A Barcoding Strategy Enabling Higher-Throughput Library Screening by Microscopy

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A Barcoding Strategy Enabling Higher-Throughput Library Screening by Microscopy

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Abstract

Dramatic progress has been made in the design and build phases of the design-build-test cycle for engineering cells. However, the test phase usually limits throughput, as many outputs of interest are not amenable to rapid analytical measurements. For example, phenotypes such as motility, morphology, and subcellular localization can be readily measured by microscopy, but analysis of these phenotypes is notoriously slow. To increase throughput, we developed microscopy-readable barcodes (MiCodes) composed of fluorescent proteins targeted to discernible organelles. In this system, a unique barcode can be genetically linked to each library member, making possible the parallel analysis of phenotypes of interest via microscopy. As a first demonstration, we MiCoded a set of synthetic coiled-coil leucine zipper proteins to allow an 8×8 matrix to be tested for specific interactions in micrographs consisting of mixed populations of cells. A novel microscopy-readable two-hybrid fluorescence localization assay for probing candidate interactions in the cytosol was also developed using a bait protein targeted to the peroxisome and a prey protein tagged with a fluorescent protein. This work introduces a generalizable, scalable platform for making
microscopy amenable to higher-throughput library screening experiments, thereby coupling the power of imaging with the utility of combinatorial search paradigms.

**Graphical abstract**

**Keywords**

synthetic biology; library screening; microscopy barcodes; coiled-coil zippers

**Introduction**

The inherent complexity of biological systems and our limited understanding of forward engineering principles usually requires the testing of a large number of designs in order to find one that yields the desired phenotype, much less learn design principles to gain predictability.\(^1\) Engineering cellular behaviors often demands the ability to measure a phenotype in individual cells. Accordingly, methods such as cytometry and fluorescence-activated cell sorting (FACS) have become powerful single-cell analytical tools that can be applied in high-throughput. A limitation of these methods is the requirement of their outputs of interest to be fluorescent or be able to link them to a fluorescent output. Many desirable cellular phenotypes such as motility, morphology, and subcellular localization are not currently amenable to these high-throughput strategies but can be clearly visualized by microscopy. However, the throughput of microscopy experiments is not sufficient for rapid design-build-test cycles.

In order to screen for these phenomena, it is often necessary to link the observable phenotype to the underlying genotype. To this end, many methods for barcoding cells for flow cytometry or microscopy have been devised, often involving the labeling of cells with unique fluorescent markers.\(^2\)–\(^5\) Two early methods of uniquely tagging cells, Brainbow and red-green-blue (RGB) labeling, use random combinations of fluorescent reporter cassettes to create a unique color for each labeled cell.\(^2\)\(^,\)\(^3\) While such approaches provide a striking visualization for distinguishing clonal cells, their stochastic nature makes it difficult to programmatically link the cell’s genotype to the observed phenotype. Another technology, fluorescent cell barcoding, uniquely labels each cell type by using a distinct dye with further multiplexing accomplished by using different concentrations of each dye to create more unique fluorescent signatures.\(^4\) While this approach allows for the pooled screening of mixed populations, it requires that each cell type be fixed then individually labeled, thereby limiting the method’s utility for live cell screening and pooled experiments of large combinatorial libraries. Recent work with surface zinc fingers has progressed toward genetically-encoded cell surface scaffolds that hybridize with fluorophore-labeled DNA.
probes. While this technique has been used to identify up to six unique genotypes in a pooled screening assay with live, immobilized cells, expanding to larger library sizes (≥10^6) would require increasing numbers of hybridization/quenching steps that would limit experimental throughput.

To address these limitations in existing methods for high-throughput cell labeling, we have developed a microscopy-readable library of barcodes (MiCodes) that enables live-cell tracking and time-lapse imaging and can be used to increase the throughput of microscopy experiments. Each MiCode consists of a combination of spectrally distinct fluorescent proteins targeted to visually discernible organelles. In this manner, a large set of unique MiCodes can be obtained with the total number of MiCodes defined by the following equation:

\[ m = f(s, c, o) = (s^c)^o \]  

where ‘\( m \)’ is the number of MiCodes, ‘\( s \)’ is the number of channel states - in this case two for OFF and ON, ‘\( c \)’ is the number of fluorescent channels, and ‘\( o \)’ is the number of organelles. A MiCode library can be physically coupled to a DNA-encoded library such that each library member is genetically linked to a unique MiCode, forming the aforementioned necessary link between phenotype and genotype. Here, we demonstrate a proof-of-concept MiCoding screen with four fluorescent proteins and four organelles using a subset of the possible \((2^4)\^4 = 65,536\) MiCodes. Micrographs can be taken to measure, in parallel with the desired phenotype, the MiCodes of all cells in the field of view.

As an example application, we tested candidate interactions among a set of synthetic coiled-coil proteins that were designed to maximize orthogonality. Coiled coils have been deployed in many synthetic biology applications as modular, high affinity protein-protein interaction parts for engineering protein assemblies in a cell.\(^6\)–\(^8\) Increased numbers of specific, orthogonal interaction pairs would enable synthetic biologists to engineer multiple independent complexes in the same cell with minimal cross-talk. Achieving interaction orthogonality appears to be challenging.\(^9\),\(^10\) For example, although orthogonality was not an explicit goal of a previous effort by Reinke and coworkers when they constructed a set of twenty-three synthetic coiled-coils with high affinity for their cognate partners, it is still instructive that a maximum set of only three orthogonal pairs was observed.\(^11\) When characterizing candidate sets of orthogonal pairs, the size of the interaction matrix that must be tested increases as \(n^2\), where \(n\) is the number of protein pairs to be tested. To address this, we MiCoded a matrix of sixty-four candidate coiled-coil interactions and developed a two-hybrid assay using subcellular localization as an output to simultaneously determine if an interaction occurred in the cytosol \textit{in vivo} for all members of this matrix. Thus, the MiCodes developed herein can enable microscopy to be used as an output for a genetically-coupled DNA-encoded library.
Results and Discussion

