S1 and S9.
A. Analysis of YGRI in Dnah11 KO in cortical pyramidal neurons

- In vivo
  - Induced mitotic recombination
  - Visual readout (single-cell MADM labeling)
  - Biased segregation patterns

Dnah11 knockout

- Biased segregation patterns remain

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B. Analysis of YGRI in different brain regions

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GFP / tdT

C. YGRI in different neuronal lineages

- Cortical pyramidal neurons
- Cortical astrocytes
- CA1 pyramidal neurons
- Cerebellar Purkinje neurons

D. YGRI in spleen

- MADM labeled spleen using Hprt-Cre driver:
  - White blood cell preparation
  - FACS
  - Gating: T cells (CD3+) B cells (CD19+)
  - YGR index analysis

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uniformly high relative frequency for G2-X events in astrocyte progenitors, which is in stark contrast to the values for the same chromosomes in projection neurons that emerge also from the same (Emx1+) stem cell lineage. Thus, sister chromatid segregation appears highly biased in a chromosome-specific manner in mitotic cortical Emx1+ progenitors. Furthermore, the rank orders of YGRI for each chromosome in different cell types were not the same, suggesting that the bias of sister chromatid segregation patterns results from a complex combination of chromosome and cell-type-specific mechanisms (Figure 7, right).

Previous studies found that cultured ESC clones that were differentiated into an neuroectoderm lineage never showed G2-X segregation (Liu et al., 2002; Armakolas and Klar 2006). These findings are in stark contrast to our in vivo results that demonstrate for all 19 mouse autosomes a substantial amount of G2-X segregation in at least 4 distinct neural cell lineages. Furthermore, our analysis of Dnah11 indicates that the involved molecular mechanisms likely differ (at least for chr7) when comparing cell culture to intact brain tissue. We cannot fully explain the cause of the differences in results obtained in cell culture and in vivo, but systemic and/or tissue-wide acting mechanisms could be involved (Knouse et al., 2018).

The phenomenon of biased sister chromatid segregation appears to be evolutionarily conserved (Pimpinelli and Ripoll 1986; Beumer et al., 1998). In asymmetrically dividing male germ line stem cells in Drosophila, sister chromatids of X and Y, but not autosomes, are segregated non-randomly (Yadlapalli and Yamashita 2013). In such a context, SUN-KASH proteins, proposed to anchor sister chromatids to the centrosome, seem to be involved, besides regulators of DNA methylation (Yadlapalli and Yamashita 2013). Although the underlying molecular mechanisms may or may not be conserved, it will be intriguing to assess the physiological function in future studies and experimentally approach the hypothesis postulating that biased sister chromatid segregation could be a mechanism to instruct the cell fate of nascent daughter cells during asymmetric stem cell division (Bell 2005; Armakolas et al., 2010; Yadlapalli and Yamashita 2013). Because MADM enables both clonal lineage tracing with concurrent genetic manipulation, such an approach promises high potential to systematically address the physiological role of biased sister chromatid segregation in the future.

**Genome-wide MADM mice library for single-cell genetic mosaic analysis**

**Genetic dissection of cell-autonomous gene function and system-wide effects**

The MADM technology enables a variety of genetic in vivo paradigms to study a broad spectrum of cell and developmental processes (Zong et al., 2005; Luo 2007; Muzumdar et al., 2007; Hippenmeyer 2013; Hippenmeyer et al., 2013). One exclusive application of the MADM system is the feature enabling the genetic dissection of the relative contributions of cell-autonomous and extrinsic systemic and/or tissue-wide components to the overall cellular phenotype upon the loss of candidate gene function (Hansen and Hippenmeyer 2020). In fact, single-cell phenotypes in classical conditional or full knockout mutants often reflect a combination of both cell-autonomous gene function and environment-derived cues that may remedy or exacerbate any observed phenotype. It is thus important to qualitatively and quantitatively determine the relative contribution of the intrinsic and extrinsic components to the overall loss of the gene function phenotype. To this end, the MADM system offers an unmatched experimental solution. The candidate gene function can be either ablated in a very sparse mosaic (or single clones) or tissue wide in all cells. Yet, in both paradigms, single-cell MADM labeling enables the high-resolution quantitative phenotypic analysis (Joo et al., 2014; Beattie et al., 2017; Laukoter et al., 2020b; Takeo et al., 2021). The MADM lines in conjunction with the above paradigms thus potentially permit the systematic dissection of the level of cell autonomy of any gene function in a given tissue, provided appropriate Cre driver lines exist. Insights at the single-cell resolution as obtained from MADM-based approaches in combination with systematic candidate gene interrogation (Beattie et al., 2017; Laukoter et al., 2020b) likely will have implications for our general understanding of diseases including neurodevelopmental disorders (D’Gama and Walsh 2018; Jayaraman et al., 2018; Buchsbaum and Capello 2019; Pinson et al., 2019; Subramanian et al., 2020).

