

Brainbow: New Resources and Emerging Biological Applications for Multicolor Genetic Labeling and Analysis

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ABSTRACT Brainbow is a genetic cell-labeling technique where hundreds of different hues can be generated by stochastic and combinatorial expression of a few spectrally distinct fluorescent proteins. Unique color profiles can be used as cellular identification tags for multiple applications such as tracing axons through the nervous system, following individual cells during development, or analyzing cell lineage. In recent years, Brainbow and other combinatorial expression strategies have expanded from the mouse nervous system to other model organisms and a wide variety of tissues. Particularly exciting is the application of Brainbow in lineage tracing, where this technique has been instrumental in parsing out complex cellular relationships during organogenesis. Here we review recent findings, new technical improvements, and exciting potential genetic and genomic applications for harnessing this colorful technique in anatomical, developmental, and genetic studies.

KEYWORDS *in vivo* imaging; lineage tracing; neural circuitry; clonal analysis; fluorescence microscopy

VISION is arguably the most powerful sensory system in humans. Complex quantitative information portrayed in a visual display is made understandable to the brain by a highly precise visual system, which is accustomed to processing multivariate information present throughout an extremely complex visual field from moment to moment. Visualization tools are therefore particularly useful in the study of dynamic biological systems. In the developing embryo or regenerating tissues for example, cells proliferate, differentiate, and disperse into mature positions. In the nervous system, neurons form complex networks, with thousands of connections potentially overlapping within a small volume (Lichtman and Denk 2011). Analyzing the structure of one of these complex systems through time and/or space is challenging, if not impossible, without a powerful approach for distinguishing among many different individual cellular components. Per-

haps the most useful visual modality for tracking gene function and individual cell behavior within these contexts is color.

Following the isolation of green fluorescent protein (GFP) from *Aequorea victoria* in 1962 (Shimomura *et al.* 1962), fluorescent proteins have been utilized in a wide array of biological systems to label tissues, cells, organelles, or individual proteins (pioneered by Chalfie *et al.* 1994). Modifications to GFP have changed its excitation and emission spectra such that new colors could be added to the biological fluorescence palette (e.g., Tsien 1998; Campbell *et al.* 2002; Shaner *et al.* 2005; Ai *et al.* 2007; Goedhart *et al.* 2012), while unique fluorescent proteins have been identified in other organisms (Matz *et al.* 1999; Shaner *et al.* 2004, 2007; Merzlyak *et al.* 2007). These developments have allowed for genetic targeting of multiple fluorescent proteins (FPs) to visualize different cell types or proteins that interact with one another.

A major limitation in labeling studies has been that cells belonging to one cell type (as defined by a common gene expression pattern) are typically labeled by the same color. Since like cells are often in close proximity with one another, it is difficult to resolve morphology or movement of individual cells. In anatomically complex tissue such as the nervous system, tracking cellular movement and neuronal connections is particularly challenging. This problem can be

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solved in part by labeling very sparsely (e.g., Luskin *et al.* 1988; Walsh and Cepko 1988; Lee and Luo 2001; Noctor *et al.* 2001; Zong *et al.* 2005), but the scarcity of labeled cells makes it difficult to study interactions between cells or neurites (Luo 2007; Jefferis and Livet 2012). The lack of unique cellular identifiers is also limiting for lineage tracing in developmental studies, which depends on the ability to assign large pool of cells to a common progenitor. As a potential solution to these difficulties, the Brainbow multicolor labeling approach was designed and implemented to generate a wide array of fluorescent colors that serve as unique identification tags in living cells (Livet *et al.* 2007).

Basic Principle—How to Get Many Colors and What They Mean

The Brainbow strategy capitalizes on the fact that the three primary colors, red, green, and blue, can combine to generate all colors in the visual spectrum. For example, a television screen combines only red (R), green (G), and blue (B) into a multicolor “RGB” display. Brainbow achieves the same effect by combining three or four distinctly colored FPs and expressing them in different ratios within each cell. The resulting color combinations are unique to each Brainbow-expressing cell and can therefore serve as cellular identification tags that can be visualized by the light microscope (Livet *et al.* 2007; Lichtman *et al.* 2008).

Many different Brainbow and Brainbow-like strategies are now used, utilizing recombinase-mediated DNA excision or DNA inversion (Figure 1A). In DNA excision-based Brainbow (e.g., Brainbow 1.0), three separate FPs are arranged sequentially in the transgene along with two pairs of Cre recombinase recognition sites (Lox sites) that flank the first and second FPs (Figure 1B). The two pairs of Lox sites (*loxP* and *lox2272*) can be recognized by Cre only in identical pairs (i.e., *loxP* with *loxP* and *lox2272* with *lox2272*). Before recombination, only the first color in the array is expressed (termed the “default” color). Following Cre recombination, one of the three FPs will be exclusively expressed by that copy of the cassette. This strategy can be expanded to four FPs by utilizing a third pair of Lox sites (Livet *et al.* 2007).

In DNA inversion-based Brainbow (e.g., Brainbow 2.0; Figure 1C), two matching Lox sites are positioned such that they face each other. Cre inverts (or “flips”) the interspaced DNA as opposed to excising it. In this strategy, two FPs are aligned in head-to-head orientation such that Cre-mediated inversion leads to expression of one of those two colors. By combining excision and inversion, it is also possible to utilize four FPs (e.g., Brainbow 2.1; Figure 1D) (Livet *et al.* 2007).

Combinatorial expression of multiple FPs requires multiple copies of the Brainbow cassette (Livet *et al.* 2007; Lichtman *et al.* 2008). Brainbow is designed to express only one randomly selected FP from each copy of the cassette. For example, if each cell contains only one copy of a three-color construct (e.g., Brainbow 1.0), it would result in a three-color

cell population overall (Figure 1E). More complex multicolor expression results when multiple copies of the Brainbow cassette are present in each cell—either via multiple insertions into the genome or through techniques that introduce many copies as extrachromosomal elements (e.g., microinjection, viral transduction, transfection). When more than one copy of the cassette is present in the nucleus, each can act as the generator of a given “pigment” for that cell. Cre acts randomly on each copy, and thus multiple pigments may be present within each cell, and they mix together to create combinatorial hues (Figure 1F). In practice, up to ~100 colors have been distinguished using various models (Livet *et al.* 2007; Loulier *et al.* 2014). The large number of potential colors provides each cell with a specific color barcode and reduces the chance that two cells will randomly become the same color. This is particularly important for cell tracing (where color is used to follow movement or neurites) and lineage analysis (where color is used to distinguish cell populations derived from different progenitors).

In addition to Cre-Lox excision and inversion strategies, other approaches have been developed to create colorful cellular identification tags. One alternative is to use FLP recombinase and *FRT* recognition sites, which are functionally equivalent to Cre recombinase and Lox sites, respectively (e.g., Flybow and Flpbow) (Hadjieconomou *et al.* 2011; Cai *et al.* 2013). Another strategy is to utilize multiple single-FP vectors simultaneously (Figure 1G). Each vector is stochastically expressed to create combinatorial and diverse hues (Boldogkoi *et al.* 2009; Weber *et al.* 2011; Worley *et al.* 2013; Garcia-Marques *et al.* 2014; Garcia-Moreno *et al.* 2014). This approach is generally more suitable for somatic labeling, as generating and maintaining transgenic lines carrying multiple single-FP transgenes is more challenging. For example, TIE-DYE in *Drosophila* requires two balancer chromosomes to maintain four transgenes (see below) (Worley *et al.* 2013).

