

# Evolving a Plant Pattern Recognition Receptor to Gain Resistance to a Pathogen-Derived Effector by Using a Novel Reverse Yeast Two-Hybrid System

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# Abstract

As the first layer of plant immunity, plants use pattern recognition receptors (PRRs) to perceive pathogens. To counteract PRRs, pathogens can secrete effectors that inhibit PRR's functions. In this study, we aim to enhance crop's disease resistance by engineering PRRs to elude the attacks from effectors, using a directed evolution approach. As a proof of concept, we worked on disrupting the interaction between effector AvrPtoB derived from destructive bacterial pathogen *Pseudomonas syringae*, and an ubiquitous PRR, CERK1. To achieve this, we developed TRUST-rY2H, a reverse yeast two-hybrid (rY2H) system that is **T**runcation-**R**esistant and **U**niversal, using **S**elf-cleavage peptide **T**echnology (TRUST). This system can eliminate the high rate of uninformative truncation mutations in rY2H, and can be universally applied to different genetic protein-protein interaction assays. Combining this novel rY2H system with a computational prediction method, we successfully identified mutations disrupting AvrPtoB-CERK1 interaction. An *in planta* screening system was also developed to identify *bona fide* AvrPtoB-resistant CERK1. Our study provides not only a novel strategy to generate transgene-free disease-resistant crops, but also a by far the best rY2H system to disrupt and study protein-protein interactions.

## Highlights

- Pioneered using directed evolution technology to create disease resistant crops
- Proposed a novel crop-protecting strategy by engineering PRRs to elude attacks by effectors
- Developed TRUST-rY2H, a novel rY2H system to disrupt protein-protein interactions.
- Identified mutations disrupting CERK1-AvrPtoB interaction using TRUST-rY2H and machine learning-based computational design.
- Designed a screening method to rapidly test the activity and AvrPtoB resistance of CERK1 variants in plants
- Developed a reverse assay for split-ubiquitin membrane protein yeast two-hybrid (MYTH) to disrupt the interaction between membrane proteins.
- Developed fluorescent reporters for conventional Y2H and MYTH to precisely measure the strength of protein-protein interactions.

# Introduction

Plant pathogens pose a global burden on agriculture, reducing harvest by up to 40% and causing more than \$220 billion in annual losses (American Phytopathological Society, 2023). Diseased crops not only lead to reduced yield and quality but ultimately impact the economy and environment (Martins et al., 2018; Savary et al., 2017). Currently, pesticides are used to protect crops, however, they negatively impact the environment and pose health risks to humans (Pathak et al., 2022). Therefore, developing pathogen-resistant crops remains a challenge that demands innovative solutions and research.

Plant innate immunity relies on pattern recognition receptors (PRRs) that detect pathogen-associated molecular patterns (PAMPs) (Boutrot & Zipfel, 2017; Zipfel, 2014). Phytopathogens evade this immune response by introducing virulence proteins called effectors into plant cells via e.g. the type III secretion system to promote pathogenesis (Macho & Zipfel, 2015). As a result, residues on the interaction interfaces find themselves under high selective pressure, facilitating the fixation of interaction-disrupting mutations to avoid being targeted by effectors (Bishop et al., 2000; Bishop et al., 2005; Shabab et al., 2008).

Among the plant pathogenic bacteria, *Pseudomonas syringae* ranks as the scientifically and economically most important bacterial pathogen worldwide (Mansfield et al., 2012). Strains of *P. syringae* are categorised into at least 50 pathovars and express a variety of effector proteins. One well-characterised effector of *P. syringae* is AvrPtoB. AvrPtoB is very important for *P. syringae*'s virulence, without which pathogen growth showed a 10-fold reduction (Gimenez-Ibanez et al., 2009). Besides *P. syringae*, members of the AvrPtoB protein family are also present in many other pathogenic strains of *Pseudomonas*, and at least two other genera of bacterial phytopathogens, *Xanthomonas* and *Erwinia* (Oguiza & Asensio, 2005).

One target of AvrPtoB is CERK1, which is responsible for sensing chitin and  $\beta$ -glucans from the fungal cell wall, as well as peptidoglycan (PGN) from the bacterial cell wall. CERK1 plays important roles in resistance to both fungal and bacterial pathogens, without which *P. syringae* growth showed a 10-fold increase (Gimenez-Ibanez et al., 2009). CERK1 consists of an extracellular domain with three tandem-LysM motifs, a transmembrane domain, and an intracellular Ser/Thr kinase domain (Yang et al., 2022). AvrPtoB can overcome CERK1-mediated resistance by inhibiting the kinase activity and targeting CERK1 for degradation, via its physical interaction with CERK1 (Gimenez-Ibanez et al., 2009).

The yeast two-hybrid (Y2H) system is a molecular technique used to detect protein-protein interactions, like those between effectors and PRRs. In traditional Y2H, the bait protein is fused to a DNA-binding domain (DBD), and the prey protein is fused to an activation domain (AD). Interaction of prey and bait activates the expression of reporter genes, leading to cell growth (Fields & Song, 1989). For reverse Y2H (rY2H), the interaction between wild-type proteins in rY2H is selected against, leading to no growth on selective medium containing 5-fluoroorotic acid (5-FOA). However, if certain mutations disrupt interaction, the cells will grow on the selective medium (Leanna & Hannink, 1996; Vidal et al., 1996). Unfortunately, truncation of proteins is very common in rY2H colonies, e.g. more than 97% of 5-FOA resistant colonies are expected to contain alleles coding truncation mutations (Gray et al., 2007).

In this study, we aim to evolve CERK1 to gain resistance to AvrPtoB. To disrupt CERK1-AvrPtoB interaction, we developed TRUST-rY2H, a novel reverse yeast two-hybrid system that is **T**runcation-**R**esistant and **U**niversal, using **S**elf-cleavage peptide **T**echnology (TRUST). By combining TRUST-rY2H and a machine learning-based computational design approach, we successfully identified several interaction-disrupting mutations. These mutations will be subjected to a rapid *in planta* screening developed in this study to identify *bona fide* AvrPtoB-resistant mutations. In future, such mutations will be introduced into the crop's genome using precise gene editing technologies. The resulting crop variety is *P. syringae*-resistant but non-transgenic, which is feasible in real-world agriculture.

