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TITLE: Enhanced yeast one-hybrid screens to identify transcription factor binding to human DNA sequences

AUTHORS AND AFFILIATIONS: Shaleen Shrestha¹, Xing Liu¹, Clarissa Stephanie Santoso¹, Juan Ignacio Fuxman Bass^{1,2}

¹ Department of Biology, Boston University, Boston, MA

² Bioinformatics Program, Boston University, Boston, MA

Corresponding Author:

Juan Ignacio Fuxman Bass

fuxman@bu.edu

Tel: (617)-353-2448

Email Addresses of Co-authors:

Shaleen Shrestha (shaleens@bu.edu)

Xing Liu (liuxing@bu.edu)

Clarissa Stephanie Santoso (csantoso@bu.edu)

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SUMMARY: Here, we present an enhanced yeast one-hybrid screening protocol to identify the transcription factors that can bind to a human DNA region of interest. This method uses a high-throughput screening pipeline that can interrogate the binding of >1,000 TFs in a single experiment.

ABSTRACT:

Identifying the sets of transcription factors (TFs) that regulate each human gene is a daunting task that requires integrating numerous experimental and computational approaches. One such method is yeast one-hybrid (Y1H) assays, in which interactions between TFs and DNA regions are tested in the milieu of the yeast nucleus using reporter genes. Y1H assays involve two components: a 'DNA-bait' (*e.g.*, promoters, enhancers, silencers, etc.) and a 'TF-prey', which can be screened for reporter gene activation. Most published protocols for performing Y1H screens are based on transforming TF-prey libraries or arrays into DNA-bait yeast strains. Here, we describe a pipeline, called enhanced Y1H (eY1H) assays, where TF-DNA interactions are interrogated by mating DNA-bait strains with an arrayed collection of TF-prey strains using a high density array (HDA) robotic platform that allows screening in a 1,536 colony format. This allows for a dramatic increase in throughput (60 DNA-bait sequences against >1,000 TFs takes two weeks per researcher) and reproducibility. We illustrate the different types of expected results by testing human promoter sequences against an array of 1,086 human TFs, as well as examples of issues that can arise during screens and how to troubleshoot them.

INTRODUCTION:

A central problem in the biomedical field is determining the mechanisms by which each human gene is regulated. Transcription is the first step in controlling gene expression levels, and it is regulated by sets of transcription factors (TFs) that are unique to each gene. Given that humans encode for >1,500 TFs^{1,2}, identifying the complete set of TFs that control the expression of each gene remains an open challenge.

Two types of methods can be used to map TF-DNA interactions: TF-centered and DNA-centered methods³ (**Figure 1A**). In TF-centered methods a TF of interest is probed for binding to genomic DNA regions or to determine its DNA binding specificity. These methods include chromatin immunoprecipitation (ChIP) followed by high-throughput sequencing, protein binding microarrays, and SELEX⁴⁻⁶. In DNA-centered methods a DNA sequence of interest is probed to determine the set of TFs that bind to the DNA sequence. The most widely applied of such methods is yeast one-hybrid (Y1H) assays, in which interactions between TFs and DNA regions are tested in the milieu of the yeast nucleus using reporter genes⁷⁻⁹.

Y1H assays involve two components: a 'DNA-bait' (e.g., promoters, enhancers, silencers, etc.) and a 'TF-prey', which can be screened for reporter gene activation^{9,10} (**Figure 1B**). The DNA-bait is cloned upstream of two reporter genes (*LacZ* and *HIS3*) and both DNA-bait::reporter constructs are integrated into the yeast genome to generate chromatinized 'DNA-bait strains'. The TF-prey, encoded in a plasmid that expresses a TF fused to the activation domain (AD) of the yeast Gal4 TF, is introduced into the DNA-bait strain to fish for TF-DNA interactions. If the TF-prey binds to the DNA-bait sequence, then the AD present in the TF-prey will lead to the activation of both reporter genes. As a result, cells with a positive interaction can be selected for growth on plates lacking histidine, as well as overcoming a competitive inhibitor, 3-Amino-1,2,4-triazole (3-AT), and visualized as blue colonies in the presence of X-gal. Because the potent yeast Gal4 AD is used, Y1H assays can detect interactions involving transcriptional activators as well as repressors. In addition, given that TF-preys are expressed from a strong yeast promoter (*ADH1*), interactions can be detected even for TFs that have low endogenous expression levels, which are challenging to detect by ChIP^{11,12}.