The use of multiple colors within one cell population allows for a shift in the types of questions that can be asked using standard reporter constructs. Labeling strategies often use a given promoter to drive one-color expression for all members of that particular cell type, which distinguishes that cell type from others (i.e., cell type 1 is a given color and cell type 2 is a different color). While this strategy is ideal for questions that investigate the behavior of cells at the population level, it essentially homogenizes a given cell population, obscuring differences or interactions between like cells. Brainbow labeling, however, is fundamentally different in that it distinguishes *among* like cells (i.e., individual cells of a given cell type are now many different colors). This approach allows one to address a very different type of question regarding the function of *individual* cells within the population (as opposed to population behavior) and is ideal for following individual cells over time and space, as well as for tracing projections in the nervous system. Another important property of Brainbow is that it is a genetic labeling technique, and the result of stochastic DNA recombination is inheritable. Therefore an initial pool of progenitor cells that

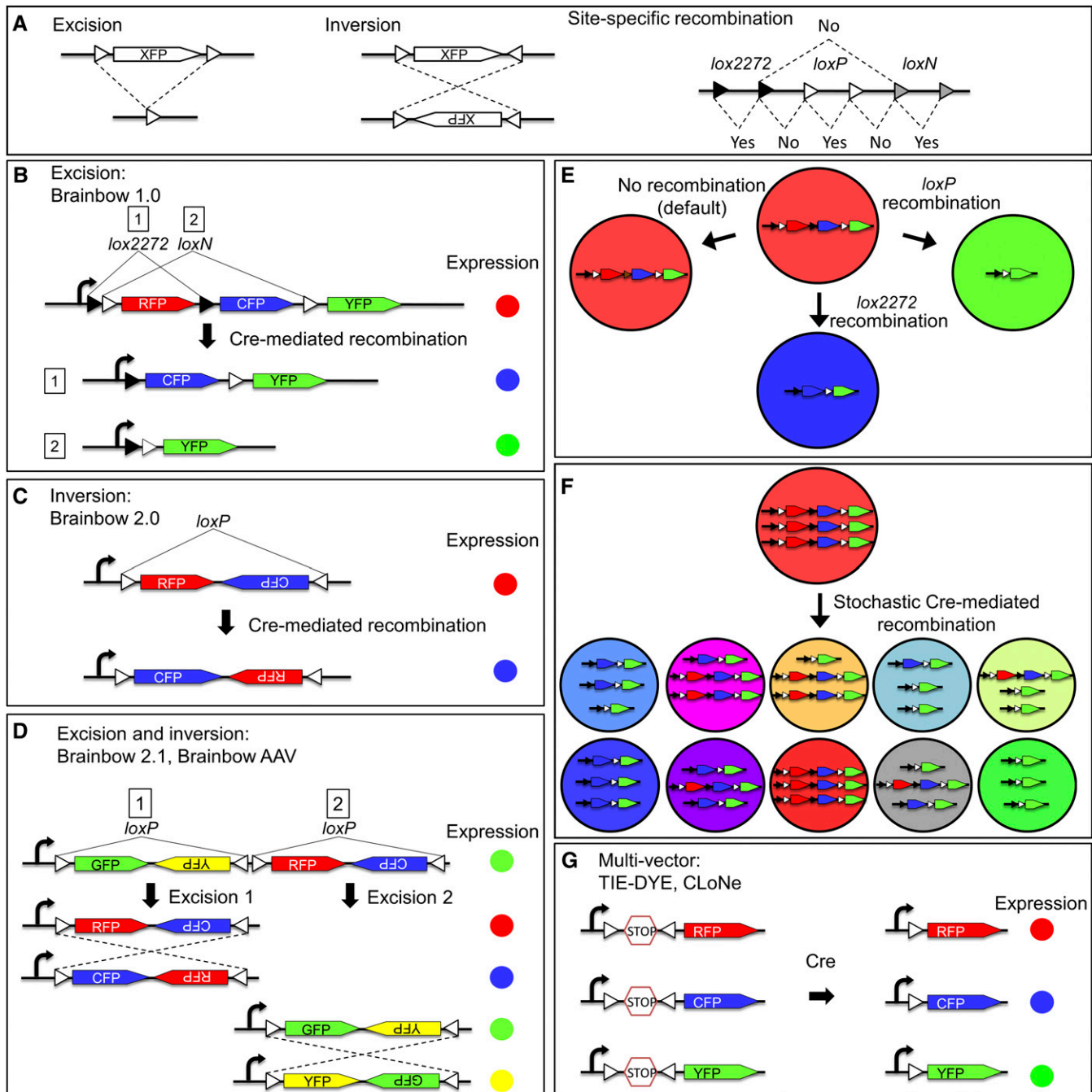


Figure 1 Principles of Brainbow labeling. (A) Cre recombinase can perform excision or inversion of DNA flanked by Lox sites (triangles), depending on the orientation of the Lox sites. Different lox sites such as lox2272 (black triangle), loxP (white triangle), and loxN (gray triangle) function identically but are incompatible with each other. (B) Excision-based Brainbow. Fluorescent proteins (FPs) are flanked by two pairs of mutually incompatible Lox sites. In the absence of recombination, RFP is expressed. Recombination results in excision expression of either CFP (event 1) or YFP (event 2). (C) Inversion-based Brainbow. FP expression can be changed between RFP and CFP by DNA inversion. (D) In Brainbow 2.1, DNA excision leads to selection of either the GFP/YFP pair or the RFP/CFP pair. DNA inversion then decides which FP of the pair is expressed. Brainbow AAV works similarly. (E) For each copy of Brainbow, only the first FP in the array is expressed. Therefore in a cell population with a single Brainbow transgene, cells can be RFP⁺ (no recombination, i.e., “default”), CFP⁺, or YFP⁺. (F) When multiple copies of Brainbow are present in a cell, each copy recombines independently. Three copies of Brainbow can generate 10 distinct colors and more copies will generate even greater color diversity. (G) Combinatorial multicolor labeling can also be achieved by using multiple vectors, each carrying a single FP. As the expression of each FP is stochastic, the color profile within each cell is different. B and F are modified from Pan *et al.* (2013).

is labeled in specific colors produces labeled progeny that reflect their cellular lineage (Figure 2). In other words, all cells within a clone will have the same color (Snippert *et al.*

2010; Hadjiconomou *et al.* 2011; Hampel *et al.* 2011; Rinkevich *et al.* 2011; Blanpain and Simons 2013; Pan *et al.* 2013; Worley *et al.* 2013; Loulier *et al.* 2014). Applications of

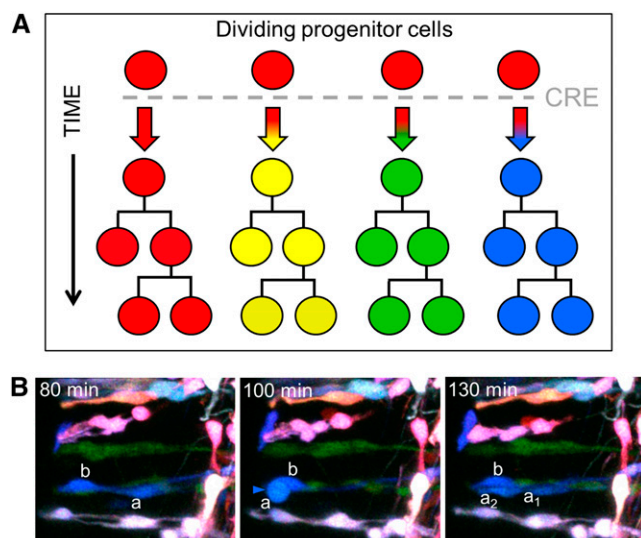


Figure 2 Brainbow for clonal analysis. (A) A uniform population of dividing progenitor cells becomes multicolor upon Cre recombination. Following recombination, each dividing cell produces progeny that share its unique color, thus color coding its resulting clone. (B) This type of Brainbow labeling was used *in vivo* to follow dividing radial progenitor cells in the chick spinal cord over time. Over a period of 50 min shown here, one member of the blue clone (cell a) divides, producing two daughter cells (a1 and a2). Panels in B are reprinted from Loulier *et al.* (2014) with permission from Elsevier.