I. Construction of MiCodes

Identifiable microscopy-readable barcodes require clearly discernible morphological features. As such, we targeted organelles that produced the following phenotypes: a large ring (plasma membrane), a smaller ring (vacuole membrane), a large single punctate spot (nucleus), and peripherally-clustered small punctae (F-actin patches) (Fig. 1A). To achieve these phenotypes, we tested a series of organelle targeting tags (Supp. Table S1) described in the Yeast GFP fusion Localization Database, (yeastgfp.yeastgenome.org/) as fusions to each of the fluorescent proteins used in this study (Supp. Table S2) under a range of expression levels. The targeting tags showing the clearest organelle targeting with minimal cytosolic background fluorescence were the C-terminal peptide tag CIIC for the plasma membrane, zinc transporter ZRC1 for the vacuole membrane, histone HTA2 for the nucleus, and actin binding protein ABP1 for F-actin patches (Supp. Table S1). To scale the diversity of possible barcodes, we next increased the number of fluorescent channels under which each targeting tag could be visualized. The compatible four channels mRuby2 red fluorescent protein (RFP), Venus yellow fluorescent protein (YFP), mTurquoise cyan fluorescent protein (CFP), and mTagBFP2 blue fluorescent protein (BFP) were used (Supp. Table S2). The expression level yielding a distinct fluorescence signal at the desired location with minimal background cytosolic fluorescence was determined. For initial expression level optimization, we used five promoters spanning approximately three orders of magnitude in strength to drive expression of each of the organelle targeting tags fused to GFP. In most cases, we found moderate expression levels (pRPL18B) to be best for yielding a clear positive signal for the given target organelle while minimizing diffuse, cytosolic fluorescence (Supp. Fig. S1, Supp. Table S3). In general, the precise optimal expression level of each tag will also likely be dependent on the imaging protocol used since minimization of cytosolic background is critical for unambiguous MiCoding.

Synthesis of the MiCode was done using a MoClo Golden Gate assembly strategy and the constituent parts of the MiCode were integrated into the URA3 and LEU2 loci of the chromosome (Supp. Fig. S2). In preliminary design iterations, we observed undesired homologous recombination among the MiCode constructs due to repeated use of the same promoters, fluorescent proteins, and terminators. Accordingly, alternate promoters with similar transcriptional strengths to pRP18B, but with sequence-independent nucleotide composition, were used to drive the various targeting tag/fluorescent protein fusions (Supp. Table S3). Similarly, different terminators were used for each fusion protein (Supp. Table S4). Finally, the organelle targeting tags (Supp. Table S1) and the fluorescent protein reporter genes were recoded to make an alternate primary sequence for Venus, mTurquoise2, and mTagBFP2 to further reduce the risk of recombination (Supp. Table S2).

A first test of the compatibility of the four organelle tags was done by targeting a unique fluorescent protein to each organelle in the same cell – specifically, RFP to the nucleus, YFP to the vacuolar membrane, CFP to the plasma membrane, and BFP to actin (Fig. 1A). The resultant cell showed the expected MiCode phenotype when visualized under each non-overlapping fluorescent channel. A few arbitrary MiCodes were then synthesized as a...
further test of feasibility (Fig. 1B). In this small set, we included strains that had organelles simultaneously targeted with two distinct fluorophores as well as organelles without a targeted fluorescent protein. In each of these MiCode examples, the correct genotype was accurately discernable. The most ambiguous phenotype we observed was for vacuolar membrane targeting, which was sometimes similar in appearance to the nucleus. With experience, however, the vacuolar membrane and nucleus could be clearly distinguishable by the intensity of fluorescence in the central region of the organelle or by the quantity of structures (Supp. Fig. S3).

II. Development of a Two-Hybrid Fluorescence Localization Assay for Measuring Protein-Protein Interactions in the Yeast Cytosol

As a first proof-of-concept of applying MiCodes for improved throughput of analyzing a cellular process, we decided to test protein-protein interactions in the cell. Although MiCoding genetic libraries could be useful for a variety of outputs best measured by microscopy, we used subcellular localization as a readout of protein-protein interactions in vivo using a novel two-hybrid assay. Many analytical methods can measure these interactions in vitro such as fluorescence anisotropy, pulldown assays, co-immunoprecipitation, and surface plasmon resonance. Although these methods have proven immensely powerful, they provide indirect measurements and do not capture the cytosolic context of the cell (i.e. measure the protein-protein interaction of interest in the presence of potential off-target interactions in the cytosol).

In our two-hybrid fluorescence localization assay, we used the ability of peroxisomes to import folded proteins, including protein assemblies, to directly probe whether two proteins interact in the cytosol. Import efficiency likely will depend on the size of the complex formed, though the peroxisome import complex has been shown to be capable of accommodating cargo with diameters as wide as 9 nm. A bait protein was tagged with a UV-photoactivatable (PA) GFP and a three amino acid (SKL) C-terminal peroxisome targeting tag 1 (PTS1), while a candidate prey protein was tagged with a fluorescent reporter protein (RFP) (Fig. 2A). The UV-photoactivatable GFP provided the ability to verify that the bait protein was both expressed and properly targeted to the peroxisome without interfering with MiCode analysis (Supp. Fig. S4). An interaction between the bait and prey led to red punctae resulting from co-import of the RFP-tagged prey protein along with the PTS1-tagged bait protein into the peroxisome (left panel, Fig. 2A). The absence of an interaction resulted in diffuse red fluorescence in the cytosol (right panel, Fig. 2A). We used a binary scoring metric for each cell analyzed where observation of a distinct punctate phenotype was scored as a 1 and a completely diffuse phenotype was scored as a 0. Since MiCoding individual strains allows for the assaying of many cells in parallel, the final scores represent averages of multiple cells for increased confidence. Because this assay utilizes the red channel to measure interactions via peroxisome targeting, we used the remaining YFP, CFP, and BFP channels for the MiCode.