Most published protocols for performing Y1H assays are based on introducing TF-preys into the yeast DNA-bait strains by transforming pooled TF-prey libraries followed by selection, colony picking, and sequencing to identify the interacting TF, or by transforming individual clones^{8,9}. These are time-consuming protocols, limiting the number of DNA sequences that can be tested per researcher. A recent improvement of Y1H assays, called enhanced Y1H (eY1H), has dramatically increased the screening throughput by using a high density array (HDA) robotic platform to mate yeast DNA-bait strains with a collection of yeast strains each expressing a different TF-prey^{10,13} (**Figure 1C**). These screens employ a 1,536 colony format allowing to test most human TFs in quadruplicate using only three plates. Further, given that TF-DNA interactions are tested in a pairwise manner, this approach allows for comparing interactions between DNA-baits (such as two noncoding single nucleotide variants) and between different TFs or TF variants^{11,12,14}.

Using eY1H assays we have delineated the largest human and *Caenorhabditis elegans* DNA-centered TF-DNA interactions networks to-date. In particular, we have identified 2,230 interactions between 246 human developmental enhancers and 283 TFs¹². Further, we have

employed eY1H assays to uncover altered TF binding to 109 single nucleotide noncoding variants associated with genetic diseases such as developmental malformation, cancer, and neurological disorders. More recently, we used eY1H to delineate a network comprising 21,714 interactions between 2,576 *C. elegans* gene promoters and 366 TFs¹¹. This network was instrumental to uncover the functional role of dozens of *C. elegans* TFs.

The protocols to generate DNA-bait stains and evaluate the levels of background reporter activity have been reported elsewhere¹⁵⁻¹⁷. Here, we describe an eY1H pipeline that can be used to screen any human genomic DNA region against an array of 1,086 human TFs. Once a yeast DNA-bait strain is generated and a TF-prey array is spotted onto the corresponding plates, the entire protocol can be performed in two weeks (**Table 1**). More importantly, the protocol can be parallelized so that a single researcher can screen 60 DNA-bait sequences simultaneously. To demonstrate the protocol we screened the promoters of two cytokine genes CCL15 and IL17F. In addition, we show results from failed screens to illustrate the types of problems that may arise when performing eY1H assays and how to troubleshoot them.

PROTOCOL:

1. Preparations

1.1. Sc –U –H plates (150 mm Petri dishes). These plates will be used for growing the DNA-bait yeast strains.

1.1.1. Dissolve the drop-out mix, the YNB, adenine hemisulfate, and the ammonium sulfate in 920 mL of water, and pH to 5.9 with NaOH 5M (approximately 1 mL per liter of media) (see **Table 2** for composition). Pour into a 2-L flask and add a stir bar.

1.1.2. In a second 2-L flask, add the agar to 950 mL of water (do not add a stir bar as it will cause the agar to boil over in the autoclave).

1.1.3. Autoclave for 40 min at 15 psi on liquid cycle.

1.1.4. Immediately pour the contents of the first flask, including the stir bar, into the agar containing flask. Add the glucose, mix well on a stir plate and cool to 55°C in a waterbath.

1.1.5. Add the leucine and the tryptophan to the media. Mix well on a stir plate and pour into 150-mm sterile Petri dishes (~80 mL per dish). Dry for 3-5 d at room temperature, wrap in plastic bags, and store at room temperature for up to 6 months.

1.2. YAPD rectangular PlusPlates. These plates will be used for growing the lawn for the DNA-bait strain and for mating with the TF array collection.

1.2.1. Dissolve powders (see **Table 3** for composition), except for agar, in 950 mL of water in a 2-L flask and add a stir bar.

1.2.2. In a second 2-L flask, add the agar to 950 mL of water (do not add a stir bar as it will cause the agar to boil over in the autoclave).