Brainbow generally make use of colors as cellular identification tags or as markers of parentage (Loulier *et al.* 2014; Roy *et al.* 2014). While Brainbow can help to make sense of a densely labeled tissue, it also is useful in sparsely labeled regions where two or more cells need to be distinguished.

Brainbow Resources: Mouse, Fly, Fish, and Beyond

One of Brainbow's strengths is its broad applicability. A number of Brainbow adaptations have been developed in recent years for different tissues and model organisms such as mouse, rat, chick, zebrafish, fruit fly, and plants. These include both germline transgenic approaches and somatic labeling approaches (e.g., nongermline transgenic). We summarize these two approaches below and in Table 1 and Table 2.

Germline approaches—Brainbow transgenic lines

Mouse: A number of transgenic Brainbow mouse lines have been generated, under the control of neuronal (Thy1.2) or ubiquitous promoters (Table 1). Neuronal lines include the original Brainbow lines (Livet *et al.* 2007), which are now available at The Jackson Laboratory (Bar Harbor, ME). Additional mouse lines were generated recently to expand upon the techniques used in the original neuronal lines (Cai *et al.* 2013) (see *New Improvements to Brainbow*). For ubiquitous expression, the R26-Confetti and R26-Rainbow lines were generated by knocking in Brainbow constructs into the ROSA26 locus (Figure 3B) (Red-Horse *et al.* 2010; Snippert *et al.* 2010; Rinkevich *et al.* 2011). The ROSA26

locus is well suited for constitutive and ubiquitous expression, and both lines have been used extensively for lineage studies (R26-Confetti line is available at The Jackson Laboratory). It is worth noting that for these lines, there is only one copy per haploid genome and it is therefore limited to four (R26-Confetti) or three (R26-Rainbow) distinct colors after recombination. Non-knock-in lines include “Rainbow” (Tabansky *et al.* 2013), “Ubow” (Ghigo *et al.* 2013), “Cytbow”, and “Nucbow” (Loulier *et al.* 2014). Transgenesis with pronuclear injection results in single- as well as multicopy genomic insertions and therefore has greater potential for color diversity (Livet *et al.* 2007).

Zebrafish: In zebrafish, our colleagues and we developed a set of zebrafish Brainbow tools (named “Zebrabow”; Figure 3C) (Pan *et al.* 2013), which include ubiquitous and Gal4-inducible Brainbow transgenic lines, and established parameters for optimal color diversity. We also showed that the combinatorial color profiles remain constant after cellular growth and division, an important prerequisite for color-based lineage tracing. Additional zebrafish Brainbow lines have been generated by other groups for either broad (“PriZm”) (Gupta and Poss 2012) or Gal4-inducible expression (Robles *et al.* 2013). The broadly expressed Brainbow lines have been used to follow cell migration, proliferation, and growth of individual clones in the cornea, heart, and brain (Gupta and Poss 2012; Pan *et al.* 2013; Dirian *et al.* 2014). The Gal4/UAS system has been particularly useful for tracing densely fasciculated axons in the somatosensory and visual systems (Pan *et al.* 2011, 2013; Robles *et al.* 2013). These lines are readily available for public use, and the Zebrabow lines have thus far been distributed to >130 laboratories around the world.

Drosophila: Several groups have adapted Brainbow 1.0 (Hampel *et al.* 2011), Brainbow 1.1 (Forster and Luschnig 2012), and Brainbow 2.0 (“Flybow”) (Hadjiconomou *et al.* 2011) for use with the Gal4-UAS system, enabling expression in specific cell types defined by any Gal4 driver line. Forster and Luschnig (2012), for example, expressed Brainbow in the tracheal tube to reconstruct and quantify the shape and orientation of individual tracheal cells during development, which helped demonstrate a role for the tyrosine kinase Src42A in regulating the expansion of a cylindrical epithelium during development. Boulina *et al.* (2013) have adapted Brainbow specifically for live imaging in *Drosophila*, using a photo-inducible form of Cre to activate recombination *in vivo* (“LOLLIbow”; Figure 3D). Brainbow has been used further in conjunction with the manipulation of gene expression to study gene function. Worley *et al.* (2013) used a multivector, multicolor approach (“TIE-DYE”; Figure 1G and Figure 3E) to follow multiple cell lineages in the wing imaginal disc, simultaneously interfering with expression of UAS-regulated constructs (e.g., *yorkie*, *cubitus interruptus*, or *ras*) in a subset of labeled cells. Unique color expression in each cell clone allowed for clear visualization of the boundary

Table 1 Transgenic lines

Organism	Latin name	Promoter	Transgenic lines
Mouse	<i>Mus musculus</i>	Neuronal	Brainbow 1.0/1.1/2.0/2.1 (Livet <i>et al.</i> 2007), Brainbow 3.0/3.1 ^a /3.2 ^a , Flpbow 1/3 ^a , Autobow ^b (Cai <i>et al.</i> 2013)
		Ubiquitous	R26-Confetti ^b (Snippert <i>et al.</i> 2010) R26-Rainbow (Rinkevich <i>et al.</i> 2011) Rainbow (Tabansky <i>et al.</i> 2013) MAGIC ^c (Loulier <i>et al.</i> 2014) Ubow (Ghigo <i>et al.</i> 2013)
Zebrafish	<i>Danio rerio</i>	Gal4 inducible	Brainbow (Robles <i>et al.</i> 2013) Zebrabow (Pan <i>et al.</i> 2013)
		Ubiquitous	PriZm (Gupta and Poss 2012) Zebrabow (Pan <i>et al.</i> 2013)
Fruit fly	<i>Drosophila melanogaster</i>	Gal4 inducible	dBrainbow ^b (Hampel <i>et al.</i> 2011) Flybow1.0/1.1/2.0 ^a (Hadjieconomou <i>et al.</i> 2011)
		Ubiquitous	LOLLibow (Boulina <i>et al.</i> 2013)
Plant	<i>Arabidopsis thaliana</i>	Ubiquitous	TIE-DYE ^b (Worley <i>et al.</i> 2013)
		Ubiquitous	Brother of Brainbow (Wachsman <i>et al.</i> 2011)

^a Default nonfluorescent nuclear marker expression.

^b No default fluorophore expression in the absence of CRE.

^c Default nuclear-EBFP2 (equivalent to DAPI-labeling) expression.

between mutant and normal cells, revealing differences in cell–cell interactions based on perturbations in gene expression.

Plants: Brainbow has also been applied for genetic studies in *Arabidopsis thaliana*, called “Brother of Brainbow” (BOB) (Figure 3F) (Wachsman *et al.* 2011). One useful feature of this elegant approach is that expression of a gene of interest can be coupled with the default FP (e.g., nuclear YFP), thus extending Brainbow to allow for manipulation and visual determination of gene expression. The authors showed that the retinoblastoma-related (RBR) gene could be coexpressed with BOB, which *trans*-complements the RBR homozygous mutant background. Cre induction ubiquitously or in cell type-specific manners then results in clones that lose the complementing RBR transgene. This is a powerful approach for testing cell autonomous vs. nonautonomous effects and can also be applied in animals (e.g., Loulier *et al.* 2014).

Somatic Brainbow labeling approaches

In addition to transgenic lines, Brainbow can be delivered to somatic cells via DNA injection, electroporation, or viral transduction (Table 2). These nongerm-line approaches can be applied to a wide range of models, allowing for direct cross-species comparisons and applications in organisms for which it is difficult to generate transgenic lines. Furthermore, these methods do not require the time and costs required for generating and maintaining Brainbow transgenic lines.