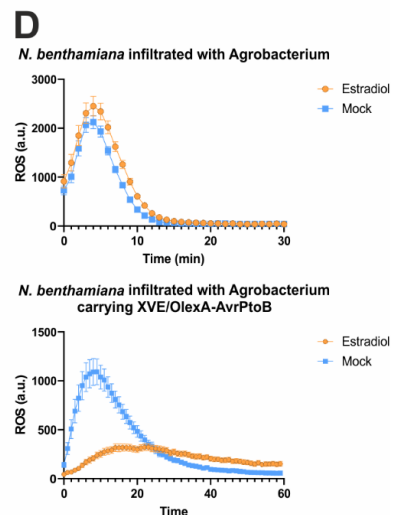
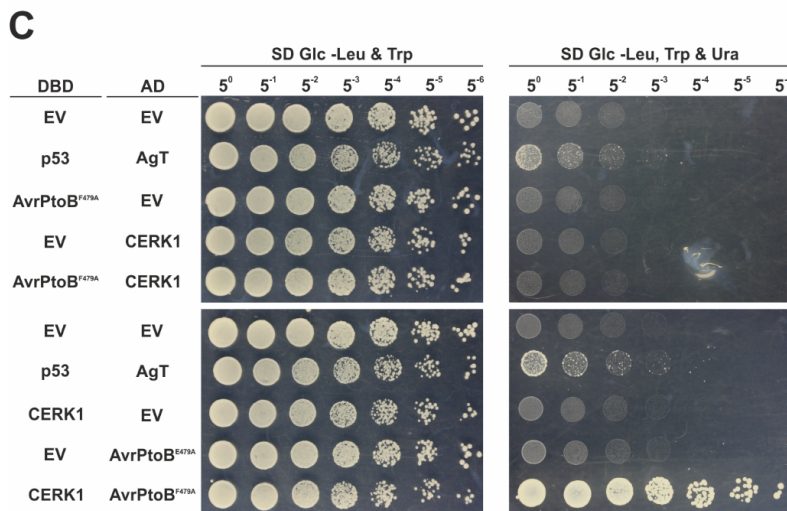
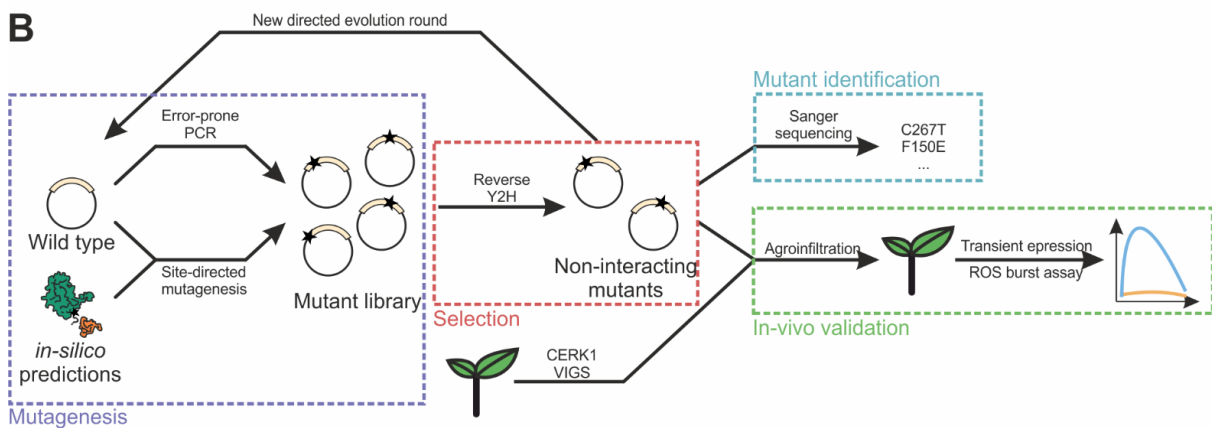
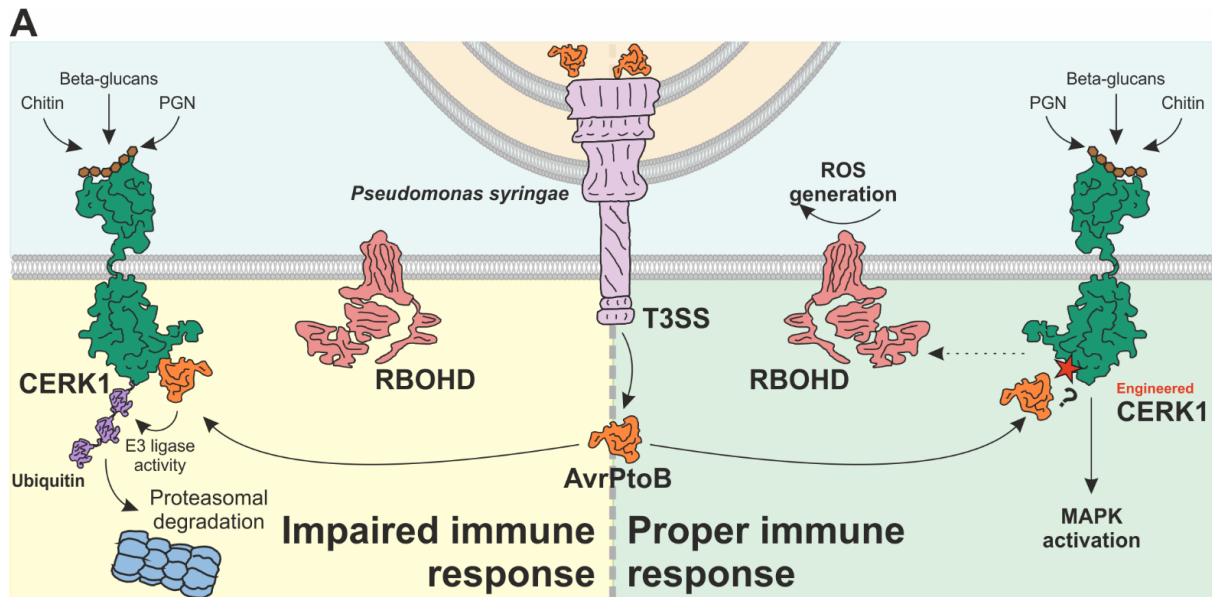
# Results

## AvrPtoB interacts with CERK1 and inhibits its function

CERK1 was validated as a direct target of AvrPtoB by Y2H assay (**Fig. 1C**). The interacting pairs CERK1 and AvrPtoB, as well as positive control p53 and large antigen T (AgT), were fused to DBD and AD respectively and expressed in yeast. On the selective plate, the combination of DBD-CERK1 with AD-AvrPtoB shows a considerably higher level of growth compared to all other combinations, demonstrating a strong interaction between CERK1 and AvrPtoB (**Figure 1C** lower part). Interestingly, when the DBD and AD are switched between the proteins (*i.e.* AD-CERK1 and DBD-AvrPtoB), the interaction was not detected (**Figure 1C** upper part), implying the DBD and AD could hinder this interaction if they are not fused favourably. This indicates a need for a Y2H system where DBD and AD can be freely switched between proteins of interest to avoid false negatives.