1.2.3. Autoclave for 40 min at 15 psi on liquid cycle.

1.2.4. Immediately pour the contents of the first flask, including the stir bar, into the agar containing flask.

1.2.5. Add the glucose, mix well on a stir plate and cool to 55°C. Pour media into the PlusPlates (~70 mL per plate) using a peristaltic pump (5 mL per sec) and a 6 mm tubing. Dry for 1 day at room temperature, wrap in plastic bags, and store in the cold room for up to 6 months.

Notes: Although the suggested media volume is 70 mL per plate, 50-80 mL per plate can be used. The three critical issues to consider when pouring PlusPlates are:

- 1) that the plates are leveled so that the agar media has the same thickness throughout the plate (use a leveled table or surface for plate pouring and do not pour in stacks of more than seven plates).
- 2) to ensure the absence of bubbles in the agar media (bubble should be popped using a sterile needle)
- 3) drying the plates for only one day and wrapping the plates in plastic bags to avoid failures in pinning yeast.

1.3. Sc –Trp and Sc –U –Trp rectangular PlusPlates. These plates will be used for growing the TF array collection (Sc –Trp) and to select diploid yeast after mating (Sc –U –Trp).

1.3.1. Dissolve the drop-out mix, the YNB, adenine hemisulfate, and the ammonium sulfate in 920 mL of water, and pH to 5.9 with NaOH 5M (approximately 1 mL per liter of media) (see **Table 4** for composition). Pour into a 2-L flask and add a stir bar.

1.3.2. In a second 2-L flask, add the agar to 950 mL of water (do not add a stir bar as it will cause the agar to boil over in the autoclave).

1.3.3. Autoclave for 40 min at 15 psi on liquid cycle.

1.3.4. Immediately pour the contents of the first flask, including the stir bar, into the agar containing flask. Add the glucose, mix well on a stir plate and cool to 55°C.

1.3.5. Add the leucine, histidine, and uracil (omit the uracil for the Sc –U –Trp plates).

1.3.6. Mix well on a stir plate and pour into rectangular PlusPlates (~70 mL per plate) using a peristaltic pump (5 mL per sec) and a 6 mm tubing. Dry for 1 day at room temperature, wrap the

plates in plastic bags, and store in the cold room for up to 3 months.

1.4. Sc –U –H –Trp + 3AT + X-gal rectangular PlusPlates. These plates will be used as readout plates for eY1H assays.

1.4.1. Dissolve the drop-out mix, the YNB, adenine hemisulfate, and the ammonium sulfate in 850 mL of water (see **Table 5** for composition). Do not pH. Pour into a 2-L flask and add a stir bar.

1.4.2. In a second 2-L flask, add the agar to 850 mL of water (do not add a stir bar as it will cause the agar to boil over in the autoclave).

1.4.3. Autoclave for 40 min at 15 psi on liquid cycle.

1.4.4. Prepare 10x BU salts (1L) by adding: 900 mL water, 70 g Na₂HPO₄·7H₂O, 34.5 g NaH₂PO₄·H₂O. Mix using a stir bar to dissolve powders and adjust pH to 7.0 using 5M NaOH. Add water to bring to 1L and autoclave.

1.4.5. Prepare the X-gal solution by adding 3.5g X-gal powder to a 50 mL plastic tube containing 42.5 mL dimethyl formamide. Add X-gal powder to dimethyl formamide to dissolve more easily (this takes 30 min). Keep stock solution in the dark (either use opaque 50 ml tube or cover in foil). Store at -20°C.

1.4.6. Immediately pour the contents of the first flask, including the stir bar, into the agar containing flask. Add the glucose and the 10X BU salts (see **Table 5** for composition), mix well on a stir plate and cool to 55°C.

1.4.7. Add the leucine, 3AT, and X-gal (see **Table 5** for composition).

1.4.8. Mix well on a stir plate and pour into the rectangular PlusPlates (~70 mL per plate) using a peristaltic pump (5 mL per sec) and a 6 mm tubing. Dry for 1 day at room temperature, wrap in plastic bags, and store in the cold room covered in aluminum foil (3AT and X-gal are light sensitive) for up to 1 month.

