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RESEARCH ARTICLE

Genome editing in *Kluyveromyces* and *Ogataea* yeasts using a broad-host-range Cas9/gRNA co-expression plasmid

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ABSTRACT

While CRISPR-Cas9-mediated genome editing has transformed yeast research, current plasmids and cassettes for Cas9 and guide-RNA expression are species specific. CRISPR tools that function in multiple yeast species could contribute to the intensifying research on non-conventional yeasts. A plasmid carrying a pangenomic origin of replication and two constitutive expression cassettes for Cas9 and ribozyme-flanked gRNAs was constructed. Its functionality was tested by analyzing inactivation of the ADE2 gene in four yeast species. In two *Kluyveromyces* species, near-perfect targeting (\geq 96%) and homologous repair (HR) were observed in at least 24% of transformants. In two *Ogataea* species, Ade⁻ mutants were not observed directly after transformation, but prolonged incubation of transformed cells resulted in targeting efficiencies of 9% to 63% mediated by non-homologous end joining (NHEJ). In an *Ogataea parapolymorpha ku*80 mutant, deletion of *OpADE2* mediated by HR was achieved, albeit at low efficiencies (<1%). Furthermore the expression of a dual polycistronic gRNA array enabled simultaneous interruption of *OpADE2* and *OpYNR1* demonstrating flexibility of ribozyme-flanked gRNA design for multiplexing. While prevalence of NHEJ prevented HR-mediated editing in *Ogataea*, such targeted editing was possible in *Kluyveromyces*. This broad-host-range CRISPR/gRNA system may contribute to exploration of Cas9-mediated genome editing in other *Saccharomycotina* yeasts.

Keywords: CRISPR/Cas9; ribozymes; Kluyveromyces lactis; Kluyveromyces marxianus; Ogataea polymorpha; Hansenula polymorpha

INTRODUCTION

Design and construction of efficient yeast cell factories for industrial production of fuels and a wide range of chemicals

is among the key developments in microbial biotechnology in the last 20 years (Nielsen et al. 2013; Chen et al. 2015; Wang, Huang and Nielsen 2017). Saccharomyces cerevisiae is currently the most intensively used yeast species for production of low-

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molecular-weight products such as alcohols, organic acids and isoprenoids (Mattanovich, Sauer and Gasser 2014). Its tolerance to high sugar concentrations and low pH, its overall robustness, its ability to grow anaerobically and especially its tractability to genetic manipulation make S. cerevisiae a convenient chassis for various biotechnological purposes (Kavscek et al. 2015; Li and Borodina 2015). Furthermore, availability of high-quality genome sequences and well-developed genetic tools (Salazar et al. 2017; Stovicek, Holkenbrink and Borodina 2017) facilitate tailoring of S. cerevisiae to specific industrial applications, either by improving existing traits or by expressing heterologous enzymes and pathways. However, its narrow temperature spectrum, limited substrate range and strong tendency toward alcoholic fermentation under aerobic conditions have stimulated interest in studying non-Saccharomyces species, often referred to as 'non-conventional yeasts', that exhibit attractive features for further improving sustainability and economic viability of biobased production (Johnson 2013).

Kluyveromyces lactis and K. marxianus are non-conventional yeasts that are generally regarded as safe (GRAS), can utilize lactose as a carbon source and have been applied for bioethanol production from cheese whey, a by-product of the dairy industry (Siso 1996). Kluyveromyces lactis has an excellent capacity for protein secretion, which has been exploited for production of several heterologous proteins (Spohner et al. 2016). Kluyveromyces marxianus can grow very fast at temperatures above 40°C and has been used for the production of bioethanol, biomass and flavor compounds (Morrissey et al. 2015). The thermotolerant and methylotrophic yeast Ogataea polymorpha (syn. Hansenula polymorpha) is a major established platform for heterologous protein expression thanks to the availability of extremely strong yet tightly controlled promoters (Gellissen 2000). In addition, this yeast has been engineered for high-temperature ethanol fermentation from various carbon sources, including xylose (Kurylenko et al. 2014), and glycerol, a by-product originating from biodiesel production (Kata et al. 2016). Ogataea polymorpha shares many characteristics with the closely related species O. parapolymorpha, which was taxonomically separated from O. polymorpha in 2010, and now includes the popular former H. polymorpha DL-1 strain (Suh and Zhou 2010).

The unique physiological characteristics of nonconventional yeast species have the potential to reduce production costs of processes that are currently performed with S. cerevisiae. However, analogous to the situation in industrial S. cerevisiae strains, this will require extensive genetic engineering. While obtaining the necessary genome-sequence information is relatively straightforward due to fast developments in whole-genome sequencing (Goodwin, McPherson and McCombie 2016), genetic modification by classical methods is still challenging mostly due to the predominant DNA-repair mechanisms of many non-conventional yeasts (Nonklang et al. 2008; Abdel-Banat et al. 2010). Unlike S. cerevisiae, these yeasts typically have a very active non-homologous end joining (NHEJ) DNA repair mechanism and use homologous recombination (HR) to a much lesser extent, which makes precise genome editing inefficient as provided linear repair DNA fragments are not efficiently integrated at the targeted genomic locus (Klinner and Schäfer 2004). Introduction of a double-strand break (DSB) in the targeted DNA locus can strongly facilitate genetic engineering, either by introduction of mutations at the cut site or by stimulating the occurrence of HR-mediated DNA repair with co-transformed repair fragments (Jasin and Rothstein 2013; Kuijpers et al. 2013b).

Over the past five years, the CRISPR-Cas9 system has emerged as a powerful and versatile tool to engineer the genomes of a wide range of organisms (Hsu, Lander and Zhang 2014). In this system, the endonuclease CRISPR associated protein 9 (Cas9) binds a guide RNA molecule (gRNA) that targets a sequence-specific site in the genome (Jinek et al. 2012). The Cas9gRNA complex then induces a DSB that is lethal unless repaired. Repair of DSBs typically occurs through NHEJ or HR, depending on the presence of a repair DNA fragment and the predominant DSB repair mechanism of the host cell (Shrivastav, De Haro and Nickoloff 2008). CRISPR-Cas9's ability to induce mutations in the target sequence has been widely exploited in the development of genetic tools for various non-conventional yeasts including K. lactis (Horwitz et al. 2015), K. marxianus (Löbs et al. 2017; Nambu-Nishida et al. 2017), O. polymorpha (Numamoto, Maekawa and Kaneko 2017), Pichia pastoris (Weninger et al. 2016), Scheffersomyces stipitis (Cao et al. 2017) and Yarrowia lipolytica (Gao et al. 2016, Schwartz et al. 2016). However, the plasmid and Cas9/gRNA expression systems available today have generally only been designed for and tested in a single yeast species, thus limiting the potential of each system to work on different yeasts with novel characteristics.

In this study, we developed a novel wide-host-range CRISPR/Cas9 system for use across several yeast species, based on plasmid-borne expression of Cas9 and a ribozyme-flanked gRNA that was recently developed for *S. pasto*rianus, a Saccharomyces hybrid highly resilient to genetic modification (Gorter de Vries et al. 2017). The system was successfully tested in *K. marxianus*, *K. lactis*, *O. polymorpha* and *O. parapolymorpha*. The results highlight the potential of cross-species CRISPR-Cas9 tools for genome engineering in yeasts.