To confirm the impaired immune response caused by CERK1-AvrPtoB interaction, a ROS burst assay was carried out (**Fig. 1D**). When we transiently transformed the plants with an estradiol-inducible AvrPtoB DNA construct (XVE/OlexA-AvrPtoB) and the treated the plants with estradiol, the ROS burst was significantly reduced, indicating that AvrPtoB indeed impairs CERK1's functions.

Confirming that AvrPtoB interacts with CERK1 and inhibits CERK1's function, we aim to evolve CERK1 to elude the attack by AvrPtoB (**Fig 1A**). The workflow is shown in **Figure 1B**: (1) identifying mutations that disrupt CERK1-AvrPtoB interaction, via either reverse Y2H assay or computational prediction; (2) test obtained mutations *in planta* to identify *bona fide* AvrPtoB-resistant CERK1 variants.



**Figure 1. AvrPtoB interacts with CERK1, inhibiting its function and leading to impaired chitin-triggered immune response.**

A) Schematic representation of molecular events in a plant cell harbouring an AvrPtoB-susceptible (left part) and an AvrPtoB-resistant CERK1 (right part).

B) Workflow for the development of AvrPtoB-resistant CERK1 variants using directed evolution.

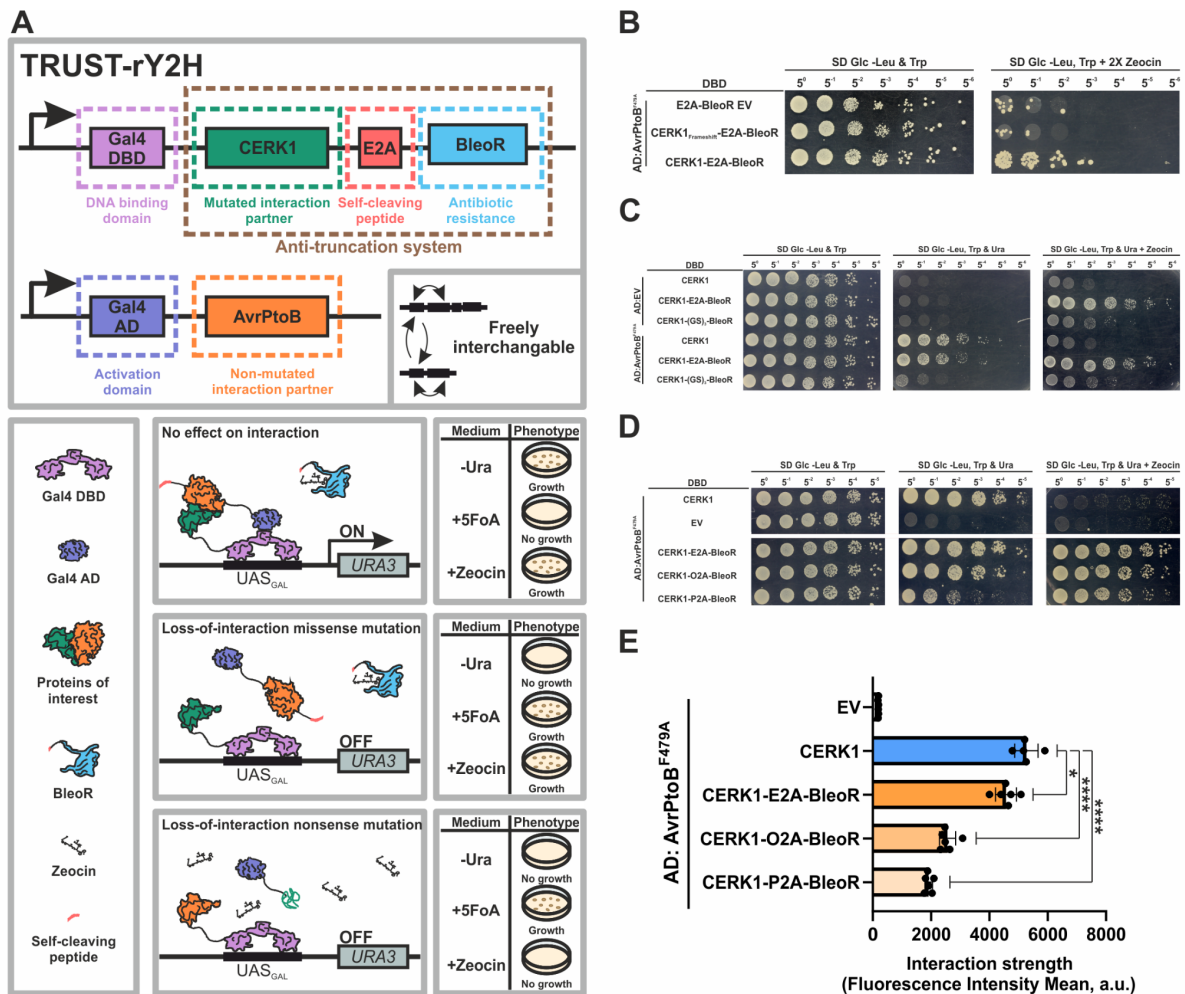
- C) CERK1 and AvrPtoB strongly interact in the Y2H assay, but only when fused to AD and DBD, respectively.
- D) Transient transformation and inducible expression of AvrPtoB attenuates the chitin-triggered ROS burst in *N. benthamiana*.

## Development of a novel reverse yeast two-hybrid system

Considering that the CERK1-AvrPtoB interaction is detectable only when CERK1 is fused to the DBD (**Fig. 1C**), and bulky C-terminal fusion might interfere with CERK1-AvrPtoB interaction, we realized that currently available reverse Y2H methods cannot achieve our goal to disrupt CERK1-AvrPtoB interaction (**Table 1**). To solve this problem, we developed a novel rY2H system, the TRUST-rY2H, which allows for the fusion of CERK1 to either AD or DBD, and selection against truncating mutations (using Zeocin-resistance gene BleoR), but with minimal interference on CERK1-AvrPtoB interaction (by sequestering CERK1 and BleoR using a 2A self-cleavage peptide) (**Fig. 2A**). The system is based on a reporter strain carrying *Ura3* gene and can thus be run in both forward and reverse directions using -Ura and +5-Fluoroorotic acid (5-FOA) media, respectively. Only full-length, in-frame translations of CERK1 allows cell growth on medium containing Zeocin, enabling truncation elimination (**Fig. 2A & B**).

We originally used the (GS)<sub>3</sub> linker for BleoR fusion, but this approach notably disrupted the CERK1-AvrPtoB interaction (**Fig. 2C** last row). We hypothesised that this disruption is likely due to the proximity of BleoR, a problem that could be solved by switching the (GS)<sub>3</sub> linker for a self-cleaving E2A peptide linker, physically separating BleoR from the CERK1-AvrPtoB complex. With the E2A peptide, we observed only a minimal interference on the interaction. Nonetheless, we tried to further optimise the system by evaluating two additional self-cleaving peptides, P2A and O2A. Both the plate (**Fig. 2D**) and flow cytometry Y2H assays have shown that the E2A peptide interferes with the CERK1-AvrPtoB interaction the least. We also used a fluorescent reporter strain to precisely quantify the effects of different 2A peptides on CERK1-AvrPtoB interaction (**Fig. 2E**). We got similar results as the plate assay, indicating that E2A is the best option.