DNA injection and electroporation: DNA injection and electroporation are widely used for somatic gene expression and can be applied to most model organisms. DNA (plasmid or BAC) is first injected adjacent to the cells of interest, and then an electrical current is applied to transfer DNA into the

Table 2 Somatic expression

Transgenesis method	Transgene name	Organism applied	Genome integration	Applications
DNA injection in embryo	Brainbow (Pan <i>et al.</i> 2011)	Zebrafish	No	Cell and axon labeling
	Brainbow (Egawa <i>et al.</i> 2013)	Mouse, chick	Yes (except for Egawa <i>et al.</i> 2013)	Cell and axon labeling, lineage analysis
Electroporation	CLoNe ^a (García-Moreno <i>et al.</i> 2014)			
	MAGIC ^b (Loulier <i>et al.</i> 2014)			
	Star Track (García-Marques <i>et al.</i> 2014)			
Lentivirus	RGB LeGO (Weber <i>et al.</i> 2011) LeGO with DNA barcode (Cornils <i>et al.</i> 2014)	Mouse, culture cells	Yes	Lineage analysis
AAV	Brainbow AAV ^a (Cai <i>et al.</i> 2013)	Mouse	No	Cell and axon labeling
Pseudorabies virus	Rainbow PRV (Boldogkoi <i>et al.</i> 2009)	Mouse, rat	No	Axon tracing, brain mapping
	PRV-263 (Card <i>et al.</i> 2011a,b)			

^a No default fluorophore expression in the absence of CRE.

^b Default nuclear-EBFP2 (equivalent to DAPI-labeling) expression.

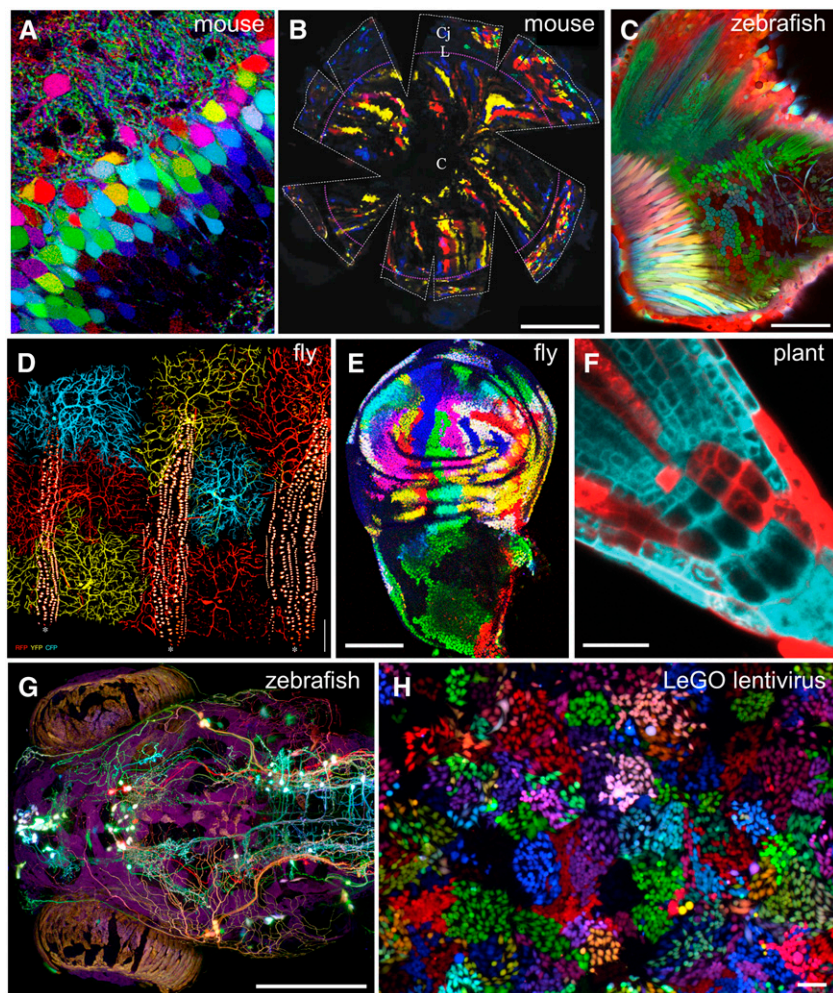


Figure 3 Brainbow transgenic lines and other approaches. (A) Neurons within the dentate gyrus of the Brainbow mouse hippocampus (line L; Image by T. Weissman and J. Lichtman). (B) Radial clones of cells in the mouse cornea from Di Girolamo *et al.* (2014), included with permission from Wiley, Copyright © 2014 AlphaMed Press. (C) Pectoral fin in “zebrabow” zebrafish, from Pan *et al.* (2013). (D) Sensory neurons in the ventrolateral body wall of a *Drosophila* LOLLI-bow larva, adapted from Boulina *et al.* (2013) with permission from Elsevier. (E) Wing-imaginal disc in TIE-DYE *Drosophila*, adapted from Worley *et al.* (2013) with permission from Elsevier. (F) Cells in *Arabidopsis thaliana* root meristem labeled using the Brother of Brainbow system from Wachsman *et al.* (2011). Image is copyrighted by the American Society of Plant Biologists and is reprinted with permission. (G) Dorsal view of larval zebrafish injected with Brainbow plasmid DNA. (H) Human embryonic kidney (HEK) cells transduced by LeGO lentivirus. Image is adapted by permission from Macmillan Publishers (Weber *et al.* 2012). Bars: B, 1 mm; C, E, G, and H, 100 μ m; D, 200 μ m; and F, 20 μ m.

cell. DNA can also be injected directly into the cytoplasm of early embryos, as is done in zebrafish (e.g., Pan *et al.* 2011) (Figure 3G). Brainbow expression with this approach is often very robust and can lead to many color combinations (due to high copy number of the transgene), but diminishes over time due to dilution of nonintegrated transgene through cell divisions. Several recent articles (Garcia-Marques *et al.* 2014; Garcia-Moreno *et al.* 2014; Loulier *et al.* 2014) have overcome this limitation by utilizing genome-integrating transposases such as PiggyBAC and Tol2, allowing long-term cell labeling and lineage tracing. Similarly, transposases also facilitate integration of DNA injected into oocytes (Kawakami 2004; Kikuta and Kawakami 2009).

Viral vectors: Viral vectors such as lentivirus, adeno-associated virus (AAV), and pseudorabies virus can carry different combinations of FPs to generate diverse color profiles in infected cells (Table 2). These vectors can all infect both dividing and quiescent cells, but have very distinct properties and applications (see below).

Lentivirus: Lentiviral vectors [e.g., lentiviral gene ontology (LeGO) vectors] are HIV-based, replication-incompetent

vectors that have been modified for gene delivery without expression of viral components or alteration of cellular metabolism (Wiznerowicz and Trono 2005). The vectors can be integrated into the host genome for stable expression and are inherited after cell division, making them suitable for clonal studies of tissue regeneration and tumorigenesis. To enable multicolor clonal labeling, Weber *et al.* (2011) made use of three different-colored LeGO vectors that express FPs in the three primary colors, red, green, and blue (Figure 3H). In this RGB LeGO system, color diversity is generated by stochastic viral insertion and FP expression in each cell. This approach has been applied *in vitro* and *in vivo* for tracking transplanted liver, bone, and blood stem cells and tumorigenic cells. Direct injection of RGB LeGO may be a potentially powerful method for *in vivo* cell labeling for long-term developmental or morphological analysis, although multicolor labeling would be restricted to the injection site, where the viral transduction rate is high (Weber *et al.* 2012; Gomez-Nicola *et al.* 2014).

AAV: AAV is suitable for long-term transgene expression and can be applied to a wide variety of species and tissues. Unlike lentivirus, AAV persists as an extrachromosomal element and does not integrate into the genome, minimizing the threat of

mutagenesis. The Brainbow AAV system consists of two types of AAV vectors that, in conjunction, can express four different FPs (Cai *et al.* 2013). Infection of multiple AAV virions in one cell is very common and results in high diversity of color. In addition, expression of FPs is Cre dependent, allowing cell type-specific Brainbow labeling. This makes it ideal for high-resolution, single-cell anatomical analyses such as defining the connections of genetically defined cell types in the brain (Cai *et al.* 2013). AAV vectors are not suitable for lineage labeling, however, because AAV episomal DNA is not replicated during mitosis and will therefore be lost after cell division (McCarty *et al.* 2004).