MATERIALS AND METHODS

Strains and growth conditions

The K. lactis, K. marxianus, O. polymorpha and O. parapolymorpha strains used in this study are listed in Table 1. For cultivation under non-selective conditions, strains were grown in 500 mL shake flasks containing 100 mL YPD medium (10 g L⁻¹ Bacto yeast extract, 20 g L⁻¹ Bacto peptone, 20 g L⁻¹ glucose, demineralized water), placed in a rotary shaker set to 30°C and 200 rpm. For antibiotic selection, media for cultivation of Kluyveromyces and Ogataea species were supplemented with 200 and 300 μ g mL⁻¹ hygromycin B, respectively. Prolonged liquid incubation for generation of gene disruptions in Ogataea species was carried out under selective conditions in 50 mL vented tubes (Cellstar CELLreactor, Greiner Bio-One, Kremsmünster, Austria), containing 25 mL YPD medium and incubated as described above. Solid medium was prepared by addition of 2% (w/v) agar. To verify the disruption of nitrate reductase gene YNR1, O. parapolymorpha was grown on synthetic medium (SM) which contained 20 g L⁻¹ glucose, 3 g L⁻¹ KH₂PO₄, 0.5 g L⁻¹ MgSO₄, 7 H₂O, 5 g L^{-1} (NH₄)₂SO₄, 1 mL L^{-1} of a trace element solution and of a vitamin solution as previously described in Verduyn et al. (1992) and on synthetic medium with nitrate (SMN) in which (NH₄)₂SO₄ was substituted with 5 g $L^{-1}\ K_2SO_4$ and 4.3 g $L^{-1}\ NaNO_3.$ For multiplexed targeting of ADE2 and YNR1 in O. parapolymorpha, SM and SMN media were supplemented with 15 mg L⁻¹ adenine to allow for growth of Ade⁻ mutants. Frozen stock cultures were prepared from exponentially growing shake-flask cultures by addition of 30% (v/v) glycerol, and aseptically stored in 1 mL aliquots at -80°C.

Table 1. Kluyveromyces and Ogataea strains used in this study.

Species	Strain	Genotype	Origin
Kluyveromyces lactis	CBS 2359	Wild type	CBS-KNAW ^A , van der Walt (1971)
	IMK829	∆ade2	This study
Kluyveromyces marxianus	CBS 5795	Wild type	CBS-KNAW ^A , van der Walt (1971)
	NBRC 1777	Wild type	NBRC ^B
	IMK830	$\Delta ade2$	This study
	CBS 397	Wild type	CBS-KNAW ^A , van der Walt (1971)
Ogataea polymorpha (syn. Hansenula polymorpha)	CBS 4732	Wild type	CBS-KNAW ^A , Yamada et al. (1994)
Ogataea parapolymorpha (syn. Hansenula polymorpha)	CBS 11895 (DL-1, ATCC	Wild type	CBS-KNAW ^A , Suh and Zhou (2010)
	26012)		
	IMD001	ku80 ^{A340AA}	This study
	IMK828	ku80 ^{A340AA} ∆ade2	This study
	IMD034	ade2 ^{C120CA} ynr1 ^{G397GT}	This study

CBS 2359, CBS 5795, CBS 397, CBS 4732 and CBS 11895 were obtained from the CBS-KNAW fungal collection (^A Westerdijk Fungal Biodiversity Institute, Utrecht, The Netherlands). NBRC 1777 was obtained from the NBRC culture collection (^B National Institute of Technology and Evaluation, Tokyo, Japan).

Table 2. Plasmids used in this study.

Name (Addgene Plasmid #)	Relevant characteristics	Origin
рҮТК079	Template for hph (Hyg ^R) open reading frame	Lee et al. (2015)
pUD423	Template for AaTEF1p-Spcas9 ^{D147Y P411T} -ScPHO5t cassette	Gorter de Vries et al. (2017
pUD527	ori kan ^R SHRA AgTEF1p-amdS-AgTEF1t SHRB	GeneArt
pUD530	ori kan ^R panARS(OPT)	GeneArt
pUD531	ori kan ^R SHRC ScTDH3p- ^{BsaI BsaI} -ScCYC1t SHRI	GeneArt
pUD532	ori kan ^R SHRI ori bla SHRA	GeneArt
pUD540	ori amp ^{R Bsal} HH-gRNA _{OpADE2} -HDV ^{Bsal}	GeneArt
pUD555	ori amp ^{R BsaI} HH-gRNA _{KIADE2} -HDV ^{BsaI}	GeneArt
pUD602	ori amp ^{R Bsal} HH-gRNA _{OpKU80} -HDV ^{Bsal}	GeneArt
pUD750	ori amp ^{R_Bsal} HH-gRNA _{OpADE2} -HDV-HH-gRNA _{OpYNR1} -HDV ^{Bsal}	This study
pUDP002 (plasmid #103872)	ori amp ^R panARS(OPT)	This study
pUDP013 (plasmid #103873)	ori amp ^R panARS(OPT) AgTEF1p-hph-AgTEF1t ScTDH3p-HH-gRNA _{OpADE2} -HDV- ScCYC1t AaTEF1p-Spcas9 ^{D147Y P411T} -ScPHO5t	This study
pUDP025 (plasmid #103874)	ori amp ^R panARS(OPT) AgTEF1p-hph-AgTEF1t ScTDH3p-HH-gRNA _{KIADE2} -HDV- ScCYC1t AaTEF1p-Spcas9 ^{D147Y P411T} -ScPHO5t	This study
pUDP046 (plasmid #107062)	ori amp ^R panARS(OPT) AgTEF1p-hph-AgTEF1t ScTDH3p-HH-gRNA _{OpKU80} -HDV- ScCYC1t AaTEF1p-Spcas9 ^{D147Y P411T} -ScPHO5t	This study
pUDP082 (plasmid #103875)	ori amp ^R panARS(OPT) AgTEF1p-hph-AgTEF1t ScTDH3p-HH-gRNA _{KmADE2} -HDV- ScCYC1t AaTEF1p-Spcas9 ^{D147Y P411T} -ScPHO5t	This study
pUDP123 (plasmid # 107269)	ori amp ^R panARS(OPT) AgTEF1p-hph-AgTEF1t ScTDH3p-HH-gRNA _{OpADE2} -HDV-HH-gRNA _{OpYNR1} -HDV-ScCYC1t AaTEF1p-Spcas9 ^{D147Y P411T} -ScPHO5t	This study

Restriction enzyme sites are indicated in superscript and gRNA target sequences are indicated in subscript. SHRs represent specify synthetic homologous recombination sequences used for plasmid assembly. Aa: Arxula adeninivorans; Sp: Streptococcus pyogenes; Ag: Ashbya gossypii; Sc: Saccharomyces cerevisiae; Op: Ogataea (para)polymorpha; Kl: Kluyveromyces lactis; Km: Kluyveromyces marxianus; HH: hammerhead ribozyme; HDV: hepatitis delta virus ribozyme. pUDP013 (gRNA_{OPADE2}) targets ADE2 in both O. polymorpha and O. parapolymorpha. The addgene plasmid code (when relevant) is indicated next to the plasmid name between brackets.

Construction of plasmids and repair DNA fragments

All plasmids used in this study are described in Table 2. The DNA parts harbored by plasmids pUD527, pUD530, pUD531, pUD532 pUD540, pUD555, pUD602 and pUD750 were *de novo* synthesized by GeneArt (Thermo Fisher Scientific, Waltham, MA, USA).

For the construction of pUDP002, an intermediate plasmid containing the Klebsiella pneumoniae hph (Hyg^R) open reading frame (ORF), expressed from the TEF1 promoter and terminator from the yeast Ashbya gossypii (Eremothecium gossypii), was constructed by 'Gibson' assembly (Gibson et al. 2009) from plasmids pYTK079 (Lee et al. 2015) and pUD527: the hph ORF was amplified from pYTK079 using primers 9837 and 9838, and inserted into a plasmid backbone which was generated by PCR amplification of pUD527 using primers 9839 and 9840. Plasmid pUDP002 was constructed by Gibson assembly from five overlapping fragments, using synthetic homologous recombination sequences (Kuijpers et al. 2013a): (i) the AgTEF1p-hph-AgTEF1t hygromycin resistance cassette was amplified from the intermediate plasmid using primers 9841 and 9842 and sequence-verified before further use; (ii) the AaTEF1p-cas9^{D147Y P411T}-ScPHO5t expression cassette (Bao et al. 2015; Gorter de Vries et al. 2017) was obtained by amplification of pUD423 using primers 3841 and 9393; (iii) the pangenomic yeast replication origin panARS(OPT) (Liachko and Dunham 2014) was amplified from pUD530 using

Locus	Target sequence (5'-3')	Position in ORF (bp)	AT score	RNA score
KlADE2	TTTCAATACCTCAAGTGCTT <u>CGG</u>	508/1710	0.65	0.70
KmADE2	GCCCATTTTTCTGCGTATAG <u>CGG</u>	537/1710	0.55	0.70
OpADE2*	CTGGAATTGATCTGCTTGGC <u>CGG</u>	120/1704	0.50	0.35
OpKU80	CATCGTTCTGCAGAAGATCA <u>TGG</u>	340/2076	0.55	0.55
OpYNR1	AGCACAGACCATAGTAACTG <u>GGG</u>	397/2580	0.55	0.55

Table 3. gRNA target sequences used in this study.