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**Figure 6. Sister chromatid segregation patterns based on YGRI in Dnah11 knockout (KO) and in different somatic cell lineages**

(A) (Left) Summary of YGRI analysis in selected MADM reporters with Dnah11 deletion (n iv mice). (Right) YGRI index for cortical projection neurons in P21 neocortex in MADM-7, MADM-12, and MADM-18 reporter lines in combination with the Emx1-Cre driver in control and Dnah11 KO (iv) mice. Note that a decrease of YGRI to 1 would indicate random sister chromatid segregation but that the YGRI was not decreased upon Dnah11 mutation. Bars represent mean ± SEM. Data show n = 3 mice for each genotype.

(B) Representative confocal microscopy images at P21 with MADM labeling (GFP, green; tdT, red) from selected MADM reporters in combination with a Nestin-Cre (cerebellum) or Emx1-Cre (cerebral cortex and hippocampal CA1 area) driver used for quantifications in Figures 5C and 6A. Arrows indicate Purkinje cells (Pcs), cortical pyramidal neurons (Pys), and CA1 pyramidal neurons (CA1 Pys). Scale bar: 100 μm.

(C) YGRI for selected MADM reporters in different neuronal lineages at P21. YGRI of cortical astrocytes and hippocampal CA1 pyramidal cells derived from Emx1+ progenitors and cerebellar Purkinje cells derived from Nestin+ progenitors significantly differ from the YGRI of cortical pyramidal neurons for most MADM chromosomes analyzed. Values represent mean ± SEM. Data show pyramidal neurons (n = 6); cortical astrocytes (n = 8); CA1 pyramidal neurons (n = 5), M5, M7, M8, M11, M12, M16, M18, and M19 (n = 6); M10 (n = 7); M17 (n = 8); cerebellar Purkinje cells M7, M8, M11, M16, M17, and M19 (n = 3); M12 and M18 (n = 4); M10 (n = 5) and M5 (n = 6 mice).

(D) (Left) White blood cell preparations from spleen in MADM reporters with Hpert-Cre at P21 were subjected to FACS. The number of green GFP+, red tdT+, and yellow GFP+/tdT+ CD3+ T cells (black) and CD19+ B cells (blue) were quantified. (Right) YGRI for six different MADM chromosomes including sparse (MADM-4), intermediate (MADM-8, MADM-15, and MADM-17), and dense (MADM-18 and MADM-19) lines. The different MADM recombinant chromosomes displayed distinct YGRI but YGRI for T cells and B cells was not significantly different for all MADM chromosomes analyzed. Bars represent mean ± SEM. Data show M8 and M10 (n = 3); M4, M17, and M19 (n = 4); M15 (n = 5 mice). Welch’s unequal variances t test, p_{M4} = 7.5E−01, p_{M8} = 7.9E−01, p_{M10} = 7.7E−01, p_{M15} = 6.4E−01, p_{M17} = 9.8E−01, p_{M19} = 5.0E−01.
One MADM application includes the property to generate cells with UPD and thus enable the study of imprinting phenotypes at a single-cell level (Hippenmeyer et al., 2013; Laukoter et al., 2020a, 2020c). In fact, technical limitations so far only allowed the investigation of UPD at the whole-animal level but lacked the resolution to obtain insights at the cellular level (Pauler et al., 2021). It will be revealing in future studies to systematically probe the cell-autonomous consequences of UPD at a single-cell level and without inducing global changes in imprinted gene expression affecting the whole animal. The library of all 19 MADM reporters will in principle enable the systematic analysis of UPD-associated cellular phenotypes in any organ, tissue, and cell type in the mouse. Importantly, the analysis of candidate gene function, i.e., loss-of-function phenotypes, can be separated from UPD-mediated imprinting phenotypes by reverse MADM breeding schemes (Hippenmeyer et al., 2010, 2013; Joo et al., 2014; Beattie et al., 2017; Laukoter et al., 2020b).