2. Spotting a TF array

2.1. Thawing the TF-prey array

2.1.1. Thaw the yeast glycerol stock plates with the TF-prey array on ice. TF-prey arrays can be generated as previously published ^{10,12,13,18}.

2.1.2. Resuspend the yeast using a 12-channel pipette within 1-3 min before the next step.

2.2. Spotting the yeast into Sc – Trp PlusPlates

2.2.1. In the HDA Rotor Robot, select multi-well 96 plates as source, PlusPlates 96 agar as target, and 96 long pin pads. Note: pin pads are not reusable and should be discarded.

2.2.2. Select the Replicate Many program to make two copies per 96-well plate. Do not use the recycle or revisit options to avoid back contamination of the frozen stocks.

2.2.3. Select the option to swirl up and down in the source to mix the yeast.

2.2.4. Bag the spotted array and incubate agar-side up at 30°C for 2-3 days.

2.3. Generating 384 colony arrays in Sc – Trp PlusPlates

2.3.1. In the HDA Rotor Robot, select the PlusPlate 96 agar plates as source, the PlusPlate 384 agar plate as target, and the 96 short pin pads.

2.3.2. Select the 1:4 Array program. In this way four 96 colony plates (each containing a different TF) will be consolidated into one 384 colony plate. Do not use the recycle or revisit options to avoid contamination between different plates.

2.3.3. Bag the plates and incubate the spotted 384-colony array agar-side up at 30°C for 2 days.

2.4. Generating 1,536 colony arrays in Sc – Trp PlusPlates

This will result in arrays containing four colonies for each TF-prey.

2.4.1. In the HDA Rotor Robot, select the PlusPlate 384 agar plates as source, the PlusPlate 1,536 agar plates as target, and 384 short pin pads.

2.4.2. Select the 1:4 assay single source program. The goal is to copy each colony into four colonies to obtain quadruplicates. Use the recycle and revisit options as it involves copying four times each colony.

2.4.3. Bag the plates and incubate the spotted 1536-colony array agar-side up at 30°C for 3 days.

2.5. Amplifying the 1,536 colony array in Sc – Trp PlusPlates

2.5.1. In the HDA Rotor Robot, select the PlusPlate 1,536 agar plates as source, the PlusPlate 1,536 agar plates as target, and 1,536 short pin pads.

2.5.2. Select the “Replicate many” program to replicate 3-4 copies. Use the recycle and revisit option, but throw out the pad when switching to a different plate of the array to avoid cross

contamination.

2.5.3. Bag the plates and incubate the spotted 1536-colony array agar-side up at 30°C for 3 days to use for mating steps (see below). After that, keep the plates at room temperature and copy again after 7 days for a new round of screening.

3. eY1H screen

3.1 Preparing DNA-bait strain lawns for mating

3.1.1. Spot the yeast DNA-bait strains on a Sc – U –H plate and grow for 3 days at 30°C.

3.1.2. Streak the yeast into a 15 cm Sc –U –H plate using a sterile toothpick, so that each plate fits 12-16 different strains. Incubate one day at 30°C.

3.1.3. Streak the yeast into a 15 cm Sc –U –H plate using a sterile toothpick, so that each plate fits 4 different strains. Incubate one day at 30°C.

3.1.4. Scrape the yeast using a sterile toothpick making sure not to scrape any agar, and add into a 1.5 tube with 500 µl of sterile water.

3.1.5. Add 10-15 sterile glass beads onto a YAPD rectangular PlusPlate. Add the yeast suspension onto the plate, and shake thoroughly in all directions for 1 min to ensure the yeast is spread through all the plate.

3.1.6. Invert the plate immediately and tap so that the beads go to the lid. Remove and recycle the beads.

3.1.7. Bag the plates and incubate agar-side down for 1-2 day at 30°C. Then proceed to the mating step.

3.2. Mating of yeast DNA-bait and TF array strains

3.2.1. Transfer the TF array to a YAPD rectangular PlusPlate with the HDA Rotor robot. Select the PlusPlate 1,536 agar as source and target, and the 1,536 short pin pad. Select the “Replicate Many” program. Each TF array plate can be used to transfer to 3-4 YAPD plates (depending on the number of plates in the array). The TF array plates used for mating must be 2-3 days old but not more as mating may be inefficient.