Target sequences are shown including PAM sequence (underlined). The gRNA for OpADE2 targets the respective genes at the same position in both O. polymorpha and O. parapolymorpha. Position in ORF indicates the base pair after which the Cas9-mediated DSB is expected, out of the total length of the ORF. AT score indicates AT content of the 20-bp target sequence. RNA score indicates the fraction of unpaired nucleotides of the 20-bp target sequence, predicted with the complete gRNA sequence using minimum free energy prediction by RNAfold (Lorenz et al. 2011).

*The same sequence was used for single (pUDP013) and for double (pUDP123) editing.

primers 9663 and 3856; (iv) the empty ScTDH3p-ScCYC1t gRNA expression cassette was obtained by digestion of pUD531 with SmaI; (v) the *Escherichia* coli pBR322 origin and *bla* gene for ampicillin selection were obtained by digestion of pUD532 with SmaI. The five fragments were gel-purified and 0.1 pmol of each fragment was assembled in a Gibson assembly reaction. An E. coli clone containing the correctly assembled plasmid (verified by digestion with PdmI) was stocked and pUDP002 (Addgene Plasmid #103872) was used as gRNA entry plasmid for subsequent plasmid construction.

The gRNA expression plasmids pUDP013 (plasmid #103873), pUDP025 (plasmid #103874), pUDP046 (plasmid #107062) and pUDP123 (plasmid #107269) were assembled in a one-pot 'Golden Gate' assembly (Engler, Kandzia and Marillonnet 2008) by BsaI digestion of pUDP002 and a synthesized donor plasmid containing the respective gRNA sequences flanked by ribozymes and BsaI restriction sites. pUDP013, pUDP025, pUDP046 and pUDP123 were constructed using pUD540, pUD555, pUD602 and pUD750, respectively. The Golden Gate assemblies were carried out in a final volume of 20 μ L, using T4 DNA ligase buffer (Thermo Fisher Scientific), 10 U BsaI (New England Biolabs), 1 U T4 DNA ligase (Thermo Fisher Scientific) and 10 ng of both pUDP002 and the respective donor plasmid. For the construction of pUDP082 (plasmid #103875), the partially overlapping primers Km-ade2-F and Km-ade2-R were annealed, extended and amplified by PCR to yield a 233-bp doublestranded DNA fragment containing the KmADE2 gRNA target sequence flanked by ribozymes and BsaI sites, which was integrated into pUDP002 by Golden Gate assembly as described above. All constructed gRNA-harboring plasmids were verified by digestion with PdmI. Plasmids pUDP002, pUDP013, pUDP025, pUDP046, pUDP082 and pUDP123 were deposited at Addgene (https://www.addgene.org/).

The ADE2 repair DNA fragments of K. lactis (962 bp), K marxianus (959 bp), O. polymorpha (960 bp) and O. parapolymorpha (960 bp) were constructed from strains CBS 2359, CBS 5795, CBS 4732 and CBS 11895, respectively, using primer sets 10723 & 10724 and 10725 & 10726 (K. lactis), 10727 & 10728 and 10729 & 10730 (K. marxianus), 10346 & 10347 and 10348 & 10349 (O. polymorpha) and 10354 & 10355 and 10356 & 10357 (O. parapolymorpha) to amplify the homology regions flanking ADE2. Both regions were then joined by overlap extension PCR using primer sets 10723 & 10726, 10727 & 10730, 10346 & 10349 and 10354 & 10357 in the case of K. lactis, K. marxianus, O. polymorpha and O. parapolymorpha, respectively. In all cases, the final repair fragment was gel-purified and further amplified to obtain quantities required for transformation, using primer sets 10723 & 10726, 10727 & 10730, 10346 & 10349, and 10354 & 10357 for K. lactis, K. marxianus, O. polymor pha and O. parapolymorpha, respectively. The primers pairs were designed to amplify the 480 bp (± 1 or 2 bp in some cases) upstream the ATG or the 480-bp terminator region downstream the stop codon of the interrupted/deleted ORF.

Design of gRNA target sequences and BsaI-flanked entry constructs

All 20-bp Cas9 gRNA target sequences used in this study are described in Table 3. Approximately 10 candidate target sequences were chosen from the first third of each targeted ORF, based on the presence of an NGG protospacer adjacent motif (PAM) site. Any target sequence with off-targets (defined as a sequence with NGG or NAG PAM and more than 15 nucleotides identical to the candidate sequence) was excluded, based on a blastn homology search against the respective yeast genome (https://blast.ncbi.nlm.nih.gov). The remaining target sequences were ranked based on AT content ('AT score'; 0 being the lowest and 1 being the highest possible AT content) and secondary structure as predicted with the complete gRNA sequence, using minimum free energy prediction by RNAfold (Lorenz et al. 2011) ('RNA score'; 0 being the lowest and 1 being the highest possible number of unpaired target sequence nucleotides). Finally, the target sequences with the highest cumulative AT and RNA score that did not exceed an AT score of 0.8 were chosen for use in this study.

To integrate ribozyme-flanked gRNAs into pUDP002, the synthetic 233-bp dsDNA gRNA entry constructs were flanked by inward-facing BsaI sites generating sticky ends (underlined) 'GGTCTCG<u>CAAA</u>' and '<u>ACAG</u>CGAGACC' at their 5' and 3' ends, respectively, compatible with BsaI-digested pUDP002. The sequence between the BsaI sites consisted of (i) the hammerhead ribozyme with the first six nucleotides being the reverse complement (c) of the first six nucleotides of the gRNA spacer ^{'c}N₆^cN₅^cN₄^cN₃^cN₂^cN₁CUGAUGAGUCCGUGAGGACGAAACGAGUA AGCUCGUC', (ii) followed by the 20-nucleotide gRNA spacer, followed by the structural gRNA 'GUUUUAGAGCUA-GAAAUAGCAAGUUAAAAUAAGGCUAGUCCGUUAUCAACUU-GAAAAAGUGGCACCGAGUCGGUGCUUUU', (iii) followed by the hepatitis delta virus ribozyme 'GGCCGGCAUGGUCCCAGCC UCCUCGCUGGCGCCGGCUGGGCAACAUGCUUCGGCAUGGCGA AUGGGAC' (Gao and Zhao 2014; Gorter de Vries et al. 2017). For multiplexed targeting of O. parapolymorpha ADE2 and YNR1 using a single expression cassette, two ribozyme-flanked gRNAs were connected in a tandem array using 20-bp linker 'GTGTAATGTCCAGAGTTGTG', and otherwise constructed as described previously (Gorter de Vries et al. 2017).