One possible limitation or confounder for the interpretation of MADM-based cell labeling and gene dosage is cellular polyplody (Øvrebø and Edgar 2018). It will thus be crucial in the future to carefully analyze in organs with polyploid cells (1) whether more than one individual recombination event can occur and (2) at which frequency. Depending on the probability of multiple recombination events, the color/tone of the overall MADM labeling, and thus gene dosage, could be distinct from the above predicted scenario for UPD and potentially offer an assay for studying gene dosage across multiple scales. In a broader context, because UPD in humans is associated with a variety of diseases (Feinberg 2007; Tuna et al., 2009; Yamazawa et al., 2010; Buiting et al., 2016) MADM-based analysis will also contribute to our general understanding of the underlying etiology of imprinting disorders at the single-cell level.

**Analysis of cellular competition at single-cell level in health and disease**

MADM can be exploited for the study of cellular competition in a developmental context (Joo et al., 2014; Takeo et al., 2021). For
instance, when the TrkC neurotrophin receptor is removed sparsely with MADM from just a few individual Purkinje cells in the cerebellum, their dendrites have fewer and shorter branches. In contrast, when TrkC is ablated from all Purkinje cells, the dendrite trees look normal. Thus, a competitive mechanism could be involved whereby the shape of the dendrite tree depends on relative differences in neurotrophin/TrkC signaling between Purkinje cell neighbors (Joo et al., 2014). Purkinje cell dendritic arbors have also been shown to depend on GluD2- and Cbln1-mediated competitive interactions (Takeo et al., 2021). Cell competition has not only been implicated in cell morphogenesis but also extensively studied in a variety of contexts. Cell competition is particularly critical for overall tissue homeostasis during growth and regeneration but also for cell mixing and tissue invasion in cancer (Merino et al., 2016; Brás-Pereira and Moreno 2018; Madan et al., 2018; Ellis et al., 2019). With the availability of MADM for all mouse autosomes, the phenomenon of cell competition can be studied holistically and for virtually any candidate gene function associated with it in diverse biological contexts in health and disease.

Although MADM technology is versatile and offers multiple applications (discussed above), the method also has certain limitations and alternative systems may be considered depending on the biological question and context (Garcia-Gonzalez et al., 2020; Hansen and Hippenmeyer 2020). For instance, the expression of the two markers relies on a single-copy insertion and thus the expression level may be lower than, for example, that in viral-based systems. Optimization of the MADM reporter expression by using amplification systems such as, for example, the TIGRE (Madsen et al., 2015; Daigle et al., 2018) might offer a solution. Unlike TIGRE-based systems, MADM currently cannot be used to monitor physiological processes, but creating split transgenes of diverse reporters in future could enable such applications in clonally related cells in wild-type and potentially in a mutant context. Although MADM can be used in combination with temporal TM/CreER induction to label individual clones emerging from dividing stem cells, the system cannot be used to sparsely label and genetically manipulate postmitotic cells. Also, MADM currently cannot be easily used for the overexpression of transgenes in a mosaic setting. However, a number of recently developed systems—such as MASTR (Lao et al., 2012), Dual ifgMosaic (Pontes-Quero et al., 2017), iSuRe-Cre (Fernández-Chacón et al., 2019), MADR (Kim et al., 2019), or BATTLE (Kohara et al., 2020) among others—allow for sparse and tunable mosaic labeling and/or genetic manipulation including overexpression of transgenes in postmitotic cells in a variety of cellular contexts. However, a clear advantage over the above methods and special property of MADM relies on the built-in control with two distinct fluorescent colors for mutant analysis and the functional assessment of gene dosage. Thus, MADM can be used for unparalleled comparative mutant versus heterozygote and control analysis at once and at a single-cell resolution in any given tissue in situ.