Pseudorabies virus: Pseudorabies virus preferentially infects neurons and can spread across synaptic junctions to label both downstream and upstream neurons. It is therefore a powerful tool to trace the functional neuronal connectivity in the developing and mature nervous system. Multicolor pseudorabies virus has been developed to further facilitate visualization of neuronal morphology and map connectivity of intersecting brain pathways (Boldogkoi *et al.* 2009; Kobiler *et al.* 2010; Card *et al.* 2011a, b). Pseudorabies virus is best suited for anatomical studies and short-term neurophysiological studies. It is not suitable for long-term cell labeling, as chronic infection leads to changes to cellular physiology and eventually death.

In summary, both germline and somatic approaches are suitable for short-term cell labeling experiments and long-term lineage analyses, but they have different strengths and weaknesses. Germline approaches have the advantage of consistent transgene copy number and more homogenous expression. It is easier to produce consistent labeling density and color diversity across different animals. In contrast, labeling density and color diversity are often more variable with somatic labeling, which allows for flexibility in terms of titrating each parameter. Brainbow transgene copy number is usually higher at the injection site and decays with distance, resulting in variable color diversity within an injected individual. The primary strength of the somatic approach is speed and flexibility. Brainbow labeling can be directly applied to strains of interest, even in animals where germline transgenics are less common (*e.g.*, rat and chick).

New Improvements to Brainbow

Since its invention in 2007, some limitations of Brainbow have been recognized (Weissman *et al.* 2011; Cai *et al.* 2013; Roy *et al.* 2014), and several groups have modified the original approach to adapt to different species and improve performance. Here we summarize some of the most notable improvements to Brainbow that make multicolor labeling more robust and more easily applicable.

Improving color balance

The first wave of Brainbow constructs work by switching expression from one (default) FP to another (alternative) FP

(Figure 1E). This ensures that all cells express at least one FP (default or alternative) for easy screening of transgenic animals. The caveat of this strategy is that color balance is dependent on recombinase activity (Figure 4). Another limitation is the perdurance of the default FP after recombination; when recombination occurs after the onset of Brainbow expression, there is accumulation of the default FP that needs to be degraded for the cell to display its appropriate genome-specified hue. This can be a potential issue for lineage tracing, as color may change over time within the same lineage (Pan *et al.* 2013; Loulier *et al.* 2014).

Such limitations can be circumvented by modulating the timing and strength of recombinase activity, but Brainbow without default expression is desirable for lineage tracing or when analyses are done soon after the onset of recombination. Several groups have now generated transgenic lines and somatic labeling tools that drive multicolor labeling in recombined cells while eliminating the default FP expression (with a transcriptional stop signal) or utilize a nuclear localized FP that can be clearly distinguished from the FP expressed after recombination (Figure 5A) (Snippert *et al.* 2010; Hampel *et al.* 2011; Cai *et al.* 2013; Loulier *et al.* 2014). In these configurations, all alternative FPs have equal chances of being expressed. Constructs with or without default expression are noted in Table 1 and Table 2.

Antibody amplification

Many commonly used FPs are derived from jellyfish (*A. victoria*) (*e.g.*, GFP, YFP, BFP, and CFP) (Tsien 1998), coral (*Discosoma* sp.) (*e.g.*, dsRed, dTomato, mOrange, and mCherry) (Shaner *et al.* 2004), or anemone (*Entacmaea quadricolor*) (*e.g.*, TagRFP, TagBFP, and mKate2) (Merzlyak *et al.* 2007). FPs derived from the same species can have distinct endogenous fluorescence spectra, but are too similar antigenically to be distinguished by antibodies. This poses a challenge for histological analyses, where endogenous fluorescence often becomes too weak after fixation. To overcome this limitation, Hampel *et al.* (2011) (dBrainbow) added unique epitope tags to each FP, so that fluorescence from each can be independently amplified via antibody labeling. An alternative approach was developed by Cai *et al.* (2013), which utilizes FPs derived from different species (PhiYFP from *Phialidium* sp., mOrange from *Discosoma* sp., GFP from *A. victoria*, and mKate2 from *E. quadricolor*) so that each FP can be recognized by antibodies specific to each FP. Both of these approaches allow for the boosting of fluorescence intensity for analysis in fixed tissue.

Improving color discrimination

From 3 to ~100 colors can be generated by Brainbow (Livet *et al.* 2007). Dividing a finite color space into increasing numbers of colors, however, requires the investigator to distinguish between closely related hues (*e.g.*, between different shades of yellow). It is therefore of great concern whether perceived differences in color (by eye or digital quantification) represent true differences in cellular identity/lineage or simply experimental variability.

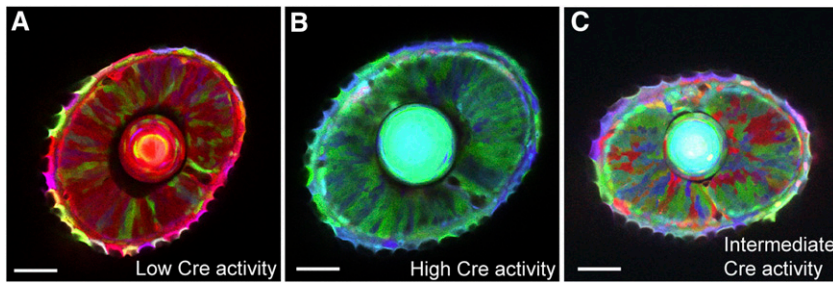


Figure 4 Tuning Cre activity to maximize color diversity. Images show larval zebrafish eyes expressing Brainbow 1.0 (Zebrafish). (A) When Cre activity is low, most copies of Brainbow express the default FP (RFP). (B) When Cre activity is high, all of the Brainbow copies are recombined, resulting in only the nondefault FPs (CFP and YFP). (C) An intermediate level of Cre activity results in much greater color diversity. Bar, 50 μm . Modified from Pan *et al.* (2013).

An elegant solution is to increase the dimension of labeling by targeting different FPs to different subcellular compartments (Garcia-Moreno *et al.* 2014; Loulier *et al.* 2014). Consider a cell with two copies of cytoplasmic Brainbow (Cytbow) and two copies of nuclear Brainbow (Nucbow). Each copy of Cytbow and Nucbow recombines independently, resulting in 7 possible cytoplasmic and 7 possible nuclear colors (including unlabeled for each) and thus $7 \times 7 = 49$ different possible color combinations overall (Figure 5B). Moreover, these 49 possibilities would be relatively straightforward to visualize and quantify, because it is technically easier to cluster 7 distinct hues for two different subcellular compartments than to cluster 49 distinct hues in one compartment. Loulier *et al.* (2014) showed that this approach can be used to distinguish different topological adjacent and intermixed clones. Similar logic potentially can be applied to trace cellular projections or specific structures. For example, axons are robustly labeled when FPs are targeted to the plasma membrane; such labeling could be combined with cytoplasmic- and mitochondria-targeted Brainbow to increase dynamic range and distinguish among axons within a more complex population of projections (*e.g.*, Livet *et al.* 2007; Cai *et al.* 2013; Loulier *et al.* 2014).

Current and Emerging Brainbow Applications

Brainbow has had significant impact on a number of diverse disciplines, including neurobiology, developmental biology, cancer, and stem cell biology. Recent developments in imaging, computation, and genomics can potentially synergize with Brainbow to allow more multifaceted and comprehensive analyses of biological systems. Here we highlight several current applications and discuss emerging avenues for applying Brainbow.