Strain construction

Yeast transformations

Kluyveromyces lactis and K. marxianus were transformed using the LiAc/single-strand carrier DNA/polyethylene glycol method (Gietz and Schiestl 2007). Overnight pre-cultures in YPD medium were used to inoculate a shake flask containing YPD medium to an initial OD_{660nm} of 0.5. Cultures were then incubated at 30 $^{\circ}C$ until an OD_{660nm} of 2.0 was reached, harvested and transformed with 200 ng of plasmid DNA and 300 or 1000 ng of repair DNA fragment in the case of K. lactis or K. marxianus, respectively. After heat shock, cells were recovered in 1 mL YPD medium for 3 h and plated on selective YPD medium. Plates were typically kept for 5 days at 30°C and then incubated at 4°C for 2 h before assessing the percentage of red Ade⁻ colonies. Ogataea polymorpha and O. parapolymorpha were transformed using the procedure for preparation of competent cells and electroporation described by Saraya et al. (2014), with modifications. All steps were performed at 30° C, OD_{660nm} of all cultures harvested for transformation was normalized to OD_{660nm} 1.2 by dilution with sterile demineralized water, all centrifugation steps were carried out for 3 min at 3000 g, the DTT incubation step was done for 20 min, and the washing step with STM buffer was performed twice with 50 mL. Electroporation was carried out with 40 μ L of a freshly prepared competent cell suspension in 2 mm gap cuvettes (Bio-Rad, Hercules, CA, USA), using 200 ng of plasmid DNA and 1 μ g of repair fragment. A Micropulser Electroporator (Bio-Rad) set to the 'Sc2' preset (1.5 kV) was used for pulse delivery. After electroporation, cells were recovered in 1 mL YPD medium for 1 h at 30°C before plating onto selective YPD medium. For the direct inoculation of prolonged liquid incubation cultures, 100 μ L of recovered cell suspension was used as inoculum. Selection plates were typically kept for 4 days at 30°C and then incubated at 4°C for a minimum of 24 h before assessing the percentage of red Ade⁻ colonies.

Molecular diagnosis of yeast mutants

For molecular analysis of K. lactis and K. marxianus mutants, colonies were grown overnight in YPD medium. Genomic DNA was extracted using the method described by Singh and Weil (2002), and used as template for PCR reactions targeting the ADE2 locus. Primer sets 10909 & 10910, and 10911 & 10912 were used for K. lactis and K. marxianus, respectively. For Sanger sequencing of putative NHEJ-corrected mutants, PCR products were purified and then sequenced using primers 10737 and SeqADE2 for K. lactis and K. marxianus, respectively. For O. polymorpha and O. parapolymorpha, genomic DNA was directly isolated from colonies, using the LiAc-sodium dodecyl sulfate method (Looke, Kristjuhan and Kristjuhan 2011). Primer sets 10380 & 10381, and 10915 & 10916 were used to check for deleted ADE2 loci in O. polymorpha and O. parapolymorpha, respectively, while sequencing of NHEJ-corrected mutants was done using primer sets 10378 & 10379, and 10386 & 10387, respectively. The deleted ADE2 locus in strain IMK828 was amplified and Sanger-sequenced using primers 10915 and 10916. Additionally for O. parapolymorpha primer pair 12257 & 12266 (Table 4) was used to amplify OpYNR1 to verify editing of this locus by Sanger sequencing.

Construction of IMK829, IMK830, IMD001, IMK828 and IMD034 To construct IMK829 (K. lactis $\triangle ade2$), strain CBS 2359 was cotransformed with 200 ng of pUDP025 (harboring gRNA_{KIADE2}) and 300 ng of a 962-bp repair DNA fragment as described above. The resulting transformants were analyzed by diagnostic PCR using primers 10909 and 10910, and a mutant exhibiting a PCR product compatible with deletion of the ADE2 ORF was isolated and renamed IMK829. To construct IMK830 (K. marxianus ∆ade2), strain NBRC 1777 was co-transformed with 200 ng of pUDP082 (harboring gRNA_{KmADE2}) and 1 μ g of a 959-bp repair DNA fragment as described above. The resulting transformants were analyzed by diagnostic PCR using primers 10911 and 10912, and a mutant exhibiting a PCR product compatible with deletion of the ADE2 ORF was isolated and renamed IMK830. For the construction of IMD001 (O. parapolymorpha ku80A340AA), the O. parapolymorpha KU80 ORF (accession XM_014078010.1) was identified by a tblastn homology search (https://blast.ncbi.nlm.nih.gov) using the S. cerevisiae YKU80 protein sequence and O. parapolymorpha CBS 11895 (DL-1) RefSeq assembly (accession GCF_000187245.1) (Ravin et al. 2013). Strain CBS 11895 was transformed with 200 ng of pUDP046 (harboring gRNA_{OpKU80}), and a single transformant was picked and used to inoculate a prolonged liquid incubation culture as described above. After 96 h, cells were plated on selective YPD medium, genomic DNA was isolated from randomly picked colonies, and the KU80 locus was amplified and Sanger-sequenced using primers 10751 and 10752. A mutant containing a single adenine nucleotide inserted between position 340 and 341 of the KU80 ORF was restreaked three times subsequently on non-selective YPD medium to remove pUDP046, and renamed IMD001. To construct IMK828 (O. parapolymorpha ku $80^{A340AA} \Delta ade2$), strain IMD001 was cotransformed with 200 ng of pUDP013 (harboring gRNA_{OpADE2}) and 1 μ g of a 960-bp repair DNA fragment as described above. After recovery, 100 μ L of transformation cell suspension was directly used for inoculation of a prolonged liquid incubation culture. After 48 h, cells were plated on selective medium and the resulting colonies inspected for occurrence of the red Adephenotype. A mutant which exhibited a PCR product compatible with deletion of the ADE2 ORF when analyzed by diagnostic PCR using primers 10915 and 10916 was restreaked thrice subsequently on non-selective YPD medium to remove pUDP013, and renamed IMK828. For the construction of IMD034 (O. parapolymorpha ade2^{C120CA} ynr1^{G397GT}), the O. parapolymorpha HPODL_02384/YNR1 ORF (accession XM_014082012.1) was identified in O. parapolymorpha CBS 11895 (DL-1) RefSeq assembly (accession GCF_000187245.1) as previously described for OpKU80. Strain CBS 11895 was transformed with 200 ng of pUDP123 (harboring dual gRNA_{OpADE2-OpYNR1}), and a single transformant was picked and used to inoculate a prolonged liquid incubation culture as described above. After 96 h, cells were plated on selective YPD medium. A set of 94 Ade- mutants were picked and grown overnight on non-selective YPD and then replica plated on SM and SMN media. A set of five transformants exhibiting both Ade⁻ (red colonies) and Nit⁻ (absence of growth on nitrate) phenotypes were randomly picked, genomic DNA was isolated and the OpADE2 and OpYNR1 loci were amplified and Sangersequenced using primer pairs 10386 & 10387 and 12257 & 12266 respectively. One sequence-confirmed double interruption mutant was restreaked three times subsequently on non-selective YPD medium to remove pUDP123, and renamed IMD034 (Table 1).

Molecular biology techniques

PCR amplification with Phusion High Fidelity Polymerase (Thermo Fisher Scientific) was performed according to the manufacturer's instructions using PAGE-purified oligonucleotide primers (Sigma-Aldrich, St. Louis, MO, USA). Diagnostic PCR was done using DreamTaq polymerase (Thermo Fisher Scientific)

Table 4. Primers used in this study.