In its current state, the system exhibits leaky behaviour, allowing even truncated mutants to grow in the presence of Zeocin (**Supplementary Fig. 2**). However, this behaviour is only present when the yeast are allowed to grow for >48h post-transformation and seems to disappear when they are plated immediately after the transformation, as would be the case in our directed evolution workflow (**Fig. 2B**).





**Table 1. Comparison of TRUST-rY2H to previous rY2H methods**

	System avoids truncations	Single step selection	Protein of interest can be fused to both the activating domain and DNA binding domain	Compatible with membrane systems (split ubiquitin system)
Original rY2H	-	-	+	-
rY2H (Vidal et al.)	+	-	+	-
rY2H (Leanna et al.)	-	-	+	-
rDY2H (Vincent et al.)	+	+	-	-
One-plus Two Hybrid system (Kim et al.)	+	-	-	-
TRUST-rY2H	+	+	+	+

**Table 2. Summary of CERK1 variants obtained from the reverse yeast two-hybrid assays.** Each line shows mutations found in a single colony. Notation: R415G means R to G substitution in 415th amino acid. Star denotes a stop codon. Mutations on the predicted interaction interface are underlined.

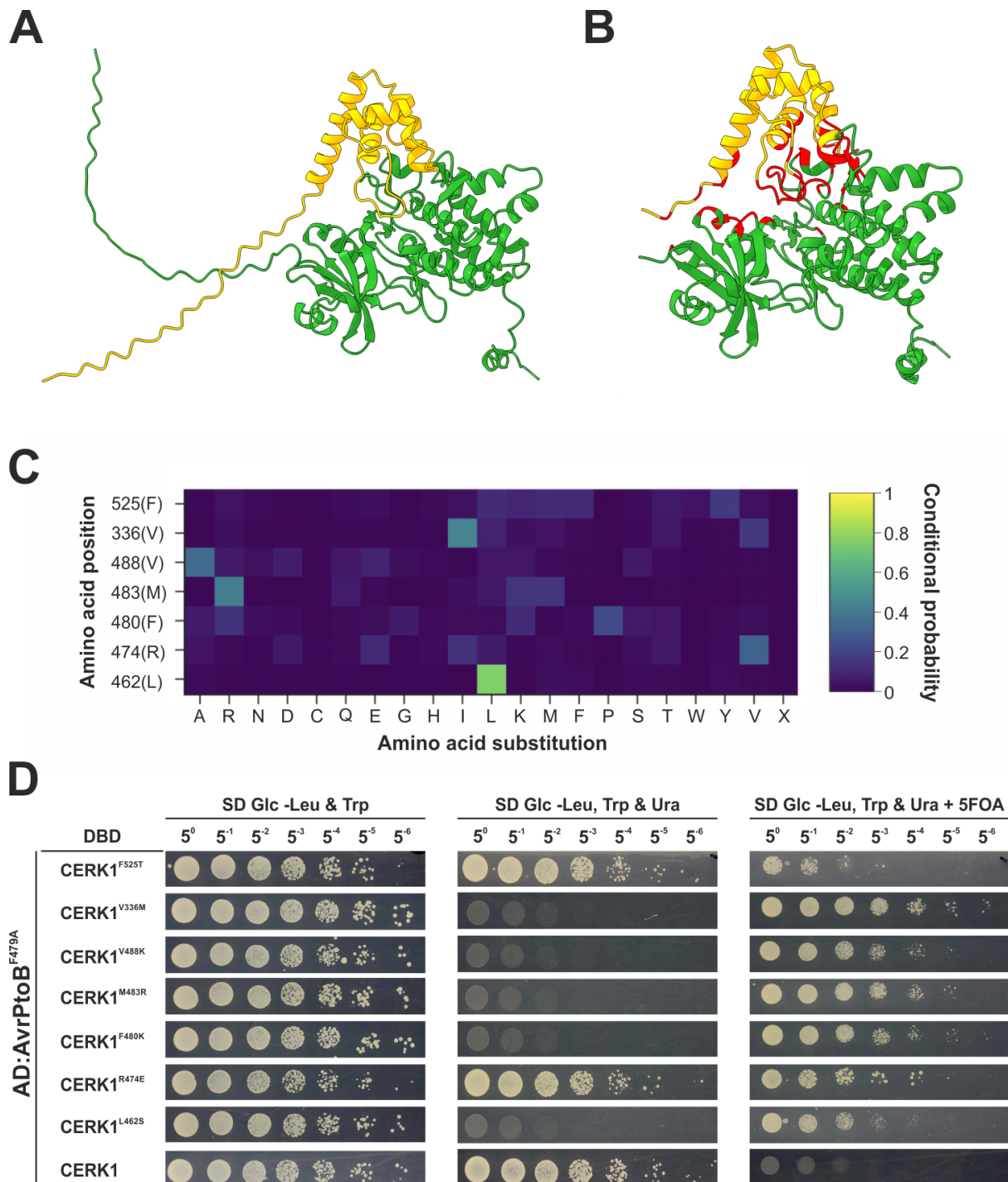
Classic rY2H without anti-truncation	TRUST-rY2H
S510P L398P G502E <u>S444F</u> <u>G461R</u> F267L, L363Q G569R C381W	R415G, A521T Y337C F533S F359S K349R
W412* L388*	K495R, K345* Q451* E575E, K364*

## Disruption of CERK1-AvrPtoB interaction using TRUST-rY2H

We performed mutagenesis on CERK1 using error-prone PCR, and subjected the mutagenesis library to TRUST-rY2H to select for mutations that disrupt CERK1-AvrPtoB interaction. For comparison, we also run a classic rY2H in parallel, which is without an anti-truncation system. Ten colonies from each screen were sequenced to identify mutations (**Table 2**). Three out of ten colonies from TRUST-rY2H were false positives with nonsense mutations (stop codons), while the classic rY2H screen produced two such false positives. Interestingly, only two mutations (S444F and G461R) were located in or close to the predicted interaction interface. This can be due to the fact that the actual binding interface might be larger than predicted, or some mutations might disrupt the global confirmation of CERK1.