Lastly, MADM technology based on gene targeting in ESCs is currently available only in mice. Future expansion of the system to other species by endonuclease-mediated transgenesis like CRISPR-Cas9 technology can be anticipated. Altogether, the genome-wide MADM resource presented in this study likely will catalyze the genetic dissection of cell-autonomous gene function and thus molecular mechanisms with single-cell resolution across a broad spectrum of biological questions in health and disease.

STAR METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.celrep.2021.109274.

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AUTHOR CONTRIBUTIONS


DECLARATION OF INTERESTS

The authors declare no competing interests.
REFERENCES


competition between symmetrically dividing Lgr5 stem cells. Cell 143, 134–144.


## STAR METHODS

### KEY RESOURCES TABLE

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RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Simon Hippenmeyer (simon.hippenmeyer@ist.ac.at).

Materials availability
All published and inaugural reported reagents and mouse lines will be shared upon request within the limits of the respective material transfer agreements. All MADM lines will be made publicly available through The European Mouse Mutant Archive (EMMA) and distributed from the University of Veterinary Medicine in Vienna or the Institute of Science and Technology Austria in Klosterneuburg.

Data and code availability
This study did not generate code and all data have been presented in Figures and Supplemental Figures. Original images will be made available upon request.
EXPERIMENTAL MODEL AND SUBJECT DETAILS

Generation, breeding and husbandry of mouse lines

Experimental procedures were discussed and approved by the institutional ethics and animal welfare committees at IST Austria, Stanford University, and at University of Veterinary Medicine Vienna in accordance with good scientific practice guidelines and national legislation (license number: IST Austria: BMWFW-66.018/0007-ll/3b/2012 and BMWFW-66.018/0006-WF/V/3b/2017; University of Veterinary Medicine Vienna: BMWFW-68.205/0023-ll/3b/2014 and BMBFW-68.205/0010-V/3b/2019). Mice with specific pathogen free status according to FELASA recommendations (Mählé Convener et al., 2014) were bred and maintained in experimental rodent facilities (room temperature 21 ± 1°C; relative humidity 40%–55%; photoperiod 12L:12D). Food (V1126, Ssniff Spezialitäten GmbH, Soest, Germany) and tap water were available ad libitum.

Mouse lines with MADM cassettes inserted in Chr. 7 (Hippinenmeyer et al., 2013), Chr. 11 (Hippinenmeyer et al., 2010), and Chr. 12 (Hippinenmeyer et al., 2013), Emx1-Cre (Gorski et al., 2003), Nestin-Cre (Petersen et al., 2002), Hprt-Cre (Tang et al., 2002), Apc-flox (Cheung et al., 2010), and iv mice [Dnah11 mutation (Hummel and Chapman 1995)] have been described previously. Nestin-Cre mice were a kind gift from W. Zhong. Body weight and signs of anemia were evaluated for genetic mosaic MADM-18GT/TG;Apc

METHOD DETAILS

Molecular biology

Generation of MADM targeting constructs

Molecular cloning and generation of recombinant DNA to construct all plasmids (incl. targeting vectors, plasmids with southern probes etc.), and all nucleic acid procedures as described below were carried out according to standard cloning protocols (Sambrook et al., 1989).