3.2.2. Transfer the lawn of a DNA-bait strain to the YAPD plates already containing the TF array with the HDA Rotor robot. Select the PlusPlate 1,536 agar as source and target, and the 1,536 short pin pad. Select the “Replicate Many” program. Use a Random off set in the Source with a radius of ~0.6 mm to avoid taking yeast from the same spot, and mix on target to facilitate contact

between yeast strains. Use the lawn containing the DNA-bait strains (3.1) as source, and the YAPD plates containing the TF array spotted in step 3.2.1 as target.

3.2.3. Bag the plates and incubate agar-side up at 30°C for 1 day.

3.3. Selection of diploid yeast

3.3.1. Transfer the mated yeast from the YAPD plates to Sc –U –Trp plates with the HDA Rotor robot. Select the PlusPlate 1,536 agar as source and target, and the 1,536 short pin pad. Select the “Replicate” program. Mix on source and on target.

3.3.2. Bag the plates and incubate agar-side up at 30°C for 2-3 days (longer incubation leads to high background reporter activity).

3.4. Transfer to readout plates

3.4.1. Transfer the diploid yeast from the Sc –U –Trp plates to the readout rectangular PlusPlates Sc –U –H –Trp + 5mM 3AT + 0.4 mM X-gal using the HDA Rotor robot. Select the PlusPlate 1,536 agar as source and target, and the 1,536 short pin pad. Select the “Replicate” program.

3.4.2. Bag the plates and incubate agar-side up at 30°C for up to 7 days.

3.5. Imaging of readout plates

For DNA-bait strains with high background reporter activity, take pictures on days 2, 3 and 4. Otherwise, take pictures at days 4 and 7. Positive interactions are identified by growth and blue color of the yeast colonies and can be manually determined, or it can be determined using image analysis software.

REPRESENTATIVE RESULTS:

Three main factors should be considered when analyzing results from eY1H assays: the background reporter activity of the DNA-bait strain, the strength of the reporter activity corresponding to TF-DNA interactions, and the number of positive colonies. The background reporter activity (*i.e.*, autoactivity) of the DNA-bait strain refers to the overall growth and color of the yeast colonies in the readout plate, even in the absence of a TF-prey. Ideally, non-autoactive strains show a background white or light brown color, with colonies for positive interactions being bigger and blue. Autoactive DNA-bait strains show yeast growth in media lacking histidine and a blue color in the presence of X-gal for all colonies in the plate, which is likely related to the binding of yeast transcriptional activators to the DNA-bait⁸. The strength of the reporter activity corresponding to the TF-DNA interactions detected (*i.e.*, the size of the

colony and intensity of blue) depends on many parameters such as the affinity, the TF expression level in yeast, the number of binding sites and distance to the yeast minimal promoters located upstream of the reporter genes, and the background reporter activity of the DNA-bait strain. For example, a weak interaction may be easily detected in a low background bait but may be difficult to detect in an autoactive or uneven background bait. It is also important to note that reporter activity levels in yeast do not necessarily correlate with the regulatory activity in human cells, as the chromatin structure, nucleosome positioning, and distance effects are different between yeast and human. Further, interactions in human are likely to be influenced by the binding of other TFs and cofactors or may be masked by functionally redundant TFs⁸. Finally, interactions are considered positive in eY1H assays when at least two of the four colonies show reporter expression above background levels. However, we have observed that ~90% of interactions identified result from all four colonies corresponding to a TF being positive^{10,12,19}. To illustrate the type of results that can be obtained using eY1H assays we screened the promoter regions (2 kb upstream of the transcription start sites) of the CCL15 and IL17F genes, against an array of 1,086 human TFs (**Figure 2**). The CCL15 promoter is an example of a non-autoactive DNA-bait where interactions, even weak ones, can be easily detected (**Figure 2A**). The IL17F promoter is an example of an autoactive DNA-bait with uneven background reporter activity, where some interactions can be detected while for several TFs it is uncertain whether the reporter activity is higher than background (**Figure 2B**).