### *In silico* mutation discovery based on machine learning

In parallel with the reverse Y2H assay, the computational approach was also used to identify mutations disrupting the CERK1-AvrPtoB interaction. To achieve this, we first predicted the structure of the CERK1-AvrPtoB complex. The first round of prediction was done using AlphaFold-Multimer (Evans et al., 2022; Jumper et al., 2021), resulting in an unsatisfactory model confidence of 0.63 (pTM = 0.75 and ipTM = 0.6) (Yin et al., 2022). As the accuracy of the binding complex is pivotal in the discovery of interaction-disrupting mutations, we explored another prediction tools, namely ColabFold (Mirdita et al., 2022). Additionally, the preliminary model showed that both CERK1 and AvrPtoB contain large, intrinsically disordered regions that are not involved in the interaction but are poorly predicted and restrict model confidence (**Fig. 3A**). The second prediction using the processed input sequences and ColabFold resulted in improved model confidence of 0.74 (pTM=0.83, ipTM=0.72).



**Figure 3. Computational prediction-based site-directed mutagenesis successfully disrupts CERK1-AvrPtoB interaction.**

A) Prediction of AvrPtoB-CERK1 binding complex from AF2. CERK1 is coloured green, and the AvrPtoB is yellow. Both are attached with a large disordered region.

B) Prediction of cropped AvrPtoB-CERK1 binding complex from Colabfold. The CERK1 is coloured green, and the AvrPtoB is yellow. The binding interface is labelled in red.

C) Conditional probabilities of 20 amino acids at selected positions calculated by Protein MPNN.

D) Experimental validation of binding interactions between CERK1-mutants and AvrPtoB.

In PyMol we roughly determined the binding interface of the best-predicted model by finding all interactions between AvrptoB and CERK1 within 3 Å (**Fig. 3B**). We further narrowed the scope of our mutation search by excluding potential ATP-binding and substrate-binding domains (Kobe & Kemp, 2003). From the CERK1 binding interface, we selected residues with the highest number of interaction partners for the site-directed mutagenesis. When choosing the specific substitutions, we opted for those most disruptive for the interaction between CERK1 and AvrPtoB, while still preserving the global protein structure, and in turn the kinase function of CERK1. Therefore, we calculated the conditional probabilities for all 20 amino acid substitutions at the seven most interactive residues using Protein MPNN (**Fig. 3C**) (Dauparas et al., 2022). Taking the chemical properties of amino acids into account, we settled on the following substitutions: L462S, F480K, V488K, F525T, R474E, M483R, and V336M.

To validate the efficacy of the in silico mutations in disrupting the interaction between CERK1 and AvrPtoB, site-directed mutagenesis was employed. We expressed and analysed 7 CERK1 variants in yeast cells, using the wild-type strain as a control (**Fig. 3D**). As expected, yeast with wild-type CERK1 demonstrated growth on SD gluc -LWU medium, implying an interaction with AvrPtoB. Out of the seven computationally predicted mutants, five (L462S, M483R, F480K, V488K, V336M) successfully disrupted this interaction as evidenced by a lack of growth on the -LWU medium.

## *In planta* screening of AvrPtoB-resistant CERK1 variants that retain kinase activity via transient expression in *N. benthamiana*

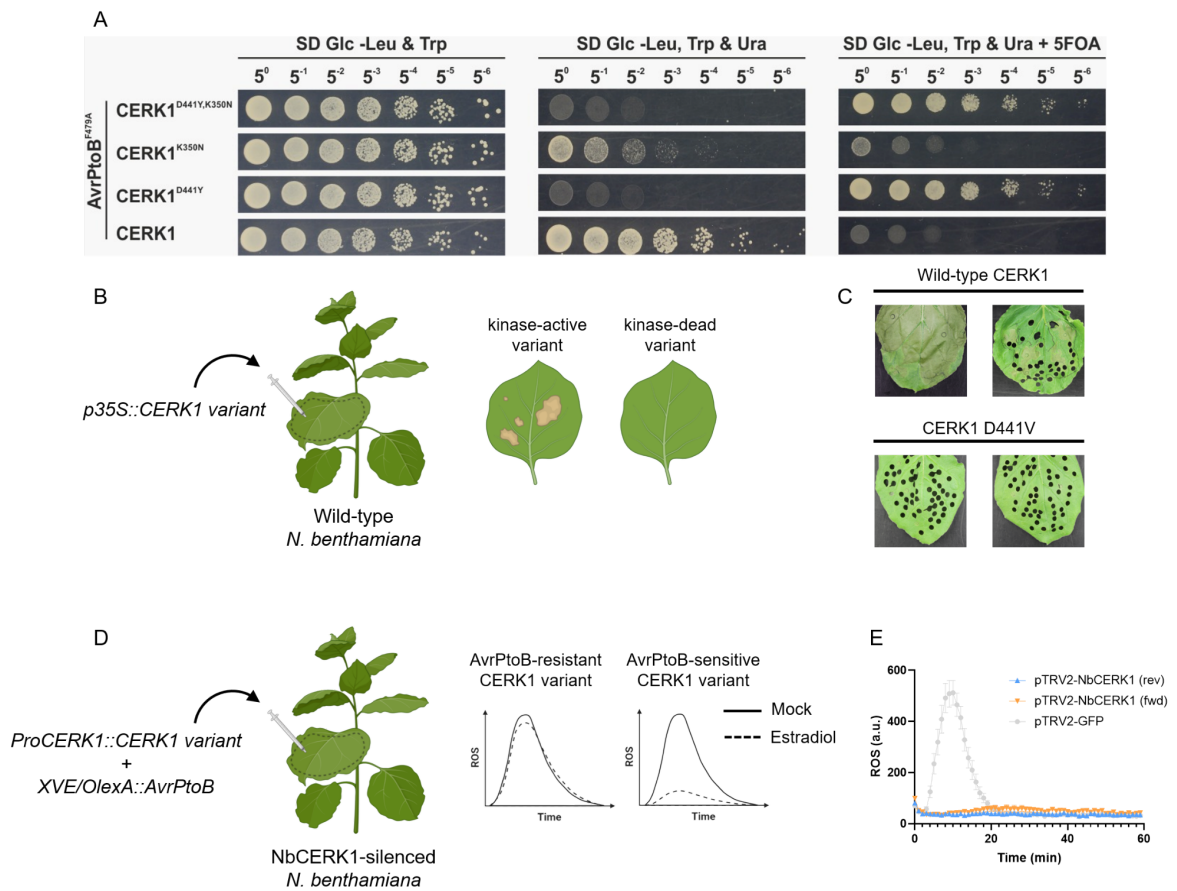
Loss of interaction with AvrPtoB can be accompanied by loss of kinase activity. This concern arises from the site-directed mutagenesis analysis of AvrPtoB-BAK1 interaction, where many structure-informed BAK1 mutants that lost interaction with AvrPtoB also lost their kinase activities (Cheng et al., 2011).