III. SYNZIPs and Design of New Specific Coiled-coils

We next sought to design coiled-coil zippers, which are short protein domains of approximately 30 amino acids that can form either coiled-coil homo- or hetero-oligomers.
with alpha-helical structure and a range of affinities. Our eventual goal was to screen a library of newly designed zippers using MiCodes in conjunction with the two-hybrid fluorescence localization assay. New orthogonal synthetic coiled-coils would further expand the synthetic biology toolbox of protein-protein interaction parts to allow more protein complexes to be formed in the yeast cytosol with minimal crosstalk. Several groups have successfully engineered synthetic coiled-coil interaction partners. In particular, Thompson et al. reported 23 synthetic coiled-coil leucine zippers that can form many distinct pairwise complexes. These zippers are referred to as SYNZIPS, a designation given to coiled-coil zippers whose biophysical characteristics and interactions with other zippers have been extensively characterized.

Reinke et al. (2010) and Thompson et al. (2012) discussed the use of both in vitro (coiled-coiled protein microarray and fluorescence polarization) and in vivo (transcription-based yeast two-hybrid and MAPK signaling) assays to measure interactions among SYNZIP protein pairs. The in vitro microarray assay detected interactions between protein pairs by printing one partner onto an aldehyde-derivatized slide and testing it for binding to fluorescent dye-labeled SYNZIP partners. The fluorescence polarization assay mixed maltose-binding protein-SYNZIP fluoresceinated proteins with increasing concentrations of unlabeled maltose-binding protein-SYNZIPs. Dissociation constants (K_d) were determined by performing non-linear least squares regression on curves of the fraction of fluorescein-labeled protein bound for various titrations of unlabeled protein. The transcription-based yeast two-hybrid (Y2H) assay used fusions of proteins to either the Gal4 DNA-binding domain or the Gal4 activation domain. Successful reconstitution of the Gal4 transcription factor, which was mediated by an interaction between partners of interest (SYNZIP pairs), drove expression of auxotrophic marker genes such as URA3 and HIS3. For this assay, colony size was used as a proxy for binding affinity. Finally, the MAPK signaling assay employed the Ste5 protein scaffold and Msg5 phosphatase, a negative modulator of the MAPK-Ste5 signaling cascade in yeast. Individual SYNZIPs were fused to either Ste5 or Msg5 and interactions were read out by expression of a GFP reporter gene controlled by Fus3 activity (Fus3 activity was inhibited by localization of Msg5 to the Ste5 scaffold). An interaction between a protein pair resulted in decreased expression of the GFP reporter. The fractional expression of GFP in comparison to a control with an Ste5 scaffold with no protein fusion was used as a quantitative metric in this assay.

To design new orthogonal coiled coils, we used a scoring function previously developed for predicting bZIP coiled-coil interactions. This scoring function predicts binding strength for a pair of sequences and is based on considering pairs and triplets of residues that would interact if the proteins formed a parallel coiled-coil dimer. The model was trained on a large experimental set of quantified bZIP coiled-coil interactions, and it showed good agreement with experimental binding data in rigorous cross-validation testing. The scoring function can be evaluated quickly because it requires only sequences as input, and it was previously used to design tight and specific binders for four human bZIP proteins. To design orthogonal pairs of coiled coil zippers for this work, sequences were assembled from a library of short 7-residue fragments (heptads) taken from known bZIP proteins, as done previously by Potapov et al. Integer linear programming (ILP) was used to solve for small sets of eight
optimal designed sequences. These sets were combined to give larger sets of 20 peptides predicted to form orthogonal interactions, as described in the Methods.

We used the reported heterodimer dissociation constants of SYNZIPs obtained from a fluorescence polarization assay (measured \textit{in vitro} with purified protein) to calibrate the peroxisome localization assay.\textsuperscript{21} Representative interactions spanning a range of affinities, some of which were also quantified in the MAPK signaling assay, are shown in Supp. Fig. S5. Two-hybrid fluorescence localization interaction scores – obtained by averaging the scores of at least five cells for each strain of a SYNZIP subset – could distinguish interactions with different reported affinities. We determined that the range of interaction affinities that could be measured by the two-hybrid fluorescence localization assay was $\sim 10$-400 nM. Based on the comparison of our interaction scores to the results of the other three assays (Supp. Figs. S5-S7), we defined an interaction score threshold of $\geq 0.4$ (K$_{d} < 200$ nM) for a strong interaction and interaction scores 0-0.4 (K$_{d} > 200$ nM) for weak interaction.

To further compare our two-hybrid fluorescence localization assay to previous assays, we tested 68 pairwise interactions between 14 SYNZIPs previously analyzed by Reinke et al.\textsuperscript{11} and by Thompson et al. (Supp. Fig. S6).\textsuperscript{21} We cloned the proteins involved in 27 interaction pairs in both bait-prey and prey-bait orientations, for a total of 54 heterodimer interactions, and additionally tested 14 homodimer interactions. Our two-hybrid fluorescence localization assay agrees moderately well with the interaction profiles reported using fluorescence polarization assays, transcription-based Y2H, and MAPK signaling assays. As shown in Supp. Figs. S6 and S7, most SYNZIPs showed similar interaction strengths for the microarray and two-hybrid fluorescence localization assays; however, there were exceptions. Of the 68 candidate SYNZIP interactions tested using both of these methods, two were diagnosed as strongly interacting by the microarray assay but categorized as weakly interacting by the two-hybrid fluorescence localization assay, while six candidate interactions were diagnosed as extremely weak by the microarray assay but strong by the two-hybrid fluorescence localization assay. Overall this eight of the 68 disagreement with the microarray results (12\%) is similar to the range of variation seen between the coiled-coil microarray and transcription-based Y2H assays (10-13\%) (Supp. Fig. S7B).\textsuperscript{11} As discussed by Thompson et al. discrepancies between the in vitro and in vivo interaction assays demonstrate the need to test interactions in the context in which they will be used.\textsuperscript{21} To this end, the two-hybrid fluorescent localization assay serves as an additional platform for validation as well as discovery of protein-protein interactions in the cytosol – it should not be used as a sole measurement of a protein-protein interaction.