Mapping neuronal connectivity with Brainbow

Mapping the connectivity patterns between diverse neuron types within the brain is one of the major challenges in neuroscience. Brainbow's color diversity provides a unique way to unambiguously trace axons and identify neuronal connections over long distances. Livet *et al.* (2007) utilized Brainbow to decipher the connectivity between mossy fiber axons (originating in the brainstem and cerebral cortex) and granule neurons within the cerebellum. In total, 341 axons and 93 granule neurons in a small three-dimensional volume ($160 \mu\text{m}^2 \times 65 \mu\text{m}$) were digitally reconstructed to

demonstrate the convergence of multiple presynaptic neurons onto individual granule cells (Figure 6, A and B). More recently, Kang and Lichtman (2013) used Brainbow expression to distinguish among multiple axons reinnervating the neuromuscular junction following peripheral nerve injury, showing that regenerating axons avoid other axonal branches only if they arise from the same parent neuron. Multicolor tracing with Brainbow has also been applied in *Drosophila* (Hampel *et al.* 2011; Hadjiconomou *et al.* 2011; Boulina *et al.* 2013), zebrafish (Pan *et al.* 2011; Heap *et al.* 2013; Robles *et al.* 2013), and chick (Egawa *et al.* 2013). Notably, Egawa *et al.* (2013) used Brainbow to detect the refinement of multiple inputs into the ciliary ganglion during embryonic chick development, identifying the precise time point suitable for optogenetic manipulation of neural activity.

The present challenge is to expand brain mapping analyses from small three-dimensional volumes to the whole brain (Lichtman and Denk 2011), and recent technical advancements may pave the way. Notably, computational methods use machine learning and visualization tools for complex brain-wiring data sets (Kim *et al.* 2014; Oh *et al.* 2014), tissue-clearing techniques make the brain optically transparent (Dodt *et al.* 2007; Hama *et al.* 2011; Chung *et al.* 2013), and imaging techniques correct for light distortion in thick samples (C. Wang *et al.* 2014; K. Wang *et al.* 2014). Coupled with multicolor Brainbow labeling, these methods could allow whole brain neuronal imaging while retaining the ability to identify neurites from many individual cells. This would complement current brain mapping approaches, which either are restricted to small monochromatic volumes (*e.g.*, serial electron microscopy) (Bock *et al.* 2011; Helmstaedter *et al.* 2013; Takemura *et al.* 2013) or do not have single-cell resolution (*e.g.*, anterograde and retrograde viral tracers) (Osten and Margrie 2013). Furthermore, Brainbow-powered light microscopy can be done within intact and living tissues, allowing one to track cellular dynamics and function in real time (*e.g.*, multiple time-point assays and genomic analysis of imaged cells). It will be exciting to see what the field can achieve by combining Brainbow with the rapidly advancing suite of new imaging techniques.

Cellular dynamics and lineage tracing

While Brainbow was originally developed for use in brain mapping and connectivity studies, its application to the field of development has been particularly impactful. During development, orderly proliferation and cellular migration

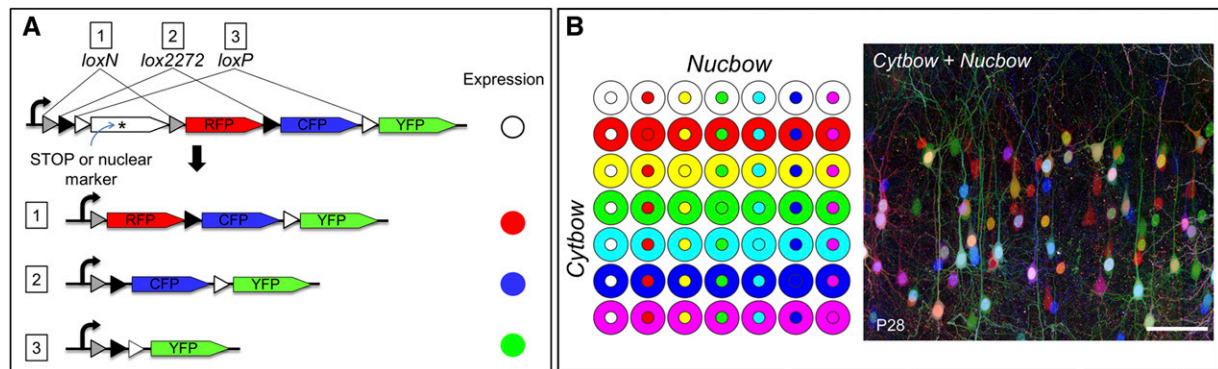


Figure 5 Improvements in color balance and discrimination. (A) A Brainbow construct with no default FP expression. After recombination, there is an equal probability of expressing RFP, CFP, or YFP, and thus color balance is not dependent on Cre activity. (B) Combining Brainbow constructs with different subcellular localization improves color discrimination. Shown in the left diagram, each cell can express both a nuclear color from Nucbow and a cytoplasmic color from Cytbow. When there are two copies of each, 49 total color combinations can be generated. Fluorescence image on the right shows a mouse cortex labeled by the Cytbow and Nucbow combination. Bar, 100 μ m. B was modified from Loulier *et al.* (2014) with permission from Elsevier.

determine tissue size and the correct assembly of component cells. Conventional viral or single FP lineage labeling methods are often unable to determine whether cells are derived from a single clone, especially for cells that undergo extensive migration. Viral vectors with genetic barcodes provide additional proof of lineage, but cannot be utilized when multiple clones are intermingled, restricting labeling to a very low density (one to two clones per animal) (Luskin *et al.* 1988; Walsh and Cepko 1988).

The diversity of Brainbow color provides an ideal method to unambiguously label multiple clones in close proximity (Figure 2). Multicolor labeling can be targeted to specific cell populations and developmental stages by utilizing a cell-type-specific promoter driving the Brainbow construct, Cre, or driving CreER, a chemically inducible form of Cre. In a series of elegant experiments, Clevers and colleagues combined intestinal stem cell-specific expression of CreER (Lgr5-CreER) and Brainbow (R26-Confetti) to investigate the dynamics of stem cell proliferation and homeostasis within the intestinal crypt (Snippert *et al.* 2010, 2014; Schepers *et al.* 2012; Ritsma *et al.* 2014). In combination with quantitative analysis, these studies suggest that stem cell homeostasis is regulated by neutral competition between dividing stem cells for a spatially limited proliferative niche and that adenoma cells are derived from Lgr5⁺ intestinal stem cells. Some other notable applications include the analysis of lineage in astrocytes and neurons (Dirian *et al.* 2014; García-Marques *et al.* 2014; García-Moreno *et al.* 2014; Loulier *et al.* 2014), coronary arteries (Red-Horse *et al.* 2010), corneal epithelial cells (Pan *et al.* 2013; Amitai-Lange *et al.* 2014; Di Girolamo *et al.* 2014), germline progenitor cells (Zhang *et al.* 2012; Komai *et al.* 2014), cleavage stage blastomeres (Tabansky *et al.* 2013), Langerhans cells (Ghigo *et al.* 2013), papillae of the tongue (Tanaka *et al.* 2013), radial glial cells (Pilz *et al.* 2013), developing nephrons (Barker *et al.* 2012), hematopoietic cells (Wang *et al.* 2013), distal digit (Rinkevich *et al.* 2011), and cardiomyocytes (Gupta and Poss 2012). Rapid developments in high-speed light microscopy

such as selective plane illumination microscopy (also called light sheet) (Keller *et al.* 2008; Amat *et al.* 2014), as well as multicolor volume microscopy (Mahou *et al.* 2012, 2014) will further improve the resolution and duration of lineage tracing experiments.

Moving beyond just color—genomic and genetic analyses

Current applications for utilizing Brainbow in anatomical, developmental, or lineage studies to date have focused mostly at the cellular level. The genomic profiles and cellular identities that determine specific neuronal connectivity or clonal behavior largely remain mysteries. We believe future studies will aim to combine *in vivo* observations with genetic analysis. The study of blood cells has led the way in this respect, where distinctly colored clones can be identified *in vivo*, isolated by flow cytometry, and then analyzed by RT-PCR and sequencing (Figure 6C). Different approaches would be suited for different tissue types. Less adherent cells, for example, would be more suitable for flow cytometry, whereas large adherent clones would be more suitable for laser capture microdissection. An attractive possibility is to combine multicolor lineage tracing with single-cell sequencing and utilize the relative amounts of different FPs to determine the cellular origin.