For CERK1, we found that two frequently used kinase-dead mutants, D441V and K350N, either lost or exhibited a reduced interaction with AvrPtoB, respectively (**Fig. 4A**), suggesting that CERK1's kinase activity and its interaction with AvrPtoB might be also closely correlated. Consequently, an additional round of screening for CERK1 variants that retain kinase activity is necessary.

To perform the screening in a rapid and high-throughput manner, we overexpressed CERK1 in *N. benthamiana* using the strong, constitutive 35S promoter, and checked for a cell death phenotype (**Fig. 4B**). This is based on the fact that cell death triggered by overexpression of CERK1 is dependent on CERK1's kinase activity (Pietraszewska-Bogiel et al., 2013; Suzuki et al., 2018), i.e., only kinase-active CERK1 variants can trigger cell death. Indeed, expression of wild-type CERK1 and not the D441V kinase-dead mutant caused cell death in *N. benthamiana* (**Fig. 4C**). This strategy is very straightforward and does not need a *cerk1* mutant of *N. benthamiana*, which takes a considerable amount of time to construct. However, the cell death phenotype is not solely dependent on the kinase activity, but also the juxtamembrane domain. As such, the lack of cell death does not mean the variant is kinase-dead, so this strategy might generate some false negatives (Zhou et al., 2020).

Active CERK1 variants will be subjected to a second round of *in planta* screening designed to screen for *bona fide* AvrPtoB-resistant CERK1 variants (**Fig. 4D**). In this round of screening, virus-induced gene silencing (VIGS) is used to silence the native CERK1 of *N. benthamiana* (NbCERK1) (Gimenez-Ibanez et al., 2009). The AtCERK1 variants are expressed using the native promoter, ensuring that AtCERK1 expression remains at a low level, insufficient to trigger cell death, which allows for the measurement of a ROS burst in response to chitin. At the same time, AvrPtoB is also expressed in an estradiol-inducible manner. By comparing the ROS bursts of mock and estradiol treatment groups, we can determine whether the CERK1 variant is resistant to AvrPtoB.

Due to time constraints, we were unable to perform *in planta* screening of variants obtained from the rY2H and computational prediction. However, our preliminary test using wild-type CERK1 and D441V kinase-dead mutant proved our method of *in planta* screening to be adequate for testing kinase-active variants (**Fig. 4C**), and the successful silencing of NbCERK1 created the chassis to screen AvrPtoB-resistant CERK1 variants (**Fig. 4E**)



**Figure 4. Design of *in planta* screening assays for kinase-active and *bona fide* AvrPtoB-resistant CERK1 mutants.**

(A) Kinase-dead CERK1 mutants lost interaction with AvrPtoB suggesting CERK1's kinase activity and capability to interact with AvrPtoB are closely correlated.

(B) The assay used to screen for CERK1 variants that retain kinase activity, which is based on overexpression of CERK1 variants and observation of cell death phenotype.

(C) Expression of wild-type CERK1 and not the D441V kinase-dead mutant caused cell death in *N. benthamiana*.

(D) The assay used to screen for AvrPtoB-resistant CERK1 variants *in planta*.

(E) Successful silencing of native CERK1 of *N. benthamiana*.

# Development of a reverse split ubiquitin-based membrane protein yeast two-hybrid system using E2A-BleoR anti-truncation system

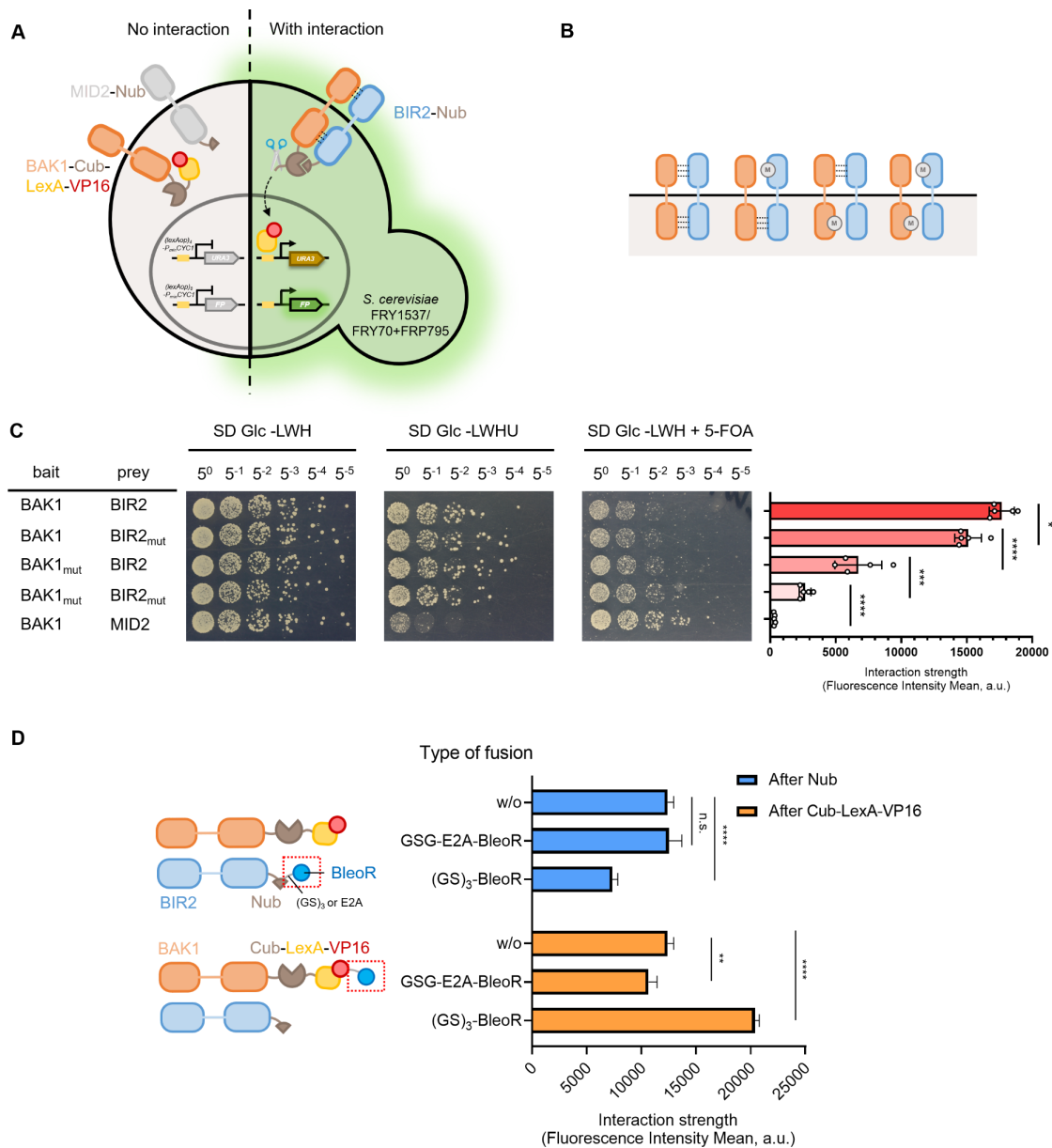
Apart from the conventional yeast two-hybrid, diverse genetic methods have been developed to test protein-protein interactions (Stynen et al., 2012). For example, the split ubiquitin-based membrane protein yeast two-hybrid (MYTH) system, which is designed to study the interaction between membrane proteins (Iyer et al., 2005). Except for the conventional Y2H, no reverse system was successfully developed for other genetic PPI assays, with there being only a failed attempt (Sahin, 2019). In terms of truncation elimination, some strategies are limited to the conventional Y2H (Kim et al., 2007; Vincent et al., 2020), some can potentially be used for other genetic PPI assays like MYTH but with the risk of impairing the interaction detection. Therefore, we strived to develop a reverse MYTH system using E2A-BleoR to eliminate truncation mutations.