Problems that can be encountered when performing eY1H assays

Although the screening eY1H protocol is straight-forward and robust, several problems can be encountered during the screen:

1) Colonies are too small and fail to transfer (**Figure 3A**): Although it is expected that some yeast expressing exogenous TFs may display slow growth given that yeast gene expression may be dysregulated, typically ~95% of TF-prey colonies display normal growth. If more than 10% of colonies fail to grow, the most frequent causes are problems with the media or with the yeast transfer. Suboptimal growth is frequently related to one of the media components losing activity (*e.g.*, uracil, histidine, or leucine), which can be solved by preparing fresh media with fresh stock solutions. Alternatively, this may also be related to a pinning offset that affects colony transfer. In this case, verify that the 1,536 pads pin the center of the yeast colonies in the source plates.

2) No yeast growth in a portion of the plate (**Figure 3B**): This issue is generally related with a failure in the mating step if the 1,536 pin pad fails to make contact with the yeast in the DNA-bait strain lawn, the TF array, or in the mating plate. In almost every case, this is due to uneven agar media level during plate pouring or due to excessive drying of the plate.

3) No interactions detected (**Figure 3C and D**): This issue is often related to either unintended inactivating mutations in the reporter genes, in particular LacZ (**Figure 3C**), or to high autoactivity that mask interactions (**Figure 3D**). To troubleshoot this problem, it is recommended to screen another independently obtained strain corresponding to the same DNA-bait.

4) The plate presents random blue spots (**Figure 3E**): This issue is often related to bacterial contamination. To solve this issue, streak the yeast to obtain individual colonies, and repeat the screen.

The above are the most frequent problems encountered when performing eY1H assays. Should other problems arise, preparing new media, confirming that appropriate settings for the HDA robot were used, and testing multiple strains per DNA-bait would likely solve most issues.

FIGURE AND TABLE LEGENDS:

Figure 1: Outline of eY1H assay screens. (A) Comparison between TF-centered and DNA-centered methods to identify protein-DNA interactions. (B) Schematics of eY1H assays. A DNA sequence of interest (promoter, enhancer, silencer, etc.) cloned upstream of the HIS3 and LacZ reporter genes is integrated into the yeast genome. The resulting DNA-bait strain is mated to a collection of yeast strains harboring TFs fused to the Gal4 activation domain (AD). Positive interactions are determined by the yeast's ability to grow without histidine and overcoming competitive inhibitor 3-AT, and turn blue in the presence of X-gal. (C) Pipeline for eY1H screens. A lawn of a yeast DNA-bait strain grown in a YAPD plate is mated in a YAPD plate to a 1,536 colony array expressing TFs fused to AD grown on a Sc – Trp plate. After one day, the yeast is transferred to a Sc –U –Trp to select for diploid yeast. After a 2-3 day incubation the yeast is transferred to a Sc –U –H –Trp + 3AT + X-gal plate (readout plate) to identify protein-DNA interactions. Each interaction is tested in quadruplicate.

Figure 2: Examples of eY1H readout plates. (A) Interactions involving the promoter of CCL15, a non-autoactive bait. Background reporter activity for this bait is low (reduced growth in the absence of histidine and absence of blue color for non-interacting TFs). (B) Interactions involving the promoter of IL17F, an autoactive bait. Background reporter activity for this bait is high (growth in the absence of histidine and background blue color throughout the plate) and uneven making it challenging to identify protein-DNA interactions. Strong, medium, and weak interactions are squared in red, orange, and yellow, respectively. The HGNC names of the interacting TFs are shown.

Figure 3: Problems in eY1H screens. (A) TF-prey array where multiple colonies failed to grow. (B) Readout plate where colonies in the lower left corner have failed to transfer. (C) Non-autoactive DNA-bait strain that does not display positive interactions. (D) Highly autoactive DNA-bait strain that does not display positive interactions. (E) Readout plate displaying multiple blue colonies due to contamination.