Genomic DNA isolation from mouse ES cells

Mouse ES cells were lysed in Lysis Buffer (1M Tris-HCl pH = 7.5, 0.5M EDTA, 5M NaCl, 20% Sarcosyl, 20 mg/ml Proteinase K) overnight at 55°C. Next day, DNA was precipitated with isopropanol for 2 hr at room temperature with agitation and then carefully transferred into a fresh tube containing TE-buffer. The tubes were left open for 10 min to allow residual isopropanol to evaporate. DNA was then incubated for 3 hrs at 37°C. Southern blot

DIG-labeled probes were generated via PCR amplification of plasmid templates containing the probe sequence using a mix of nucleotides containing Digoxigenin-11-dUTP (DIG-dUTP). The PCR reaction was next separated by electrophoresis and the corresponding band was cut and gel purified using the Monarch DNA gel extraction Kit-NEB.

Genomic DNA was digested with the corresponding enzymes overnight at 37°C and electrophoresed in 0.8% agarose for 6 hr at low voltage together with Lambda Hind III marker. Next day, the agarose gels were depurinized in 0.25M HCl, denaturated in 0.4 NaOH and transferred overnight into a positively charged nylon membrane. Next day, agarose gels were assessed under UV light to verify complete transfer of DNA to the membrane. The nylon membrane was then neutralized in 0.5M Tris-HCl (pH = 7.5) and crosslinked with UV light. The membrane was incubated in hybridization buffer (5x SSC, 2% blocking reagent, 50% Formamide, 0.1% Sarcosyl, 0.02% SDS) for 4 hr at 42°C in glass tubes in a rotating oven. In the meantime, the DIG-labeled probe was denaturated at 95°C for 10 min and then quickly chilled on ice for 5 min. The DIG-labeled probe in hybridization buffer was added to the membrane and incubated overnight at 42°C in glass tubes in a rotating oven. Next day, stringency washes were performed with Wash Solution I (2xSSC, 0.1% SDS) at room temperature, followed by Wash Solution II (0.2x SSC, 0.1% SDS) at 68°C. Next day, the membrane was blocked in blocking solution (1% blocking reagent, 0.1M Maleic acid, 0.15M NaCl) for 1 hr. Then anti-DIG AP antibody (1:20,000) in Blocking Solution was added to the membrane, incubated for 30 min at room temperature and then washed with Wash buffer (0.1M Maleic acid, 0.15M NaCl, 0.3% Tween) for 15 min. Finally, the membrane was incubated with CDP-Star (1:100) chemiluminescent substrate in CDP-Star detection buffer (0.1M Tris-HCl, 0.1M NaCl, pH = 9.5) for 5 min, wrapped in transparent film and kept in the dark for 1 hr. The pattern of probe hybridization was detected in a Peqlab FUSION SL Advance system for chemiluminescent imaging.

Generation of transgenic MADM mice

Targeting of MADM constructs to mouse ES cells by electroporation

The linearized MADM targeting constructs were introduced into C57BL/6N embryonic stem cells (Parental ES cell line C2, Stock Number: 011989-MU, Citation ID: RRID: MRRC_011989-MU, A. Nagy Basic ES Cell line) by electroporation using a Bio-Rad Gene Pulser Xcell. After selection with 150µg/ml G418, surviving clones were analyzed for correct targeted integration by Southern blot hybridization (see above). Metaphase spread chromosome counting was performed on ES cells of clones with confirmed correct targeting of the MADM cassettes before they were prepared for blastocyst injection.
Production of chimeras
Host blastocysts were produced by superovulation of BALB/cRj females by intraperitoneal (IP) injection with 5.0 IU of equine chorionic gonadotropin (Folligon; Intervet) and, 48 hr later, with 5.0 IU of human chorionic gonadotropin (Chorulon; Intervet) followed by mating with males of the same strain. Morula stages were harvested from isolated oviducts at day 2.5 days post coitum (dpc) and cultured in M16 medium overnight in an incubator at 37°C and 5% CO₂ to produce host blastocysts. About 10-15 ES cells were injected into a single blastocyst. The injected embryos were cultured for 2-3 hr to recover and then transferred into the right uterus horn of 2.5 dpc pseudopregnant RjOrl:Swiss surrogate mothers as described earlier in detail (Rulicke et al., 2004, Rulicke et al., 2006). The offspring were selected based on their chimeric coat color. High-percentage male chimeras (> 80%) were bred with C57BL/6NR) females and the offspring were selected by coat color and genotyped by PCR for the respective GT or TG MADM transgenes.