Various comparisons can be made between our two-hybrid fluorescence localization assay and the in vivo and in vitro assays described by Thompson et al.\textsuperscript{21} Although the in vitro fluorescence polarization assay provides quantitative binding affinities, these affinities may not be a precise predictor of protein interaction behavior in a cytosolic context. Y2H resolves some of these potential problems and can be assayed in a quantitative manner, but its output is an indirect measure of the binding and can be confounded by other factors such as non-specific or indirect interactions.\textsuperscript{24,25} The MAPK signaling assay is similar in this regard. The two-hybrid fluorescence localization assay provides information on the SYNZIP
binding performance in vivo with single-cell resolution. Instead of measuring the output of a linked transcriptional response, this assay directly couples a protein interaction to co-localization, giving an output that is a direct readout of steric interaction. An additional benefit of the two-hybrid fluorescence localization assay is the detection of off-target localization. In our initial screen of newly designed coiled-coils, we observed a few of our newly designed prey proteins fused to RFP mistargeting to the nucleus and peroxisome (R.C. unpublished data). In sum, our two-hybrid fluorescence localization assay expands the toolkit of protein interaction assays by providing a direct in vivo measurement of protein binding that is amenable to microscopy.

IV. Application of MiCoding Towards the Screening of a Library of Coiled-Coil Zippers

Towards enabling higher-throughput microscopy screening of candidate protein interaction pairs, we added MiCode tags to a single pair of newly designed coiled-coil proteins, 100A and 100B designed to heterodimerize (see Methods). For testing of the interaction between 100A and 100B using the two-hybrid fluorescence localization assay, we designated 100A as the bait and 100B as the prey. The newly designed bait zipper (100A) and prey zipper (100B) were cloned separately and each was incorporated into a DNA segment that included half of the eventual full MiCode (Fig. 2B and Supp. Fig. S2). In this single-pair mock assembly (i.e. the identities of the bait and prey were known a priori; however, the bait-prey binding affinity was uncharacterized by other assays), the gene expressing the bait construct was physically linked to the genes expressing the plasma membrane and actin-targeting half of the MiCode. Similarly, the gene expressing the prey construct was cloned next to the genes expressing the other half of the MiCode (nucleus and vacuole membrane targeting). The prey zipper and its half-MiCode were integrated into the LEU2 locus, followed by integration of the bait zipper and other half-MiCode into the URA3 locus. Fig. 2B shows representative strains containing either the bait-half MiCode (top panel) or prey-half MiCode (bottom panel) constructs integrated in the chromosome. A full MiCode was created through co-integration of the bait and prey half-MiCoded constructs into the same strain (Fig. 2C). Upon inspection of the fully-MiCoded test strain via microscopy, we observed punctate fluorescence in the RFP channel yielding an interaction score of 1 between bait 100A and prey 100B. The identities of both the bait and prey were verified by identifying the targeted organelles for each of the YFP, CFP, and BFP channels. In the YFP channel, the plasma membrane is visible. In the CFP channel, the vacuolar membrane is visible, while no organelle is visible in the BFP channel.

Next, we extended the MiCoding strategy to the simultaneous testing of the entire two-by-two interaction matrix for the 100A/100B zipper pairing. The complete interaction matrix contained four experimental configurations: bait/prey 100A/100A (test for 100A homodimerization), 100B/100A (test for heterodimerization in orientation one), 100A/100B (test for heterodimerization in orientation two), and 100B/100B (test for 100B homodimerization). We first tested these four interaction pairings clonally without MiCoding to avoid any possible physiological effect of the MiCode on the interaction phenotype (R.C. unpublished data). Next, we tested all four pairings simultaneously by MiCoding each configuration and screening the resulting mixed population (Fig 3). Since our assembly strategy pre-assigned bait and prey proteins each with a specific half-MiCode.
(Fig. 3C and Supp. Fig. S2), the full MiCode phenotype allows us to extract the genetic identity of the interacting coiled-coil pair. Cells representative of the four experimental configurations from the two-by-two matrix are boxed in Fig. 3A and 3B and highlighted in Fig. 3C along with the MiCodes and the decoded bait/prey identities. As shown in Fig. 3D, the interaction scores could be rapidly determined by surveying a large number of cells. We calculated the interaction score for each bait/prey configuration by averaging the binary output of the assay – interaction (1) or no interaction (0) – over all the cells corresponding to that genotype in the field of view. In addition to reporting this numerical score, we also present the number (counts) of cells of each genotype present on the micrograph used for this calculation. From this small 2×2 matrix screen, we report that protein pair 100A/100B exhibits only heterodimerization, with no detectable homodimerization. Furthermore, we note that in this test the presence of the MiCode did not affect the phenotypic output of the interaction assay.

Encouraged by these results, we screened a larger set of novel designed coiled-coils with MiCoding. Ten new protein pairs (101A/B-110A/B) were designed to form a set of orthogonal heterodimers (see Methods, Supp. Fig. S8, and Supp. Table S5). Each designed pair was pre-tested in a two-hybrid localization assay screen for heterodimerization. Eight of ten pairs showed clear evidence for interaction – a score greater than 0.4 – with the intended partner (Supp. Fig. S9). Three of 20 constructs severely inhibited cell growth, and three of the eight designed pairs gave asymmetrical interaction results with respect to the bait/prey configuration (Supp. Fig. S9). We chose four of the remaining five well-behaved coiled-coil pairs for screening in a full 8×8 matrix of 64 potential interactions (Fig. 4B).