DISCUSSION:

The robotic eY1H mating screening approach described here greatly increases the throughput to identify the set of TFs that bind to a DNA region of interest, compared to previous library screening or arrayed screening approaches based on transformation. Further, the TF-DNA interactions detected by eY1H assays are highly reproducible as 90% of interactions detected are positive for all four colonies tested per TF, and 90% of interactions retest in an independent screen of the same yeast DNA-bait strain^{10,12,19}. More importantly, TF-DNA interactions detected by eY1H validate at a 40-70% rate when tested in human reporter assays^{12,20}, in primary human cells (unpublished results), and in *C. elegans* knockout animals¹¹. This is a similar validation rate to that observed for ChIP-seq data²¹.

Although, interactions identified by eY1H are highly reproducible when retesting the same yeast DNA-bait strain, testing different yeast strains for the same DNA-bait sometimes produce different, although overlapping, sets of TF-DNA interactions. This is usually due to differences in background reporter activity between strains. In addition, testing fragments of a DNA sequence result in the detection of more TF-DNA interactions than testing the full sequence, in particular when overlapping fragments are tested. This may be related with the assay being more efficient in identifying interactions that are close to the reporter minimal promoters, and because testing overlapping fragments reduces the chances that a binding site may be occluded by yeast nucleosomes. Thus, for small scale projects, it is recommended that overlapping 0.5-1 kb fragments of a regulatory region are tested and that two independent strains are screened for each DNA-bait sequence⁸.

There are several critical steps in the eY1H screening protocol to avoid some of the issues presented in **Figure 3**. First, although most media ingredients are stable for several months (except for 3AT and X-gal) a lack of proper colony growth likely indicates that at least one of the ingredients may have lost activity and should be replaced. Second, it is important to prepare the rectangular plates so that the agar is leveled and so that they do not dry for more than one day to avoid failure in pinning when using the robotic platform. Finally, it is key to use the robotic platform programs as indicated in the protocol (revisit, recycle, mixing, etc.) for the yeast to be transferred effectively, for mating to be efficient, and to avoid cross contamination between yeast clones.

The examples we selected to illustrate the use of eY1H screens correspond to human gene promoters. However, other regulatory regions can also be tested including enhancers and silencers. For example, we have used eY1H assays to evaluate TF binding to human developmental enhancers and to *C. elegans* first introns^{12,22}. In addition, given that interactions are tested in a pairwise manner, eY1H assays can be used to compare interactions between non-coding variants, and between TF coding sequence variants. For example, using eY1H assays we identified altered TF binding to 109 noncoding variants associated with different genetic diseases, and also differential interactions profiles for 58 TF missense mutations^{12,14}. Although, this protocol focuses on evaluating TF binding to human regulatory regions, DNA regions from other species can also be tested provided that a TF-prey is available or can be generated. Indeed, TF-prey arrays have been generated for *C. elegans*¹⁰, *Drosophila melanogaster*²³, *Mus musculus*²⁴, *Arabidopsis thaliana*^{25,26}, and *Zea mays*²⁷. Thus, with increasing available resources, eY1H assays may be applied additional systems.

Although eY1H assays have been instrumental to identify the repertoire of TFs that bind to different regulatory regions in human and other species, they are not free of caveats

8,11,12,19,20,25. One of the limitations is that interactions are tested in the milieu of the yeast nucleus and, although the DNA-baits are chromatinized, the chromatin structure in yeast may not reflect the chromatin structure in the species from where the DNA-bait originated and will not reflect cell type differences observed in vivo. Thus, interactions identified by eY1H assays must be validated in reporter or other functional assays. Of note, we and others have found TF-DNA interactions detected by eY1H validate at a 40-70% rate in functional assays^{11,12,20,23}. Another limitation of eY1H assays is that it cannot detect interactions involving TFs that require post-translational modifications absent in yeast to bind to DNA, TFs that are not properly folded in yeast when fused to the AD, and TFs that are missing from the array⁸. In addition, in the current format eY1H assays do not detect interactions involving heterodimeric TFs as each yeast colony in the TF array expresses a single TF-prey. Thus, further improvements in the assay will increase the breadth of TFs that can be tested and expand the capabilities of eY1H assays to identify novel TF-DNA interactions.

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DISCLOSURES:

The authors declare that they have no competing financial interests.

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