Genotyping of MADM reporters, Apc-flox and iv mice
For primer sequences see Table S2. Forward and reverse primer 1 is specific for each MADM reporter. In the absence of MADM cassette the forward/reverse primer 1 PCR will result in the WT band as indicated. The reverse primer 2 is generic and located in the MADM cassette. The forward/reverse primer 2 PCR will result in the MADM band as indicated. The combined use of all three (forward, reverse primer 1, and reverse primer 2) in a single PCR reaction will enable the distinction of WT (single band at WT size), heterozygote (two bands, one at WT and one at MADM size), and homozygous MADM (single band at MADM size) stock mice. Note that MADM<sup>GT/C14</sup> and MADM<sup>TC14</sup>C stock mice should be maintained individually. The distinction of MADM-GT versus MADM-TG is possible by using GT-cassette (GT-for and GT-rev) and TG-cassette (TG for and TG rev) specific primers, respectively. Male mice can be identified by using Y chromosome (Ychrom for and Ychrom rev) specific primers. Presence of transgenes encoding Cre recombinase can be confirmed by using Cre primers (Cre for and Cre rev) as indicated.

Genotyping of Apc-flox mice was performed according to the protocol available at JAX. Genotyping of iv mice was performed using Takara PCR Amplification Kit followed by a 16h enzymatic digestion using TaKaRa TaqI at 65°C. Separation of wt (50bp) and mutant (100bp) band was performed on a 6% agarose gel.

Isolation of MADM-labeled tissue
Mice were deeply anesthetized through injection of ketamine/xyazine/acepromazine solution (65 mg, 13 mg and 2 mg/kg body weight, respectively), and confirmed to be unresponsive through pinching the paw. Perfusion was performed with PBS followed by ice-cold 4% PFA. Tissue was further fixed in 4% PFA overnight at 4°C. Brain, thymus, heart, lung, liver, kidney, spleen, eye and spinal cord were surgically removed and cryopreserved in 30% sucrose for 48 hr and then embedded in Tissue-Tek O.C.T. (Sakura). All samples were stored at −20°C or −80°C until further usage. Samples were sectioned in a cryo microtome at a 10 μm (liver) or 45μm (all other samples) thickness. Brain samples were collected in 24 multi-well dishes and then mounted onto Superfrost Glass Slides (Thermo Fisher Scientific), all other samples were directly mounted on glass slides.

For isolation of skin, pancreas, mammary gland intestine and colon, no perfusion was required. Mice were sacrificed by cervical dislocation and back skin was prepared for histology as previously described (Amberg et al., 2013). Briefly, back skin was shaved and surgically removed above the spine and placed on lint-free surface. Abdominal mammary glands, pancreas and small intestines were surgically removed. Small intestines and colons were cut open longitudinally and made into Swiss rolls. All samples were stored at −20°C or −80°C until further usage. Samples were sectioned in a cryo microtome at a 10 μm thickness and directly mounted onto Superfrost Glass Slides (Thermo Fisher Scientific).

Histology and immunostaining of MADM-labeled tissue
For immunofluorescence staining in skin, pancreas, mammary gland and intestine, sections were thawed at room temperature for 15 min and encircled with DAKO hydrophobic pen. Then, they were washed 3x for 5 min with PBS. Antigen retrieval was performed by adding pre-warmed citrate buffer pH = 6.0 to the samples and incubating them at 85°C for 30 min. Samples were washed 3x for 5 min with PBS, then incubated in blocking solution (10% horse serum, 0.5% Triton X-100 in PBS) for 1h at room temperature. Primary antibodies were diluted in staining solution (5% horse serum, 0.5% Triton X-100 in PBS) and added to the samples over night at 4°C. Next day, samples were washed 3x for 5min with PBS and incubated with secondary antibodies (1:1000) and Hoechst (Sigma, 1mg/ml stock, 1:10000) diluted in staining solution for 2hrs at room temperature. After washing 3x for 5min with PBS, samples were mounted with Mowiol and stored at 4°C until they were imaged at a Zeiss LSM800. Primary antibodies: Keratin 8 (Abcam, 1:100), Keratin 14 (BioLegend, 1:500), beta-Catenin (Cell Signaling, 1:100), phospho-H3(Ser10) (Cell Signaling, 1:800). Secondary antibody: donkey anti-rabbit Alexa647 (Molecular Probes). Mounted brain sections were washed 3x for 5 min in PBS, DAPI stained (1:20’000) for 10 min and then embedded in mounting medium containing 1,4-diazabicyclooctane (DABCO; Roth) and Mowiol (Roth).