To determine whether each designed protein heterodimerized exclusively with its cognate pair in the set (i.e. exhibited no interactions with other proteins in the matrix), we MiCoded and tested all possible interactions between noncognate pairs, including potential homodimers. To assemble all possible interacting pairs, we first generated the eight half-MiCoded prey strains individually. Next, we pooled the prey strains in a one-pot reaction and transformed a cocktail consisting of the eight bait half-MiCode integration cassettes to generate the full library of MiCoded interacting partners. A representative series of images for this assay, run with all variants pooled, is shown in Fig. 4A. Thirty-six images were taken in each of the four channels to acquire multiple cell counts of each of the 64 matrix members (Fig. 4B). As expected, all of the cognate pairs showed interactions in a high fraction of the cells analyzed, with the lone exception of bait/prey 101B/101A. Many proteins also showed evidence of undesired homodimerization, notably proteins 101A, 102A, 102B, and 108B. Although there was a much lower incidence of non-cognate and off-target interactions than cognate interactions, several were observed such as 101A/101A (non-cognate) and most prey coiled-coils with bait 108B (off-target). In some cases like 108B, we observed asymmetric results: bait 108B interacted showed some binding to all prey proteins, whereas prey 108B showed little or no binding to most bait proteins.

We again probed the possibility that the addition of a MiCode could perturb the assay or impede cell growth. First, we examined the interactions of designed cognate coiled-coil pairs with and without the MiCode tag (Fig. 4C). Most (seven of eight) of the cognate heterodimer pairings exhibited a similar interaction phenotype both with and without the
MiCode as expected. Notably, the interaction of bait 101B and prey 101A was much weaker in the MiCoded assay than in the un-MiCoded assay. Thus, with the exception of 101B/101A, expression of the MiCode did not seem to impact the ability to assay a positive interaction. Another consideration is the potential growth inhibition resulting from over-targeting certain organelles. For example, most strains with bait 108A were present in lower abundance (i.e. had lower cell counts) in the library than expected. All strains with bait 108A had actin targeted by both Venus and mTurquoise2, suggesting that double-targeting actin with those fluorescent proteins in combination is mildly toxic to the respective host strains causing low cell counts (Supp. Fig. S10). To acquire appreciable cell counts, we grew these problematic MiCoded strains individually using exactly the same MiCodes to measure candidate interactions reported in Figure 4B. We previously observed a similar toxicity when double-targeting the vacuole membrane using any two of YFP, CFP, or BFP, and therefore we removed all double-targeted vacuole membrane containing MiCodes (surprisingly, any combination of YFP, CFP, or BFP with RFP targeted to the vacuole membrane is not toxic, such as the strain shown in Fig. 1B). This toxicity is particularly an issue when growing the library collectively in a single batch where these MiCode-containing clones can be outcompeted by other strains. Such growth biases have been known to be present in library-based approaches. For example, mutants in pooled, high-throughput transposon mutagenesis libraries can grow at different rates based on how the transposon insertion perturbs the genetic context of each variant, thereby resulting in library members to be present at different abundances.

We present the results for two small MiCoded libraries, a 2×2 and an 8×8 matrix, that we used to screen newly designed coiled coils that were computationally optimized for orthogonality. In the 2×2 matrix, we tested the four possible interactions – two cognate and two non-cognate – with and without the MiCode and observed no significant differences in the results between the two methods. Importantly, MiCoding improved the ease of screening and allowed us to extract all the results from a single yeast transformation reaction and a small number of micrographs. Encouraged by the agreement between the MiCoded and the non-MiCoded data, we moved toward a larger set of twenty proteins (ten pairs), which we pared down to an 8×8 matrix screen based on knowledge gained from an initial non-MiCoded interaction screen of designed cognate pairs. We then performed a MiCoded screen of the 8×8 matrix to acquire data for all 64 possible interactions in order to screen for off-target and homodimer interactions. In this case, we report good, but not perfect, agreement between the MiCoded and non-MiCoded data.

Our interaction data demonstrate that achieving orthogonality in this cellular localization assay was difficult, even when proteins were designed with this explicit goal in mind. Regardless, we report two new coiled-coil pairs, 100A/100B from the 2×2 matrix screen and 107A/107B from the 8×8 matrix screen, that can be used for synthetic system designs where hetero-oligomerization without homo-oligomerization is desired. With respect to the unintended interactions, we emphasize that we have not definitively determined the sensitivity of the localization assay compared to that of other techniques. Although the overall agreement of the new assay with previously reported SYNZIP interaction data was good, it is also true that SYNZIP pairs not previously reported to interact by any of the four other assays were positive for our screen. Furthermore, it is important to note that the
interaction geometry and stoichiometry of complexes formed by the designed proteins has not been determined. The computational design procedure considered only parallel coiled-coil heterodimer structures, yet small changes in sequence can lead to changes in coiled-coil assembly. Thus, there is a possibility that the unintended complexes are higher-order homo- or hetero-oligomers, or dimers with helices aligned in an antiparallel orientation. Indeed, retrospective computational analysis indicated that pairs 101A/101A, 102A/102A, 102B/102B, 101B/108B, 102B/108B, 107A/108B might form strong interactions when the constituent alpha helices associate in an anti-parallel mode. In such a mode, charged residues at the e and g helical positions in these pairs could form attractive interhelical Coulombic interactions. This finding highlights the importance of considering multiple competing states when designing coiled coils. Recently, an improved model for scoring anti-parallel coiled-coil interactions was developed that could be used to augment the design protocol used in this work.

While finding orthogonal coiled-coiled zipper pairs from a rationally-designed matrix of interaction candidates proved difficult, the use of MiCodes greatly increased the speed of the screening process by allowing for data collection and analysis of pooled, heterogenous samples. MiCoding allowed many cells to be analyzed simultaneously in a single micrograph, which enabled a larger matrix of proteins to be assayed while still achieving replicates for analysis. In conclusion, the two-hybrid fluorescence localization screen, when coupled with MiCodes, enabled quick screening of a large matrix of interactions through mixed-sample analysis and should facilitate faster iterations of the design-build-test-learn cycle in future studies.