Name	Sequence (5' - 3')	Purpose
3841	CACCTTTCGAGAGGACGATG	Construction of pUDP002
3856	CTAGCGTGTCCTCGCATAGTTC	Construction of pUDP002
9393	TGCCGAACTTTCCCTGTATGAAGCGATCTGACCAATCCTTTGCC GTAGTTTCAACGTATGTTTTCATTTTTGCGATGCCAG	Construction of pUDP002
9663	CATACGTTGAAACTACGGCAAAGGATTGGTCAGATCGCTTCAT ACAGGGAAAGTTCGGCATCAACATCTTTGGATAATATCAGAATGAG	Construction of pUDP002
9837	ATACAGTTCTCACATCACATCCGAACATAAACAAGGATCCATG GGTAAAAAGCCTGAACTC	Construction of pUDP002
9838	ACAAGTTCTTGAAAACAAGAATCTTTTTATTGTCCTCGAG	Construction of pUDP002
0000	TTATTCCTTTGCCCTCGGAC	Construction of autopage
9839	CTCGAGGACAATAAAAAGATTCTTG	Construction of pUDP002
9840	GGATCCTTGTTTATGTTCGGATG	Construction of pUDP002
9841	ACTATATGTGAAGGCATGGC	Construction of pUDP002
9842	GTTGAACATTCTTAGGCTGG	Construction of pUDP002
Km-ade2-F	GGTCTCGCAAAGTCAAGCTGATGAGTCCGTGAGGACGAAACG AGTAAGCTCGTCGCCCATTTTTCTGCGTATAGGTTTT AGAGCTAGAAATAGCAAGTTAAAATAAGGCTAGTCC GTTATCAACTTGAAAAAGTGGCACC	Construction of pUDP082
۲m-ade2-R	GGTCTCGCTGTGTCCCATTCGCCATGCCGAAGCATGTTGCCC AGCCGGCGCCAGCGAGGAGGCTGGGACCATGCCGGCCA AAAGCACCGACTCGGTGCCACTTTTTCAA GTTGATAACGGACTAGCCTTATTTTAACTTGC	Construction of pUDP082
10723	GTAGTACCGACCTTATCCGTG	Construction of K. lactis ADE2 repair fragment
10724	GTTGTCTTAGTGAAGAAGGTGAAC	Construction of K. lactis ADE2 repair fragment
10725	TATATAATAACATCACGTTCACCTTCTTCACTAAGACAACAGCTGC CAAATTAGAAACTATCG	Construction of K. lactis ADE2 repair fragment
10726	TGTGCGTTGATATATGCCAAC	Construction of K. lactis ADE2 repair fragment
10727	ATCATAGACAGTCAGTTAGTTCCC	Construction of K. marxianus ADE2 repair fragment
10728	TTCTTTGGTCCATGATTAACAAGG	Construction of K. marxianus ADE2 repair fragment
10729	ACTACAACAATATAAACCTTGTTAATCATGGACCAAAGAAGTATTC AACTACCTCCAACAAGAAG	Construction of K. marxianus ADE2 repair fragment
10730	CAAATTTATGAAGTTTGTGCCATTTG	Construction of K. marxianus ADE2 repair fragment
10346	TGTGCACTCAATTGCAACC	Construction of O. polymorpha ADE2 repair fragment
10347	TTCCAACGACCTTTGAGTCC	Construction of O. polymorpha ADE2 repair fragment
10348	TAATTTAATTTAATTTACATGGACTCAAAGGTCGTTGGAAGTTGGC TATGAGGAATACCTTAAC	Construction of O. polymorpha ADE2 repair fragment
10349	GGGACGTTTACTGGACGG	Construction of O nolymorphy ADE2 ropair fragmont
		Construction of O. polymorpha ADE2 repair fragment
10354	CCTGATGTGCACTCAATTGC	Construction of O. parapolymorpha ADE2 repair fragment
10355 10356	CAACGACCTTCGAGTCCATC TATTAATTTAATTTAATTT	Construction of O. parapolymorpha ADE2 repair fragment Construction of O. parapolymorpha ADE2 repair fragment
	TTGGCTATGAAGAATACC	
10357	GTTTATTGGATGGCAATCTCG	Construction of O. parapolymorpha ADE2 repair fragment
10737	AATTGCATCTCTTTGTGATGTC	Sanger sequencing of K. lactis ADE2 disruptions
SeqADE2	CTGCAACTGCTTGTTCAGCC	Sanger sequencing of K. marxianus ADE2 disruptions
10378	CCAATTACAAGTACTACTTCGAG	Sanger sequencing of O. polymorpha ADE2 disruptions fw
10379	CTAGCTCCTTGGTGAAAGG	Sanger sequencing of O. polymorpha ADE2 disruptions rv
10386	ACAAGTACTACTTCGAGGAC	Sanger sequencing of O. parapolymorpha ADE2 disruptions fv
10387	CTAGCTCCTTGGTAAAGGG	Sanger sequencing of O. parapolymorpha ADE2 disruptions ry
10751	GGACGCCTGCTTAGACTTG	Sanger sequencing of 0. parapolymorpha KU80 disruptions fv
		Sanger sequencing of 0. parapolymorpha KU80 disruptions in Sanger sequencing of 0. parapolymorpha KU80 disruptions rv
10752	AGCACGGTATATTCGCACAG	• • • • • • • •
12257	CACCATGGTCGGAAGAACC	Sanger sequencing of O. parapolymorpha YNR1 disruptions fv
12266	ATGTAATTCCTCACGAACTTTGG	Sanger sequencing of O. parapolymorpha YNR1 disruptions rv
10909	TCTTCGTCGCCATTTATTGTTGAG	Diagnosis of K. lactis ADE2 deletions fw
10910	CTATTGCGGTTCGCTCTTCC	Diagnosis of K. lactis ADE2 deletions rv
10911	ATTCGCCGAATCTGACGTG	Diagnosis of K. marxianus ADE2 deletions fw
10912	TGGTGTGCAGACGGATAAAC	Diagnosis of K. marxianus ADE2 deletions rv
10380	AGGTGCTCAAACACAAAGAG	Diagnosis of O. polymorpha ADE2 deletions fw
10381	TCGTATCTCGTAAGTTGATTTAGG	Diagnosis of O. polymorpha ADE2 deletions rv
10915	CCGTCTGAACGGAATGATGTC	Diagnosis and Sanger sequencing of O. parapolymorpha ADE2 deletions fw
10916	CCCTCAACTGCAGACACATAG	Diagnosis and Sanger sequencing of O. parapolymorpha ADE2 deletions rv

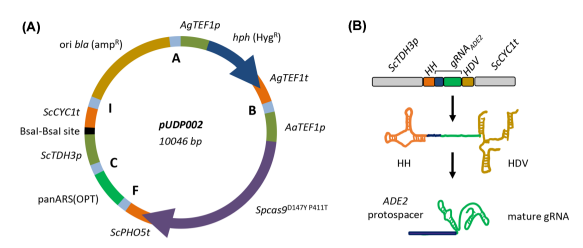


Figure 1. Components of the pUDP genome editing system. (A) Map of pUDP002 (Addgene plasmid #103872), a wide-host-range gRNA entry plasmid. pUDP002 is composed of a hph (Hyg^R) selection marker cassette under control of the TEF1 promoter from Ashbya gossypii conferring hygromycin resistance, Spcas9^{D147Y P411T} under control of the TEF1 promoter from Arxula adeninivorans, the pangenomic yeast replication origin panARS(OPT), a BSaI cloning site for entry of gRNA constructs, and a pBR322-derived *E*. coli origin and bla gene for ampicillin selection. A, B, F, C and I indicate 60 bp synthetic homologous recombination sequences used for 'Gibson' assembly of the plasmid. (B) Representation of the ribozyme-flanked gRNA expression cassette design. Guide RNAs (represented by gRNA_{ADE2}) were flanked on their 5' by a hammerhead ribozyme (HH represented in orange) and on their 3' by a hepatitis delta virus ribozyme (HDV represented in bronze). When integrated into pUDP002, this construct is under control of the RNA polymerase II promoter TDH3 and the CYC1 terminator from S. *cerevisiae*. Upon ribozyme self-cleavage, a mature gRNA comprised of the guiding protospacer (in blue) and the structural gRNA fragment (in green) is released.

and desalted primers (Sigma-Aldrich). All primer sequences are shown in Table 4. DNA fragments obtained by PCR were separated by gel electrophoresis. Gel purification was carried out using the Zymoclean Gel DNA Recovery Kit (Zymo Research, Irvine, CA, USA). PCR purification was performed using either the GenElute PCR Clean-Up Kit (Sigma-Aldrich) or GeneJET PCR purification kit (Thermo Fisher Scientific). Gibson assembly was done using the NEBuilder HiFi DNA Assembly Master Mix (New England Biolabs, Ipswich, MA, USA) according to the manufacturer's recommendations. Restriction digest with PdmI and SmaI was performed using FastDigest enzymes (Thermo Fisher Scientific), according to the manufacturer's instructions. *Escherichia* coli strain XL1-blue was used for plasmid transformation, amplification and storage. Plasmids were isolated from E. coli with the GenElute Plasmid Miniprep Kit (Sigma-Aldrich).