Another approach for using Brainbow to manipulate gene activity is to link color to the expression of specific transgenes (Figure 6D) (Wachsman *et al.* 2011; Worley *et al.* 2013; Loulier *et al.* 2014). If a gene is fused with (or expressed in tandem with) one particular FP, then cells with detectable levels of the FP can then be followed to measure the effect of gene expression and to test for cell autonomous *vs.* non-cell autonomous effects. For example, Loulier *et al.* (2014) designed a Brainbow construct in which one of the FPs (CFP) is coexpressed with a gene that regulates mitotic spindle orientation (dominant-negative LGN). Four days after electroporation of the dominant-negative LGN–Brainbow construct into mouse embryonic cortex, it was found that cells expressing CFP (and thus dnLGN) were less likely

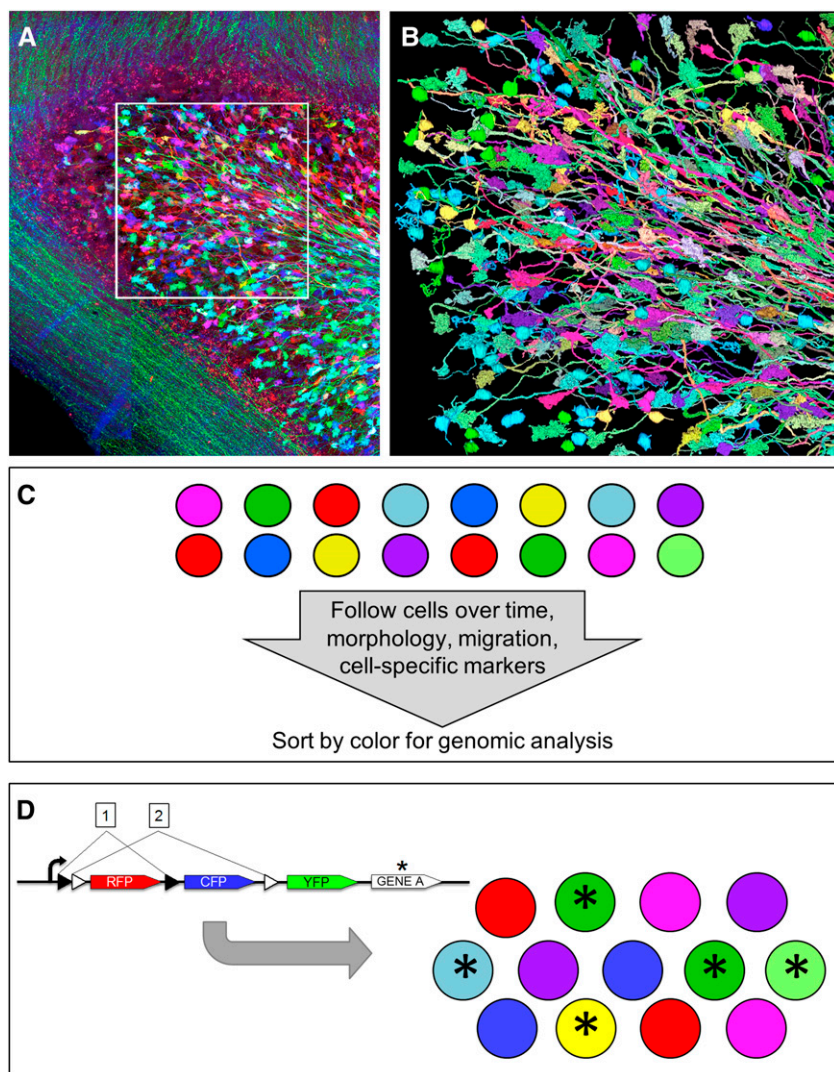


Figure 6 Current and emerging applications. (A) A cerebellar folium from the Brainbow mouse line H was imaged using confocal microscopy. Three-dimensional volume ($160 \mu\text{m}^2 \times 65 \mu\text{m}$) indicated in the box was segmented using semiautomated methods and reconstructed digitally, as shown in B. (B) Digital reconstruction of 341 axons and 93 granule neurons from volume marked in A. A and B are modified from Livet *et al.* (2007). (C) Multicolor cells can be followed over time in living tissue and then sorted by color (e.g., FACS) for sequencing or gene expression analysis. (D) In this schematized construct, a particular gene (gene A*) is coexpressed with YFP (following excision at loxP site 2). In the resulting cell population at right, only cells expressing any level of YFP will also express the gene of interest.

than CFP-negative cells to be located in the ventricular zone and more likely to have moved to the cortical plate, consistent with predicted roles for LGN in development (reviewed in Pevre and Morin 2012). Expanding upon this approach, FP intensity could even be quantified to assay dosing and combinatorial effects of gene expression. Theoretically, each FP could be fused with a separate gene (e.g., gene 1 with YFP, gene 2 with RFP, and gene 3 with CFP). In this case, the hue of each cell would represent a specific ratio of gene expression. Such techniques will expand Brainbow's potential in genetic studies.

Practical Tips for Brainbow Optimization and Analysis

There are several useful step-by-step guides for Brainbow labeling and we encourage readers to consult them for specific hardware requirements and protocols (Pan *et al.* 2011; Weissman *et al.* 2011; Mahou *et al.* 2012; Shimosako *et al.* 2014). Here we highlight several challenges and key issues that we and other users have encountered.

Maximizing color diversity

Two main factors are important for determining color diversity: (1) the copy number of the Brainbow transgene and (2) the timing and level of Cre activity. In general, as more copies of the Brainbow DNA construct are present in cells, higher expression levels and more color combinations result (mixture of more pigments); however, too many copies per cell can result in reduced color diversity (since all cells have all pigments). For example, Cai *et al.* (2013) observed reduced color diversity at the Brainbow AAV injection site (likely due to too many copies present in each cell) with maximal color diversity farther away from the injection site (fewer copies per cell). In transgenic lines that have incorporated multiple copies in a tandem array, there is some evidence that recombination between matching pairs of Lox sites extends across insertions, leading to a decrease in copy number and reduced color diversity following Cre activity (Loulrier *et al.* 2014; J. Livet, personal communication). The ideal number of colors is not the same for every

experiment. Some studies that investigate regions with many overlapping cells may require a large number of distinct colors, while a handful of colors may be appropriate in studies that consider less dense regions.

The timing and amount of recombination also determine clone size and color balance and need to be titrated appropriately for each system. For Brainbow vectors with default expression (*i.e.*, expression of a FP in the absence of Cre, such as Brainbow 1.0), tunable Cre expression is necessary to adjust color balance (Figure 4). For Brainbow constructs with no default expression (*e.g.*, Brainbow 3.0), color balance is not dependent on the level of Cre activity. Furthermore, the timing of Cre activity determines when progenitor cells are labeled: if recombination occurs very early in a lineage when only a few progenitor cells are present, the entire resulting cell population may inherit only those few same colors. Delaying recombination allows for a larger progenitor pool to recombine separately, generating more colors for the resulting cell population.

Color constancy

Since recombination is random, color will vary from one animal to the next. For a given promoter, however, the same cell populations should be labeled. If individual cells are being followed over time or space, it is crucial to keep constant the image acquisition settings, since small changes from one imaging session to the next can change color appearance significantly. If tracking development, the growth of the organism can send the cells of interest deeper into the tissue, affecting the relative scatter of each FP's emitted wavelength. In this case, the use of nearby contextual landmarks to ensure cellular identity is useful. If relying only upon color to assign neuronal identity (for example, concluding that a light pink cell body in one part of brain corresponds to a light pink axon in a different region, without tracing it there), a rigorous controlled approach must be used for acquiring images and quantifying color (Livet *et al.* 2007; Weissman *et al.* 2011; Cai *et al.* 2013).

