

# Directed yeast two hybrid screening for MEKK1 interaction partners

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## Method Article

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# Abstract

## Introduction

This protocol utilizes the Y190 yeast strain, that characteristically turns from white to red during growth due to *ade*- mutation. Baits were subcloned into the pAS1cyh2 vector, and Preys subcloned into pACT2 \ (5).

## Reagents

Y190 yeast strain YPAD medium 0.1M LiAc \ (Sigma) ssDNA \ (2.0 mg/ml) PEG 3350 \ (Sigma) Sterile H<sub>2</sub>O 42 °C waterbath SC minus plates \ (-LEU, -TRP) SC minus plates \ (-LEU, -HIS, -TRP), supplemented with 50 mM 3AT and 20 µg/ml X-gal

## Procedure

**\*\*Transformation\*\*** **\*\*1\*\*** Inoculate 10 ml of YPAD medium with a single Y190 colony and incubate overnight at 30 °C. 55 ml of fresh YPAD are then cultured overnight to an OD<sub>600</sub> of 0.3. Cells are then incubated by shaking at 200 RPM and 30 °C until they reach an OD<sub>600</sub> of 0.7. **\*\*2\*\*** Harvest cells at 2000 RPM for 5 min at room temperature. **\*\*3\*\*** Aspirate the media, wash cells in sterile water, and spin at 2000 RPM at 25 °C for 5 min. **\*\*4\*\*** Aspirate water and resuspend cells in 1 ml of 0.1 M LiAc and transfer suspension to a fresh microfuge tube. Pellet cells at 6000 RPM for 15 s and aspirate the supernatant. Next, resuspend the pellet in 500 µl of 100 mM LiAc. **\*\*5\*\*** Boil a 1 ml sample of ssDNA \ (2.0 mg/ml) and chill on ice. Mix the yeast cell suspension and transfer 50 µl samples into fresh tubes for the interactions to be tested. **\*\*6\*\*** Pellet cells at 6000 RPM and aspirate supernatant. **\*\*7\*\*** Add the following gently over the cell pellet: 240 µl of PEG 3350 \ (50% w/v), 36 µl of 1.0 M LiAc, 25 µg of ssDNA \ (2.0 mg/ml), and 50 µl sterile water containing plasmid DNA for the interaction screen \ (0.1 – 2.0 µg). Vortex mixture until cell pellet is completely mixed into suspension. **\*\*8\*\*** Incubate the suspension at 30 °C for 30 min. Transform the yeast by heat shock at 42 °C for 15 min, mix suspension carefully and pellet cells at 6000 RPM for 15 s. **\*\*9\*\*** Finally, resuspend yeast cells in 200 µl of sterile water for plating. **\*\*Plating and selection\*\*** **\*\*10\*\*** Plate yeast onto SC minus plates \ (-LEU, -TRP) and grow at 30 °C for 3 – 5 days. **\*\*11\*\*** Streak single colonies onto SC minus plates \ (-LEU, -HIS, -TRP) supplemented with 50 mM 3AT and 20 µg/ml X-gal. **\*\*12\*\*** Yeast are then grown for 3 – 5 days and scored for growth and color change.

## Timing

5 d minimum

## References

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