V. Scaling the Size of a MiCoded Library

In this study, we used only a small subset of the potential MiCode library as required by the size of our experiment. However, it should be feasible to conduct experiments with considerably larger libraries where each individual member is uniquely MiCoded. As the size of the MiCode library is scaled, however, a few considerations should be made. First, care should be taken to choose organelle-targeting tags that both produce a phenotypically distinct MiCode and confer no physiological growth defect to the host. Expression of additional organelle-targeting tags will increase the risk of toxicity. Although we optimized expression level of the organelle targeting tags for a widefield fluorescence microscope, it is possible that the use of a more sensitive microscope or method such as confocal microscopy could achieve a higher signal-to-noise ratio and potentially allow the expression levels of these targeting tags to be lowered below the toxicity threshold, even with multiple targeting. Another consideration is the difficulty in correctly identifying organelles that can look similar. Clearly, this is the foremost consideration for incorporating additional organelles into a MiCode set. Nuclear and vacuolar membrane targeting is an example of an easily misdiagnosed targeting, although experience in identification considerably improves the accuracy of MiCode identification (Supp. Fig. S3).

The size of MiCode libraries could be exponentially increased through the addition of channel states, organelles, or fluorescent channels – provided they are clearly discernable from the other organelles and fluorescent channels. For example, a strategy for increasing
the number of channel states is to use a UV-photoactivatable (PA) fluorescent protein. All other fluorescent proteins could be photobleached followed by UV stimulation of the PA fluorescent proteins. Preliminary tests of PA fluorescent proteins were promising (R.C. unpublished data). If these PA proteins could be implemented without interfering with the ON/OFF states of other fluorescent proteins, this could give each fluorescent channel three states (ON/OFF/PA) instead of the two described above. Further expansion of the number of channel states could be achieved by precise modulation of fluorescence intensity, giving both a STRONG and WEAK ON states.

Additional fluorescent channels could be incorporated through peptide-fused quantum dots as fluorescent protein analogues. Quantum dots are fluorescent nanoparticles with precise emission wavelengths that can be tuned to allow additional channels to be multiplexed. When conjugated with organelle-targeting peptides, quantum dots have been shown to be taken up by live cells and targeted to organelles. A scaffold molecule, protein, DNA, or RNA, could be expressed and targeted to organelles to recruit different patterns of quantum dots yielding unique MiCodes with linkages to known genotypes. The MiCoding platform can also potentially be coupled with other in vivo surface labeling probes to further increase barcode diversity.

Increased numbers of organelles may be achievable, particularly if the confidence in accurate identification is improved by combining fluorescent microscopy with organelle staining methods using compatible fluorescent dyes or analytical approaches such as confocal microscopy that can improve signal over background. For example, there are additional organelles such as the mitochondria that we have not fully explored which could be compatibly-targeted. It may even be feasible to subdivide an organelle, such as the nucleus and nucleolus.

Returning to the equation presented in the Introduction:

\[ m = f(s, c, o) = (s^c)^o \quad (1) \]

where ‘m’ is the number of MiCodes, ‘s’ is the number of channel states, ‘c’ is the number of fluorescent channels, and ‘o’ is the number of organelles. Here, we leveraged the flexibility of MiCode library design and assembly with our prior knowledge of morphologically distinct fluorescent protein-targeting tag combinations to create a subset of easily identifiable MiCodes that satisfied our screening requirements. With this strategy, there were:

\[
m = \#(\text{bait MiCodes}) \times \#(\text{prey MiCodes})
\]

\[
= (s_{\text{GFP,b}} \wedge o_{\text{GFP,b}} \wedge s_{\text{CFP,b}} \wedge o_{\text{CFP,b}} \wedge s_{\text{BFP,b}} \wedge o_{\text{BFP,b}}) \times (s_{\text{GFP,p}} \wedge o_{\text{GFP,p}} \wedge s_{\text{CFP,p}} \wedge o_{\text{CFP,p}} \wedge s_{\text{BFP,p}} \wedge o_{\text{BFP,p}})
\]

\[
= (2^2 \times 2^2 \times 2^2 \times 1) \times (2^2 \times 2^2 \times 2^2 \times 1)
\]

\[
= 1,024 \text{ possible MiCodes, where } b=\text{bait and } p=\text{prey}
\]
If all fluorescent channels are used for all organelles, then \( m = (s \land c) \land o = (2 \land 4)^4 = 65,536 \) possible MiCodes. If the system can be expanded using the techniques described above to increase the number of fluorescent channels by one, the MiCode library size can be expanded to \((2 \land 5)^4 = 1,048,576\). Adding a third state to these fluorescent channels can increase this further: \((3 \land 5)^4 = 3,486,784,401\).

The cloning of such large libraries will likely require one-pot assembly strategies rather than the individual construction of each library member. Using a scheme analogous to the Combinatorial Genetics En Masse (CombiGEM) approach for assembling higher-order combinations of DNA barcoded constructs, it should be possible to construct large-scale combinatorial libraries of MiCoded constructs (Supp. Figs. S11 and S12). These DNA-barcoded MiCode constructs could then be combined with a set of DNA-barcoded assay constructs (i.e. bait-prey constructs for the leucine zipper assay) and the two barcodes can be associated using Illumina next-generation sequencing (Supp. Fig. S13). In this manner, the MiCode and experimental phenotype can be visualized in parallel on the microscope and the unique identity of the library member can be extracted using the DNA barcodes associated with the MiCode and experimental genotype.

For analysis of larger libraries, it should also be feasible to design algorithms for automating MiCode image acquisition and identification. Publicly-available cell segmentation software as well as custom scripts can be used to perform automated cell segmentation of MiCoded cells through edge detection and morphology-based filtering with up to 90% accuracy. It should be feasible to automate MiCode identification (i.e. calling different organelles) by using morphology-based filtering thresholds derived from geometric analysis of a known MiCode dataset. For example, the actin and nucleus organelles could be distinguished by their respective radii. Even so, more sophisticated algorithms applying supervised machine learning on pre-identified MiCode micrographs will likely be necessary for robust image analysis identification and represent an area of future focus.