Whole-genome sequencing of O. parapolymorpha IMD001

Genomic DNA of O. parapolymorpha IMD001 was isolated with the QIAGEN Genomic-tip 100/G kit (Qiagen, Hilden, Germany) from a stationary-phase overnight shake-flask culture grown on YPD medium, according to the manufacturer's instructions. Genomic DNA was quantified using a Qubit® 2.0 Fluorometer (Thermo Fisher Scientific). Ten micrograms of genomic DNA was sequenced by Novogene Bioinformatics Technology (Yuen Long, Hong Kong, China) on a HiSeq 2500 (Illumina, San Diego, CA, USA) with 150-bp paired-end reads using TruSeq PCR-free library preparation (Illumina). In order to verify complete absence of plasmid pUDP046, sequencing reads were mapped to the sequence of pUDP046 and to the genome of O. parapolymorpha CBS 11895 (accession GCF_000187245.1) (Ravin et al. 2013) using the Burrows-Wheeler Alignment tool and further processed using SAMtools (Li et al. 2009; Li and Durbin 2010). The absence of sequences from pUDP046 was confirmed by visualizing the generated .bam files in the Integrative Genomics Viewer software (Robinson et al. 2011). Sequencing data are available at NCBI (https://www.ncbi.nlm.nih.gov/) under BioProject PRJNA413643.

RESULTS

pUDP: a plasmid-based wide-host-range yeast CRISPR/Cas9 system

The pUDP system was designed to enable Cas9-mediated genome editing in different yeast species by simple transformation with a single plasmid. To this end, DNA parts encoding the plasmid origin of replication, cas9 expression cassette, selection marker and gRNA expression cassette were chosen to function in a wide range of yeast species (Fig. 1). To ensure replication of pUDP plasmids, the pangenomic yeast origin of replication panARS(OPT) was used. This origin of replication, which was inspired by a K. lactis chromosomal ARS, has been shown to function in at least 10 different species of ascomycetous yeasts, including the species used in this study (Liachko and Dunham 2014). The cas9^{D147Y P411T} nuclease variant (Bao et al. 2015) was expressed under control of the TEF1 promoter from Arxula adeninivorans (Blastobotrys adeninivorans), which enabled strong constitutive expression in various yeasts (Steinborn et al. 2006; Terentiev et al. 2004). Similarly, the K. pneumoniae hph gene encoding hygromycin B phosphotransferase, which conferred hygromycin resistance in a wide range of microorganisms, was expressed under control of the Ashbya gossypii (Eremothecium gossypii) TEF1 promoter (Wach et al. 1994) that showed activity in a range of yeasts (Mazzoni, Serafini and Falcone 2005; Kim et al. 2010). To avoid RNA modifications that interfere with biological function, transcription of gRNAs for genome editing is commonly controlled by RNA polymerase III promoters. By analogy with expression of gRNA in mammalian cells (Cong et al. 2013; Mali et al. 2013), the RNA polymerase III SNR52 promoter was used in S. cerevisiae (DiCarlo et al. 2013; Mans et al. 2015), but accurate annotation and characterization of this promoter might not be available for other yeast species. Therefore, self-processing ribozyme-flanked gRNAs were expressed from an RNA polymerase II promoter (Gao and Zhao 2014; Ryan et al. 2014). This concept has already been successfully applied in several different organisms (Gao et al. 2016; Gorter de Vries et al. 2017; Weninger et al. 2016). In this system, the gRNA is

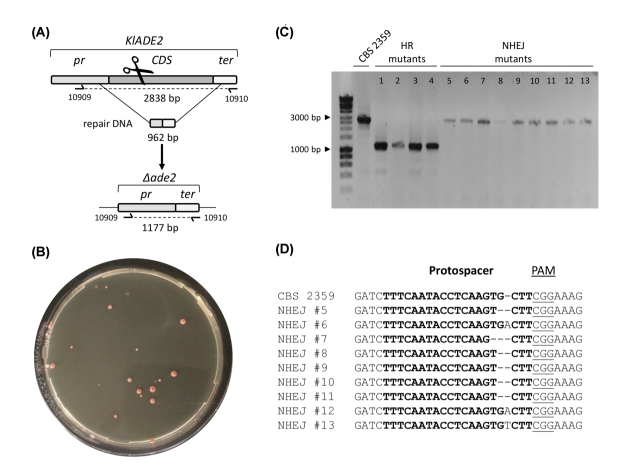


Figure 2. Efficient gRNA targeting in K. lactis CBS 2359 enables marker-free gene deletion. (A) Schematic representation of ADE2 editing upon transformation of CBS 2359 with pUDP025 (gRNA_{KLADE2}) and a repair DNA fragment. The primers for diagnostic PCR of transformants are indicated. (B) Colony morphology of CBS 2359 upon transformation with pUDP025 and a marker-free 962 bp repair fragment. (C) Diagnosis of 13 randomly picked red Ade⁻ transformants of CBS 2359 upon transformation with pUDP025 and a 962-bp marker-free 962 bp repair fragment. Four transformants (IR mutants 1–4) showed a PCR product of 1177 bp corresponding to the deleted allele. The control labeled CBS 2359 and nine transformants (NHEJ mutants 5–13) showed a PCR product of 2838 bp corresponding to the wild-type allele. (D) Sanger sequencing results of purified PCR fragments from nine Ade⁻ mutants (corresponding to mutants 5–13 in panel C) derived from the transformation of CBS 2359 with pUDP025 and repair fragment.

flanked by a hammerhead and hepatitis delta virus ribozyme at its 5' and 3' end, respectively, resulting in precise release of mature gRNA after self-cleavage. To ensure sufficient gRNA transcription, the strong glycolytic TDH3 promoter from S. *cerevisiae* was used. These elements were combined in plasmid pUDP002, which could subsequently serve as entry plasmid to insert any desired gRNA. To streamline integration of ribozyme-flanked gR-NAs, pUDP002 contained two different BsaI restriction sites between ScTDH3p and ScCYC1t, enabling directional insertion of synthetic BsaI-flanked gRNA constructs (Gorter de Vries *et al.* 2017).

To test the genome-editing efficiency of the pUDP system, the ADE2 gene was targeted in all four non-conventional yeasts used in this study: K. lactis, K. marxianus, O. polymorpha and O. parapolymorpha. The ADE2 gene encodes a phosphoribosylaminoimidazole carboxylase, also referred to as AIR carboxylase, involved in adenine biosynthesis. Besides causing adenine auxotrophy, loss-of-function mutations in ADE2 result in a red-color phenotype due to accumulation of the oxidized form of 5-amino imidazole ribonucleotide (AIR). This allows detection of ADE2 targeting by simple visual inspection of transformation plates (Roman 1956). Therefore, gR-NAs were designed targeting ADE2 in each species based on available genome data (Table 3), and gRNA-harboring plasmids pUDP025, pUDP082 and pUDP013 for deletion of ADE2 in K. lactis, K. marxianus and O. polymorpha/O. parapolymorpha, respectively, were constructed.