In general, any factor that affects the FPs differentially can lead to an undesired change of a cell's overall hue. Short wavelengths (*e.g.*, blue light) scatter more readily than longer wavelengths (*e.g.*, red), and thus the same cell may appear different when it is located superficially as opposed to deep (*i.e.*, it may appear more blue toward the surface). Bleaching can also affect each FP to different extents, since each has its own photostability. In general, *in vivo* preparations are less vulnerable to bleaching of fluorescence expression.

Color quantification

In most cases it is necessary to quantify the diverse colors observed by eye, particularly when conclusions are drawn based on similarity or dissimilarity of color, *e.g.*, assigning lineage relationships, or as readout of gene expression. In image processing software such as Image J and Photoshop, colors are displayed as three channels: red, green, and blue. This is known as the RGB color model. With three-FP Brainbow,

RFP is usually designated as red, YFP as green, and CFP as blue. When values from each channel are directly plotted on a three-dimensional graph, there is substantial variability along a diagonal intersecting zero (Figure 7A). This reflects variability in brightness, which depends on cellular topography, imaging depth, and promoter activity. It is therefore preferable to normalize brightness level and measure the relative proportions of each FP. This can be done with a ternary graph that has three axes, each representing the percentage of a color (red, green, or blue; Figure 7B) (Loulier *et al.* 2014). Another approach is to convert RGB values to hue, saturation, and brightness (also known as the HSB or HSV color model). Color values can be plotted as a two-dimensional hue vs. saturation graph, which is independent of brightness (Figure 7, C and D). Ratiometric color quantification has been extended to up to five FPs (Malide *et al.* 2012).

For lineage analysis, cells with the same color are likely derived from the same progenitor. It is important to keep in mind, however, that it is still possible for two cells with a distinct lineage to arrive at the same hue by chance (Figure 7E). Wider color diversity greatly reduces the likelihood that unrelated cells will have the same color, but additional verification is necessary when color combinations are few (*e.g.*, single genomic insertion of Brainbow or suboptimal Cre activity) or when cells from different clones are intermingled (see Blanpain and Simons 2013). One powerful way to test whether single-color cells belong to the same clone is to compare the average number of cells per single-color clone (clone size) at different labeling densities (Figure 7E): if cells with the same color are clonally related, the average clone size should be the same regardless of labeling density, similar to what has been done in retroviral clonal analysis (Galileo *et al.* 1990).

Conclusions

Brainbow is a tremendously powerful tool for visualizing the dynamics of large numbers of cells, unraveling neural circuits, and piecing together lineage relationships. Emerging applications can now combine visual observations with genetic and genomic analyses. Particularly intriguing are approaches that use color to first visualize and identify a given cellular population and then quantify gene expression in those cells, in addition to approaches that manipulate gene function by linking expression of FPs with given transgenes, thus genetically targeting a subset of clearly identifiable cells.

The Brainbow toolbox has greatly expanded in recent years. A variety of transgenic Brainbow lines are available for use in common model systems such as mouse, *Drosophila*, and zebrafish. Somatic labeling approaches such as microinjection and viral vectors further extend Brainbow to model systems that are less amenable to transgenesis. New improvements to the original Brainbow constructs have increased its practical use significantly, especially in terms of color detection and discrimination.

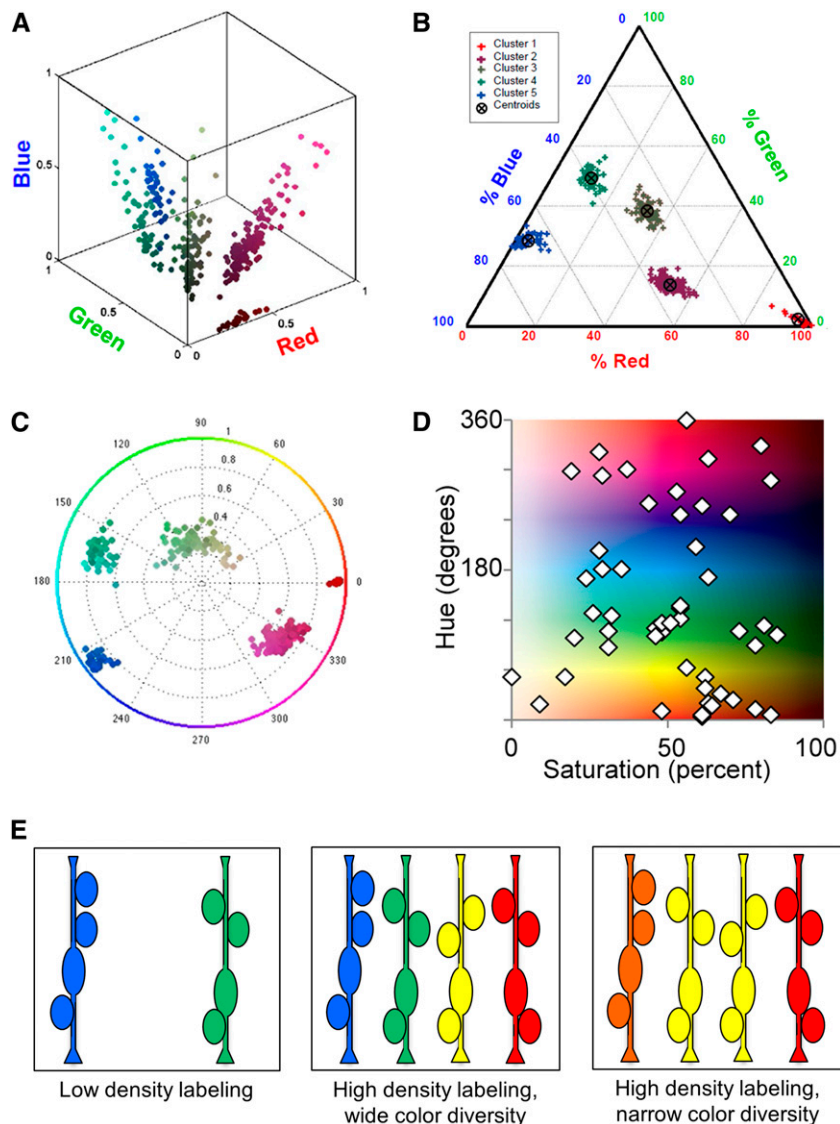


Figure 7 Quantification of color. A–C compare the same data set with different quantification methods. (A) When color is plotted in a three-dimensional RGB graph without correction for brightness, fluorescence values of different cells are widely scattered along the diagonal. (B) When colors are expressed as a ratio along three color axes, color distribution becomes more clustered. (C and D) Color can also be normalized by plotting hue (0°–360°) against saturation (0–100%). This can be displayed either as a circular plot (C) or as an XY distribution (D). A–C are modified from Loulier *et al.* (2014) with permission from Elsevier. D was modified from Pan *et al.* (2013). (E) Schematized clones of related cells are shown, labeled at low and high densities. If the average number of cells per clone is four in low-density labeling (left), this average number should also be four in high-density labeling (center). Narrow color diversity (right) could result in labeling of neighboring clones the same hue (two yellow clones in the center). In this case the average clone size would appear larger.

While Brainbow initially gained attention for its beauty, it has now proved to be a wellspring of biological insights, for example into neuronal connectivity patterns (Livet *et al.* 2007; Egawa *et al.* 2013; Heap *et al.* 2013; Kang and Lichtman 2013; Robles *et al.* 2013), dynamics of stem cell proliferation and organ homeostasis (Snippert *et al.* 2010; Rinkevich *et al.* 2011; Gupta and Poss 2012; Schepers *et al.* 2012; Tabansky *et al.* 2013), and genetic regulation of single cells *in vivo* (Wachsman *et al.* 2011; Forster and Luschig 2012; Loulier *et al.* 2014). We believe the new resources and applications will make Brainbow an increasingly valuable research tool, and we look forward to seeing exciting (and beautiful) Brainbow genetic studies in the future.

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