**VI. Summary**

Overall, MiCoding should prove useful for a variety of outputs measurable by microscopy but not amenable to high-throughput FACS. In this work, we first created tags for morphologically-distinct organelles, then expanded the MiCode library size by adding several spectrally-unique fluorescent channels. The diversity of a MiCoded library can be further scaled by increasing the number of channel states through photoactivatable proteins, subcellular tags through further subdivision, and fluorescent channels using quantum dots.

In addition, the MiCoding platform should prove generalizable and be applicable to other eukaryotic hosts. Possible future studies that could benefit from MiCoding are the exploration of genetic circuitry controlling neutrophil chemotaxis, the developmental tracking of cell lineages in small multicellular organisms such as Caenorhabditis elegans, and mechanobiological studies. For example, MiCoding could allow for the rapid surveying of genome-wide gene knockout or CRISPRi-mediated knockdown libraries at the single-cell level by enabling the identification of potentially undiscovered morphologies in pooled screens across a variety of conditions. This work represents a first step towards these
exciting applications, allowing outputs clearly observable by microscopy to be analyzed in higher-throughput.

**Methods**

**MiCode Design and Construction**

All cloning was performed using the Yeast Toolkit Golden Gate assembly strategy. Single-gene plasmids containing a single organelle and fluorescence tag fusion or a single SYNZIP were transformed in TG1 chemically competent *E. coli* and were selected on LB containing ampicillin (100 mg/L). In subsequent reactions, the single-gene plasmids were assembled together into half-MiCode plasmids. As described earlier, we rationally designed the half-MiCode plasmids such that they linked the SYNZIP to pre-assigned MiCodes. These half-MiCode plasmids were transformed into TransforMax EPI300 electrocompetent *E. coli* and were selected on LB containing kanamycin (25 mg/L).

In all the constructs, any homologous regions that could potentially lead to unwanted recombination were eliminated. Each cassette within the system was expressed using different promoters of equivalent expression level and different terminators of equivalent termination efficiency. Additionally, an alternate version with different codon usage was generated for each of the MiCode tags and each of the fluorescent proteins. The codons were randomly chosen based on *S. cerevisiae* codon usage frequencies, and any rare codons were removed.

**SYNZIP Design**

We aimed to design several pairs of coiled-coil proteins (N pairs, 2N proteins) such that the first protein in a pair forms a strong interaction with the second protein from the same pair but forms only weak interactions with any other protein in the designed set. Interactions in sets of pairs that satisfy these criteria are called orthogonal interactions. To design orthogonal coiled coils, we used a scoring function previously developed for predicting bZIP coiled-coil interactions. As in prior work, sequences composed of 7-residue fragments from known bZIP proteins were optimized with integer linear programming, with constraints imposed to favor desired patterns of binding and non-binding. Specifically, we minimized the sum of the predicted binding scores for cognate interactions:

\[
S = \sum_{i=1}^{N} S(A_i, B_i) \tag{2}
\]

where \(N\) is the number of designed pairs and \(S(A_i, B_i)\) is the predicted binding score between protein A and protein B in the \(i^{th}\) designed pair. Simultaneously, constraints were imposed to ensure that non-cognate interactions were weak. In particular, to prevent formation of homodimers, we imposed constraints \(S(A_i, A_i) > S_{cutoff}\) and \(S(B_i, B_i) > S_{cutoff}\) for each designed pair \(i=1,N\), where \(S_{cutoff}\) was a predefined value chosen to ensure that homo-dimer interactions were at least 100-fold weaker compared to cognate interactions. Similar constraints were imposed for other non-cognate interactions between sequences from different pairs (off-target interactions).
Simultaneously optimizing a large number of designed sequences and their interactions is a computationally demanding task even when using a simple scoring function. We were able to optimize four pairs of sequences at a time using a subset of the heptad library. To design a larger set of pairs, we repeated the design procedure many times, every time using a random subset of heptads for building designs. Following this, a set of 10 pairs was obtained by taking many four-pair sets and searching for larger sets of pairs with low potential for off-target interactions (Supp. Table S5, Supp. Fig. S5). An initial set of 10 predicted orthogonal pairs was reduced to 4 pairs selected for further testing, based on an initial screen of the predicted heterodimers using the novel two-hybrid fluorescence localization assay (Supp. Fig. S6). In this initial screen, toxic proteins that reduced cell growth, and asymmetric-strength protein pair interactions for which prey-bait and bait-prey cognate interactions differed in strength, were eliminated.

Strains and Media

All experiments were conducted in a BY4741 *S. cerevisiae* strain background (MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0). Multi-gene MiCode and zipper assay plasmids containing Leu2 and Ura3 auxotrophic markers were integrated into the respective chromosomal loci. Multi-gene plasmids were linearized with NotI then transformed using a standard lithium acetate transformation protocol. Following heat shock, the cells were selected and grown on synthetic dropout media (6.7 g/L Difco Yeast Nitrogen Base without amino acids; 2 g/L US Biological Drop-out Mix Synthetic. Minus Leucine, Uracil, or Leucine and Uracil without Yeast Nitrogen Base; 20 g/L Dextrose). All *S. cerevisiae* strains were grown at 30°C.

To form a complete MiCode, a bait and prey multi-gene plasmid were co-integrated to form a full MiCode strain. This process was used to create the strains shown in Figs. 1 and 2. The full library (Fig. 3) was created through a pooled co-integration with all the bait multi-gene plasmids. Each member the pooled set of bait plasmids was standardized to 1.5 pmol of DNA. Prior to the pooled library reaction, each individual prey MiCode strain was verified through fluorescence microscopy. In summary, to form the full library two successive rounds of integrations were performed, first selecting on leucine drop-out media then on uracil and leucine drop-out media.