Efficient CRISPR/Cas9 targeting enables gene deletion in Kluyveromyces species

To test the effectiveness of the pUDP system in K. lactis, gRNA_{KIADE2} was inserted into pUDP002 and the resulting plasmid pUDP025 was used to target ADE2 in K. lactis CBS 2359. Transformation with pUDP025 without a repair DNA fragment yielded a total of 35 transformants of which 19 (54%) exhibited a red Adephenotype. In the presence of a 962-bp repair fragment which targeted the ADE2 promoter and terminator (Fig. 2A), the transformation of pUDP025 yielded 26 red transformants out of a total 27 (96%) (Fig. 2B). In comparison, transformation of the backbone plasmid pUDP002, which does not express a gRNA, generated a number of transformants that was ca. 35-fold higher, none of which displayed a red phenotype. This difference in transformation efficiency already provided information about the quality of the gRNA and the editing. Unless repaired a chromosomal DSB should be lethal, therefore this 35-fold drop in transformation efficiency would represent the fraction of transformants that did not successfully repair the CRISPR-Cas9 induced break and subsequently died from it. Out of the 26 Ade⁻ transformants obtained after co-transformation with pUDP025 and repair

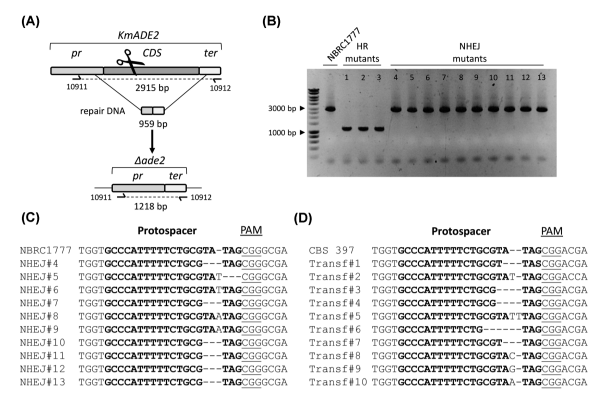


Figure 3. Efficient gRNA targeting enables marker-free gene deletion in haploid K. marxianus NBRC 1777 and gene disruption in diploid K. marxianus CBS 397. (A) Schematic representation of the ADE2 editing upon transformation of NBRC 1777 with pUDP082 (gRNA_{KmADE2}) and a repair DNA fragment. The primers for diagnostic PCR of transformants are indicated. (B) Diagnosis of 13 randomly picked red Ade⁻ transformants of NBRC 1777 upon transformation with pUDP082 and a 959-bp marker-free repair fragment. Three transformants (LR mutants 1–3) showed a PCR product of 1218 bp corresponding to the deleted allele. The control labeled NBRC 1777 and 10 transformants (NHEJ mutants 4–13) showed a PCR product of 2915 bp corresponding to the wild-type allele. (C) Sanger sequencing results of purified PCR fragments (or mutants 4–13 in panel B) derived from the transformation of NBRC 1777 with pUDP082 and repair fragment. (D) Sanger sequencing results of purified PCR fragments of 10 randomly picked red Ade⁻ mutants derived from the transformation of CBS 397 with pUDP082 and repair fragment.

fragment, 13 red colonies were randomly picked and analyzed by diagnostic PCR, revealing two distinct groups. Nine colonies exhibited a normally sized ADE2 locus, while four colonies (31%) showed a PCR-fragment size compatible with complete deletion of the KlADE2 ORF (Fig. 2C). Subsequent Sanger sequencing of the amplified fragments derived from the nine clones exhibiting an ADE2 wild-type size band identified the presence of indels at the targeted site. These indels reflected imperfect repair of the Cas9-induced DSB via NHEJ, resulting in introduction of loss-of-function mutations and disruption of the Cas9 target site (Fig. 2D). These results demonstrated that the pUDP system could be used for efficient targeting of the ADE2 gene in K. lactis CBS 2359, resulting in repair by either HR or NHEJ DNA repair mechanisms.

The pUDP system was similarly tested in K. marxianus by targeting ADE2 in the haploid strain NBRC 1777 and the diploid strain CBS 397, using plasmid pUDP082 expressing gRNA_{KmADE2}. The transformation of K. marxianus NBRC 1777 with pUDP082, without and with a 959-bp repair DNA fragment resulted in 13 and 30 transformants, respectively. Of these, 13 out of 13 and 29 out of 30 transformants were red, indicating successful disruption of ADE2 in both cases. To determine which repair mechanism contributed to the repair of the Cas9-induced DSB in the presence of the repair fragment, 13 randomly picked red transformants were subjected to diagnostic PCR. Similar to the results obtained in K. lactis, two groups of transformants were identified. Three transformants (24%) showed a genotype corresponding to the repair of the DSB with the repair fragment (Fig. 3A and B), while the remaining transformants exhibited a PCR result that was undistinguishable from that of the NBRC 1777 ADE2 wild-type locus. Sanger sequencing of the Cas9 target site again revealed the presence of indels, leading to nonsense mutations in the ADE2 coding sequence (Fig. 3C).

When the diploid strain CBS 397 was transformed with pUDP082 without repair DNA fragment, 106 out of 133 (80%) of the colonies were red, while 117 out of 143 (82%) colonies showed the red phenotype when a 959-bp repair fragment was provided. In comparison, a control transformation with pUDP002 yielded 262 transformants without any red phenotypes. From the transformation with pUDP082 and repair fragment, 10 red colonies were randomly picked and analyzed by diagnostic PCR, resulting in PCR products that were identical in size to the native ADE2 locus in all cases. Sanger sequencing of the same mutants confirmed the link between the red-colored phenotype and occurrence of small indels at the Cas9 cut site, which likely introduced loss of function mutations in ADE2 (Fig. 3D). Interestingly, Sanger sequencing also indicated that the editing occurred identically at both KmADE2 alleles as a clear and continuous sequence was observed over the cut site. These results demonstrated that the ADE2 gene can be efficiently targeted by the pUDP system in both K. marxianus strains, with DNA repair mediated by HR or NHEJ mechanisms in the haploid strain NBRC 1777 and NHEJ in the diploid strain CBS 397.

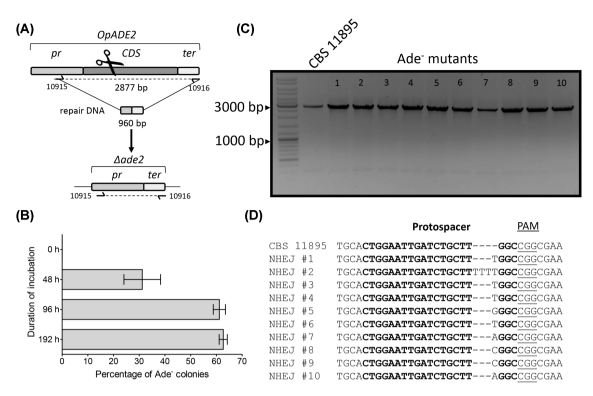


Figure 4. Prolonged liquid incubation enables gene disruption in O. *parapolymorpha* CBS 11895 (DL-1). (A) Schematic representation of the potential ADE2 editing upon transformation of CBS 11895 with pUDP013 (gRNA_{OpADE2}) and a repair DNA fragment. (B) Percentage of red Ade⁻ colonies observed based on phenotypic screening of CBS 11895 + pUDP013 (gRNA_{OpADE2}) transformants when plated directly after transformation (0 h), or after prolonged liquid incubation of established transformants in selective medium (for 48, 96 and 192 h) and subsequent plating. Mean and standard deviation were calculated with a total of 4411, 4710 and 4301 colonies (obtained after 48, 96 and 192 h), respectively) from two liquid incubation cultures started with single CBS 11895 transformants which were obtained in independent transformations. No red Ade⁻ mutants were observed when cells were plated directly after transformation. (C) Diagnosis of 10 randomly picked red Ade⁻ colonies of CBS 11895 after transformation with pUDP013 and 96 h of subsequent liquid incubation under selective conditions. All 10 transformants (NHEJ mutants 1–10) and the control labeled CBS 11895 showed a PCR product of 2877 bp corresponding to the wild-type allele. (D) Sanger sequencing results of purified PCR fragments from 10 red Ade⁻ colonies (corresponding to mutants 1–10 in panel B) derived from the transformation of CBS 11895 with pUDP013 and 96 h of subsequent liquid from the transformation of CBS 11895 with pUDP013 and 96 h of subsequent liquid from the transformation of CBS 11895 with pUDP013 and 96 h of subsequent liquid from the transformation of CBS 11895 with pUDP013 and 96 h of subsequent liquid incubation under selective conditions.