To enable the reverse selection, a *Ura3* reporter gene, which is not found in common MYTH strains, is needed. Therefore we repurposed strain FRY1537 from Ottoz et al., which carries a *Ura3* gene driven by a synthetic promoter that can be activated by the LexA-VP16 synthetic transcription factor, as a reporter strain for our rMYTH system. A fluorescent reporter strain FRY70+FRP795 (similar to FRY482 from Ottoz et al., 2014) was also constructed to precisely quantify the interaction on a single-cell level using flow cytometry (**Fig. 5A**).

As a proof-of-concept, we used two well-studied single-pass membrane proteins in the plant immune systems, BAK1 and BIR2, to test our rMYTH system and intracellular domains (**Fig. 5A**). BAK1 and BIR2 have been reported to interact with each other through both extracellular and intracellular domains (Ma et al., 2017; Halter et al., 2014). As the residues on the interaction surfaces were previously identified, we generated mutations on some of these residues to disrupt either extracellular or intracellular interaction or both of them (**Fig. 5B**). This resulted in BAK1-BIR2 pairs with different levels of interaction strength, which were quantified by the fluorescence readout (**Fig. 5C** bar plot). As shown in the plate assay results (**Fig. 5C**), there was a positive correlation between the interaction strength and the cell growth on the plate without uracil, while the 5-FOA plate showed an opposite trend. These results suggest the success of repurposing FRY1537 as a reporter strain for rMYTH.

We then tested whether the E2A-BleoR anti-truncation strategy can be also applied to rMYTH (**Figure 5D**). E2A-BleoR and (GS)<sub>3</sub>-BleoR were fused after Cub-LexA-VP16 for the bait construct and after Nub for the prey construct, respectively. The fusions of E2A-BleoR after Nub and Cub-LexA-VP16 had no effect and a neglectable negative effect on BAK1-BIR2 interaction, respectively. For (GS)<sub>3</sub>-BleoR, a 50% decrease of interaction strength was observed when it was fused after the Nub, suggesting that the covalent fusion severely interfered with the detection of interaction. When (GS)<sub>3</sub>-BleoR was fused after the Cub-LexA-VP16, an increase in the interaction strength was observed, which implies that (GS)<sub>3</sub>-BleoR can cause false positive results, perhaps because of the homodimerization of BleoR (Dumas et al., 1994).

The rMYTH system can be potentially used to evolve BAK1, the co-receptor shared by many pattern recognition receptors in plants (Yasuda et al., 2017), to become resistant to plasma membrane-localised *P. syringae* effector AvrPto. Only MYTH, but not the conventional Y2H system is capable of detecting AvrPto-BAK1 interaction in yeast (Halter et al., 2014).



**Figure 5. Development of TRUST-rMYTH, a reverse split ubiquitin-based membrane protein yeast two-hybrid system with E2A-BleoR truncation elimination module.**

(A) Schematic diagram of the rMYTH design used for selection of loss-of-interaction mutations on membrane proteins. (B) BAK1 and BIR2 with interaction-disrupting mutations on extracellular or (and) intracellular domains are used to test the rMYTH system. (C) Cell growths on -LWHU and -LWH + 5-FOA plates were positively and negatively correlated with the interaction strength measured by flow cytometry using the fluorescent reporter strain. (D) E2A-BleoR can be fused after either Nub or Cub-LexA-VP16 without or only mildly affecting BAK1-BIR2 interaction. Differences between sample means in the flow-cytometry data were assessed using one-way ANOVA with Dunnett's post-hoc test; <0.01 (\*), <0.0001(\*\*\*\*)



# Discussion

Compared to existing rY2H systems, our TRUST-rY2H is more robust as seen through the advantages covered in Table 1. Although carefully designed (removing the start codon of BleoR, out-of-frame empty vector), TRUST-rY2H didn't manage to eliminate all the truncation mutations (Table 2). This is perhaps due to the leaky expression of BleoR, which might be caused by multiple methionine codons at the C-terminus of CERK1 and upstream sequences with IRES property, or non-canonical translation initiation (Monteuuis et al., 2019). To overcome this, we plan to introduce D25A mutation to BleoR to compromise its activity by ~ 50% (Dumas et al., 1994), which has been proven useful to counteract the leaky expression (Tominaga et al., 2021).

Surprisingly, among the ten sequenced colonies from the rY2H without anti-truncation selection, only two carried truncation mutations. This truncation rate (20%) is far lower than reported (>97%) (Barr et al., 2004, Gray et al., 2007). One possibility is that some of the hits are false positives. This can be because of mutations disrupting DBD, promoter, and *Ura3* gene, which can also lead to a 5-FOA-resistant phenotype. Therefore, we need to confirm these mutations by performing a round of forward Y2H or fully sequence the plasmids extracted from yeast. Another possibility is that CERK1-AvrPtoB is sensitive to mutations, i.e., among all the potential interaction-disruption mutations, there are many more non-truncation mutations than truncation mutations. The result could also be due to stochastic colony picking and requires more colonies to achieve statistical robustness.

We also experimented with a mutation discovery method entirely based on machine learning (Bryant et al., 2022; Homma et al., 2023). With the predicted models, we were able to pinpoint the binding interface and design point mutations that prohibit binding interactions, which were experimentally validated. Based on the experimental results, five out of the seven predicted mutations disrupted the CERK1-AvrPtoB interaction, suggesting the success of our computational approach.

Although testing variants in *N. benthamiana* by ROS burst assay is simple and fast, it is not sufficient to prove that CERK1 mutants retain their full activity. Therefore, stable expression lines should be constructed by complementing *A. thaliana cerk1* mutant or *fec* (*fls2 efr cerk1*) mutant with obtained CERK1 variants, and multiple immune outputs should be evaluated, for example, activation of mitogen-activated protein kinases (MAPKs) (Yamada et al., 2016), expression of the immune marker genes, and callose deposition. (Gimenez-Ibanez et al., 2009). Ultimately, disease resistance should be quantified by inoculating *Pseudomonas syringae* to aforementioned CERK1 variants complementation lines and measuring pathogen growth.