I-SceI Integration

In order to increase the efficiency of these chromosomal integrations, as well as lower the percentage of multiple integration events, we first inserted an I-SceI homing endonuclease recognition site into the *URA3* locus that would later be targeted for integration. The I-SceI landing pad was integrated into the *URA3* locus by inducing double-stranded break with CRISPR/Cas9 and repairing the site with a PCR cassette containing the I-SceI recognition site. Transformation and expression of SceI is expected to create a double-stranded break in the chromosome and increase the efficiency of homologous recombination at this target site. During the second transformation, the pooled multi-gene MiCode plasmids were co-transformed with a markerless plasmid to transiently express I-SceI. Testing determined that 40 fmoles of this plasmid per integration reaction was the optimum amount and that it increased integration efficiency by 9.7-fold.
Microscopy

Images were taken on either a Zeiss Axio Observer D1 bright field or Zeiss LSM710 confocal microscope. The Axio Observer D1 offered greater throughput but sacrificed the spatial resolution that the LSM710 could provide. Samples were prepared from a mid-log culture (grown for ~4 hours to 0.6 OD600). Cultures were pelleted in a microcentrifuge at 8000rpm for 1min, then resuspended in a tenth of the original volume in PBS. The liquid was covered with a coverslip coated with concanavalin A: a 2mg/mL solution of concanavalin A was added onto the coverslip, incubated for 10 minutes, and then washed with PBS. Then, 2.5uL of the concentrated culture was added to a glass slide and covered with the concanavalin A-coated coverslip.

Axio Observer D1 images were taken at 1000ms exposure for the mKate channel, 1000ms for the Venus channel, 200ms for the mTurquoise2 channel, 1000ms for the mTagBFP2 channel, and 25ms for the DIC image. Zeiss LSM710 images were taken at a scan speed of 4 and an averaging mode of 8. Gain for each channel was set to sub-saturation values for each sample. Microscopy files were viewed and then exported to images in Fiji under an auto exposure setting. Several images were taken per slide, and the fields of view were spaced to prevent photobleaching. Each micrograph was contrast-adjusted for clarity.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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References


Figure 1. MiCode identification. (A) Cartoon yeast cells and fluorescent micrographs depicting the various subcellular organelle tags and fluorescent proteins used in constructing a MiCode. Four organelle tags are used: nucleus (HTA2), vacuolar membrane (ZRC1), plasma membrane (CIIC tag), and actin (ABP1). Four fluorescent proteins are used: mRuby2 (RFP), Venus (YFP), mTurquoise2 (CFP), and mTagBFP2 (BFP). (B) Several examples of randomly-selected MiCodes for demonstration purposes. All scale bars are 10 μm, and all micrographs were contrast-adjusted for clarity.
Figure 2.
Two-hybrid fluorescence localization assay for probing protein-protein interactions. (A) Cartoon yeast cells and fluorescent micrographs depicting strong and weak binding scenarios. The bait protein is also fused to a photoactivatable GFP protein (not shown for clarity) that is used for verifying peroxisome localization of the bait protein (Supp. Fig. S4). The strong binding example strain is bait/prey SYNZIP2/SYNZIP1 with a reported K_d of <10 nM, while the weak binding example is of SYNZIP18/SYNZIP21 with a reported K_d of >300 nM. (B) Cartoon yeast cells and fluorescent micrographs of two strains each containing a half-MiCode, either only the bait or prey half. The cells were digitally magnified above the microscope's 1000× magnification to better illustrate the organelle targeting. Because the prey is tagged with RFP, when only it is expressed, there is diffuse red fluorescence. Below the micrographs, we present gene locus maps showing the half-MiCode and bait/prey expression cassettes in situ in the yeast genome. Grey regions in the maps indicate spacer sequences. The order of the fluorescent proteins and localization tags reflects their position as a result of the Golden Gate cloning strategy (Supp. Fig. S2). (C) Cartoon yeast cell and fluorescent micrographs of a strain containing a full MiCode with both the bait and prey. In this case, there is a strong observable interaction between bait/prey 100A/100B. All scale bars are 10 μm, and all micrographs were contrast-adjusted for clarity.
Figure 3.
An example library that links the leucine zipper assay to pre-assigned MiCodes. (A) Leucine zipper assay with cells representative of the four MiCode phenotypes indicated. (B) Merged color channels used for MiCode identification. (C) Isolated examples of the four MiCode phenotypes possible in this small library. In the MiCode column, the half-MiCodes used for the prey and bait are boxed in orange. (D) Interaction scores and interaction counts are based on solely the images in Figs. 3A and 3B. The interaction scores across cells were computed by scoring all cells in a micograph on binary scale: 0 for a lack of a visible interaction and 1 for any visible punctae. The counts value corresponding to each bait/prey pairing represents the number of cells used to calculate the average binding score. The values were then averaged over the given number of cells to yield the final score. The micographs were contrast-adjusted for clarity.
Figure 4.
MiCodes applied to assaying an 8×8 array of coiled-coils. (A) A representative image of the 8×8 library that contains a mixed population of all bait-prey interactions. (B) Tabulated data for the two-hybrid fluorescence localization assay experiment showing interaction scores and interaction counts. These data are averaged across thirty-six micrographs. These results were supplemented with micrographs of single monoclonal strains for bait/prey 107A/107B, 107B/102B, 108A/101A, 108A/101B, 108A/102A, 108A/102B, 108A/107A, 108A/107B, 108A/108A, and 108A/108B, whose results were added to pool. Unsupplemented data are shown in Supp. Fig. S10. (C) Tabulated data for the no-MiCode control strains that was used for validating the cognate heterooligomer interactions. Each of the eight cognate coiled-coil interactions was assayed using the two-hybrid fluorescence localization assay. The micographs were contrast-adjusted for clarity.