Flow cytometry
Mice were sacrificed by cervical dislocation and spleens were collected in ice-cold PBS. Spleens were minced through a 70μm cell strainer. The strainers were then flushed with 10ml PBS-FBS (1x PBS, 2% FBS) and cell suspensions were centrifuged for 6min at 1,200rpm. Cell pellets were resuspended in 1ml ACK lysis buffer (GIBCO) and incubated for 30sec. Lysis reaction was stopped by
adding 10ml PBS-FBS. Cells were centrifuged for 6 min at 1,200 rpm. Pellets were resuspended in 1ml PBS-FBS and transferred to 5ml round-bottom FACS tubes via a 70μm cell strainer. Tubes were filled up with PBS-FBS and centrifuged for 6 min at 1,200 rpm. Cells were incubated with Fc block (BD Biosciences) for 5 min and then incubated with 100μl of antibody mastermix for 30min on ice. Antibodies CD3 HorizonV451 (eBioscience) and CD19 APC (eBioscience) were diluted 1:200. Finally, 4ml of PBS-FBS were added and cells were centrifuged for 6 min at 1,200 rpm. Flow cytometric sorting of GFP⁺, tdT⁺ and GFP⁺ tdT⁺ cells was performed on a BD AriaIII. Analysis was performed using FlowJo.

**Analysis of MADM-labeled brains and peripheral tissue**

Representative images were acquired at an inverted LSM800 or LSM880 confocal microscope (Zeiss) using 10X/20X objectives or 40X/63X oil objectives for acquisition of higher magnification images of immunostained tissue. Images where then processed using Zeiss Zen Blue software and Photoshop (Adobe). Images for quantification were acquired at an inverted LSM800 or LSM880 confocal microscope (Zeiss) or SlideScanner VS120 (Olympus) using 10X objective and processed via custom scripts in ImageJ. Tiled images, encompassing the entire region of interest were imported into Photoshop software (Adobe) and the boundaries for the region of interest were traced. MADM-labeled cells were manually counted based on respective marker expression.

Adenomas in MADM-18;HprtCre;Apc mice intestine or colon were classified based on pathological criteria described previously (Behrens et al., 1998; Fodde et al., 2001; Barker et al., 2009) such as nucleic dysplasia (enlarged and elongated nuclei, strong nuclear staining), invasion of adenomatous epithelium into the lamina propria and up into the villus, coverage of adenomatous epithelium by a normal surface mucosa, polypoid lesion morphology with depressed center, mitotic figures within the adenomatous epithelium.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

See Table S3 for complete information regarding quantifications and statistics used in this study. Table S3 includes all graphed values, including SEMs, p values, and exact values of n. Statistical analysis was performed in the software Prism8 (GraphPad). Evaluation of data was performed by the Welch’s unequal variances t test (Figures 4, 5B, 6A, and 6D), Welch’s ANOVA (Figures 5E, 6C, and 6D) or two-way ANOVA (Figure 6C). Data expressed as ratio was log-transformed prior to the statistical test. For Figures 4 and 5B, n was defined as the density of green/red cells per mm³ from one animal resulting from the quantification of 4-20 sections. For Figures 5 and 6, n was defined as the YGR index for one animal resulting from the quantification of 10-24 sections (Figures 5E, 6A, and 6C), or from FAC-sorted cells from one animal (Figure 6D). The YGRI was defined as the ratio of yellow cells divided by the average of green and red cells.