Our strategy can be used to engineer not only the resistance (R) genes, but also the susceptibility (S) genes, which are exploited by pathogens. Since S genes are often involved in physiological processes of plants, complete knock-out or disruption of a S gene sometimes leads to fitness cost (Koseoglou et al., 2022). Therefore, using rY2H technology to design gene variants resistant to effectors meanwhile retaining its physiological functions can be another potential application scenario of our strategy.

# Materials and methods

## Yeast two-hybrid

For the Y2H plate assay 4-5 medium-sized yeast colonies were picked and thoroughly resuspended in 50  $\mu\text{L}$  of Milli-Q  $\text{H}_2\text{O}$ . The obtained yeast suspensions were adjusted to an  $\text{OD}_{600}$  value of 0.2, serially diluted by a factor of 5, and plated on the appropriate selection medium. The base medium was SD glucose with -Leu & Trp as control, -Leu, Trp & Ura to select for interactions, -Leu, Trp & 5-Fluoroorotic acid ( $\gamma = 1 \text{ mg mL}^{-1}$ ) to select for non-interactions. Zeocin ( $\gamma = 1 \text{ mg mL}^{-1}$ ) was added to the plates to select against truncated and out-of-frame mutations. The MYTH system is based on the same principle, using media that are additionally -His.

## Flow cytometry

For each group, 6 colonies are picked, resuspended in 1.5 mL yeast SD medium and cultured overnight in 96-deep well plates (Corning®) sealed with 3M™ Micropore™ Surgical Tape. For measurement using flow cytometry, the overnight cultures are diluted 5-fold using PBS (Phosphate-buffered saline) buffer and loaded in the flow cytometer. The measurement of yeast fluorescence using flow cytometry followed the protocol described in Ottoz et al., 2016. The data analysis was performed using FlowJo software, using a one-way ANOVA with Dunnett's posthoc test to assess differences between sample fluorescence means.

## Agrobacterium-mediated transient expression in *Nicotiana benthamiana*

*N. benthamiana* was co-infiltrated with agrobacterium containing a plasmid coding for AvrPtoB, and a plasmid containing P19. 100 ng of plasmid was added to electrocompetent agrobacterium GV3101, which were electroporated at 1.4 kV. 2 h of recovery in 500 ml LB medium was carried out at 28 C. The bacteria were then plated and grown overnight in selective medium. The following day, a liquid culture was started using 5-6 colonies from the plate. The next morning, the bacteria were centrifuged and resuspended in the infiltration buffer. The  $\text{OD}_{600}$  of the cultures was adjusted to 0.5. P19 culture was added to each trial culture at a ratio of 1:5, with a final  $\text{OD}_{600}$  of 0.5. The mixed cultures were then incubated for 1-3 hours at room temperature while shaking. *N. benthamiana* leaves were then infiltrated with the culture mixture using a syringe. The plant was left at room temperature overnight before the leaves were brushed with estradiol (100 mM in DMSO, diluted with water to 50 mM) using cotton swabs to induce expression of AvrPtoB. Negative controls were incubated in 10-50 mM DMSO to mimic experiment groups.

## ROS burst assay

Leaf discs were harvested from *Nicotiana benthamiana* using a 4-mm diameter biopsy punch. To decrease the background signal, leaf discs were incubated overnight on 100  $\mu$ L of distilled water in white 96-well-plates (Greiner Bio-One) wrapped with aluminum foil. Chitin stock solution (10 mg/mL) was diluted for 100 times in to the ROS assay solution (100  $\mu$ M L0-12 (FUJIFILM), 20  $\mu$ g mL<sup>-1</sup> horseradish peroxidase (HRP)). Prior to ROS measurement, the water was removed and replaced with ROS assay solution with or without the addition of elicitors. Immediately following the addition of the assay solution, light emission was measured from the plate using Tecan SPARK microplate reader.

## Binding complex prediction with AlphaFold2

The CERK1-AvrPtoB binding complex was first predicted using AlfaFold-Multimer V2.3.1 using default parameters with cloned protein sequences as input. Based on the obtained prediction, intrinsically disordered regions in CERK1 and AvrPtoB which don't participate in the binding complex were removed from the input sequences. A second round of predictions was performed using ColabFold V1.5.1 with the new input sequences, recycling number set at 15, and using templates. The predicted models were evaluated by the model-predicted TM scores using the following formula:

$$\text{model confidence} = 0.8 \text{ ipTM} + 0.2 \text{ pTM}$$

## Analysis of binding interface

The binding interface was determined in PyMol by predicting all possible interactions within 3 Å. The seven most interactive CERK1 residues in the binding interface were selected for site-directed mutagenesis. The selection specifically avoided the residues close to the potential ATP-binding as well as the substrate-binding domain to preserve the CERK1 kinase function. Additionally, glycine and proline were avoided to reduce potentially negative effects on protein stability and folding.

## Discovery of structure-preserved mutations

The final mutations were discovered with the help of a protein message-passing neural network (ProteinMPNN). Which takes the most confidence predicted CERK1 structure as input and enables the sequence redesign based on the given structure. The redesigned amino acid sequence is expected to refold into the given backbone. In our work, the redesign only happened at the selected residue positions. The output returns the conditional probabilities of all 20 amino acids at given positions. The higher conditional probability has an amino acid at a given position, it is more favourable in the defined structure. Considering both chemical properties and conditional probabilities, we have chosen the rational mutations, which could prohibit interactions with AvrptoB and preserve the initial CERK1 structure.

## Error-prone PCR

GoTaq polymerase (Promega) was used according to the corresponding protocol. Mutation rates were calculated using the [ThermoFisher fidelity calculator](#). In order to have a single point mutation predicted across the CERK1 sequence, regular conditions sufficed. The annealing temperatures were adjusted to the primers and the thermocycler was run for 35 cycles. The PCR product was purified as normal.

## Virus-induced gene silencing (VIGS)

Electrocompetent agrobacterium were transformed as described above with either pTRV1 or pTRV2-CERK1. The bacteria were then recovered and cultured overnight as described above. The following day, the cultures were adjusted to an OD600 of 2.0 and pTRV1 and pTRV2-CERK1 cultures were mixed in a 1:1 ratio. The mixture was then incubated for 1-3 h at room temperature while shaking. The true leaves of 1-week-old *N. benthamiana* were then infiltrated with the culture mixture using a syringe. The infiltrated plants were grown for 3-4 weeks at (greenhouse conditions). A ROS burst assay can be performed to validate the gene silencing.

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