CRISPR/Cas9 editing enables gene deletion in Ogataea species

To test the effectiveness of the pUDP system in O. parapolymorpha, plasmid pUDP013 harboring gRNA_{OpADE2} was used to transform strain CBS 11895 (DL-1). Transformations were performed with or without a 960-bp repair DNA fragment designed to mediate HR at the promoter and terminator regions of the OpADE2 gene (Fig. 4A). In presence of the repair fragment, an average of 298 \pm 50 colonies per transformation were obtained, while a lower average number of 64 \pm 9 colonies were counted after transformation without the repair fragment. In contrast to the situation in Kluyveromyces species, none of these colonies exhibited a red Ade⁻ phenotype, although the ability of the transformants to grow on selective medium indicated that the plasmid was present. The heterologous origin of genetic parts on pUDP002 might result in a suboptimal expression of the necessary components of the CRISPR-Cas9 machinery in O. parapolymorpha. To allow more time for expression of Cas9 and gRNA, two randomly picked transformants (CBS 11895 + pUDP013) from independent experiments were incubated for a longer period in selective YPD medium. These prolonged liquid incubation cultures were sampled after 48, 96 and 192 h, and samples taken at each of these time points were plated on selective YPD medium. Already after 48 h of incubation, $31 \pm 7\%$ of the plated colonies exhibited the red Ade- phenotype. This fraction increased to $61 \pm 2\%$ and $63 \pm 2\%$ after 96 and 192 h of incubation, respectively (Fig. 4B). From a plate obtained after 96 h of liquid incubation, 10 red colonies were randomly picked and subjected to diagnostic PCR. All 10 transformants showed a band compatible with the size of the wild-type ADE2 locus (Fig. 4C). As previously found in K. lactis and K. marxianus, Sanger sequencing of the ADE2 gene in these 10 transformants (NHEJ #1 to #10) confirmed the presence of short indels at the Cas9 target site, which introduced nonsense mutations within ADE2 (Fig. 4D). The presence of the repair fragment did not change this outcome: prolonged liquid incubation cultures that were started from colonies obtained by co-transformation of pUDP013 and the repair fragment gave rise to identical proportions of Ademutants as those described above. For the closely related species O. polymorpha, the obtained results were qualitatively comparable. Transformation of strain O. polymorpha CBS 4732 with pUDP013 did not result in colonies with a red phenotype, neither in the presence nor in the absence of a 960-bp repair DNA fragment. Also, no mutations in ADE2 were detected as evaluated by Sanger sequencing of nine randomly picked (white) colonies from a transformation without the repair fragment. When two independent transformants (CBS 4732 + pUDP013) were incubated in liquid medium, plating after 192 h yielded 9 \pm 1% of red colonies. These results indicated that the pUDP system promoted less efficient genome editing in the O. polymorpha strain CBS 4732 than in O. parapolymorpha CBS 11895, although the system could be utilized for NHEJ-mediated disruption of the ADE2 gene in both yeasts.

In contrast to S. cerevisiae (Baudin et al. 1993) and Kluyveromyces species (this study), the HR-mediated DNA

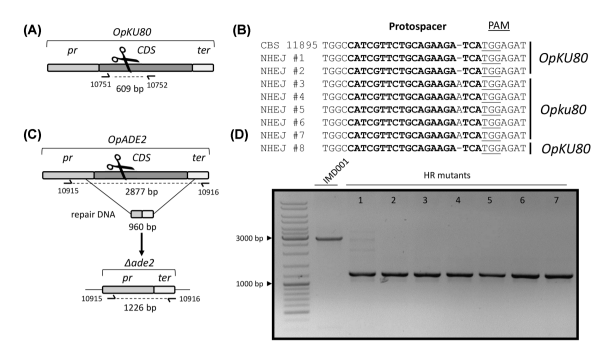


Figure 5. Construction of IMD001, an *O. parapolymorpha* ku80 strain that enables low efficiency gene deletion. (A) Schematic representation of the *OpKU80* editing upon transformation of CBS11895 with pUDP046 (gRNA_{OpKU80}). The primers for diagnostic PCR and sequencing of transformants are indicated. (B) Sanger sequencing results of purified PCR fragments from eight randomly picked colonies derived from the transformation of CBS 11895 with pUDP046 (gRNA_{OpKU80}) and 96 h of subsequent liquid incubation. The transformants NHEJ #1,NHEJ#2 and NHEJ#8 displayed a wild-type sequence, while mutants NHEJ #3 to #7 included frameshift mutations (C) Schematic representation of the ADE2 editing upon transformation of *O. parapolymorpha* IMD001 (CBS 11895 ku80) with pUDP013 (gRNA_{OpADE2}) and a repair DNA fragment. The primers for the diagnostic PCR of transformants are indicated. (D) Diagnosis of all seven red Ade⁻ colonies obtained (from a total of ca. 1900 colonies) upon transformation of IMD001 with pUDP013 and a 960-bp marker-repair fragment, and subsequent liquid incubation in selective conditions for 48 h, started directly from the transformation recovery culture. All mutants showed a PCR product of 1226 bp corresponding to the deleted allele. The control labeled IMD001 showed a PCR product of 2915 bp corresponding to the wild-type allele.

repair mechanism in Ogataea species is relatively inefficient (Klinner and Schäfer 2004). The removal of KU80 in O. polymorpha has been reported to result in a strong reduction of NHEJ in return favoring the occurrence of HR-mediated DNA repair (Saraya et al. 2012). To delete KU80 in O. parapolymorpha, plasmid pUDP046 harboring gRNA_{OpKU80} was constructed and used to transform strain CBS 11895 (Fig. 5A). Following the liquid incubation procedure established for disruption of OpADE2, a single transformant was picked, incubated in selective YPD medium for 96 h and plated on selective YPD plates. Of eight colonies randomly picked and subjected to Sanger sequencing of the KU80 locus, five (NHEJ #3 to #7) contained a single adenine nucleotide inserted at the Cas9 targeting site between position 340 and 341 of the OpKU80 coding sequencing (A340AA), resulting in a loss-of-function mutation (Fig. 5B). One of these transformants was isolated by restreaking three times on non-selective medium, and renamed IMD001 (Opku80^{A340AA}).

While strain IMD001 was not able to grow on selective medium indicating loss of pUDP046, complete curation of the plasmid had to be verified in order to use IMD001 for subsequent genetic interventions with pUDP plasmids. Circular plasmids with limited homology to the nuclear DNA have been reported to integrate into the genome of *Ogataea* (*Hansenula*) yeasts, despite the presence of an origin of replication on the plasmid (Kunze, Kang and Gellissen 2009). In particular, upon cultivation under selective conditions, plasmids were found to have integrated into the nuclear genome with high variations in copy number in strain CBS 11895 (DL-1) (Kang *et al.* 2001). To verify that no pUDP046 plasmid sequences had recombined into the chromosomal DNA of strain IMD001, its genome was sequenced using Illumina 150-bp pair-end short reads that were mapped onto a reference comprised of an *O. parapolymorpha* CBS 11895 genome assembly (accession number: GCF_000187245.1) and the sequence of pUDP046. No reads were found to map onto the pUDP046 sequence, thus confirming the absence of unwanted integration of the transformed plasmid into the nuclear genome and the suitability of *O. parapolymorpha* IMD001 as a host strain for further genetic engineering.

To test the impact of the KU80 disruption on the pUDP system in O. parapolymorpha, plasmid pUDP013 (gRNA_{OpADE2}) was used to transform strain IMD001 together with the 960-bp repair DNA fragment. As previously observed for the wild-type strain O. parapolymorpha CBS 11895, no red Ade- transformants were observed on the transformation plates. In an attempt to improve occurrence of homology-mediated repair, a liquid incubation culture was directly inoculated with the transformation mix, grown for 48 h and then plated on selective YPD medium. On these plates, only 7 out of \sim 1900 colonies exhibited a red Ade⁻ phenotype, representing a <1% targeting efficiency. Diagnostic PCR performed on these seven transformants revealed that all harbored a deleted ADE2 version compatible with HRmediated repair of the CRISPR-Cas9-induced DSB (Fig. 5C and D). One of the obtained mutants was further analyzed by Sanger sequencing, which confirmed scarless integration of the repair fragment. Longer liquid incubation times of up to 192 h did not increase the incidence of red colonies after plating. These results demonstrated that the KU80 disruption abolished NHEJmediated DSB repair in O. parapolymorpha IMD001, and that the CBS 11895 strain background possesses a basal HR-mediated DNA repair activity that can be utilized for precise genome

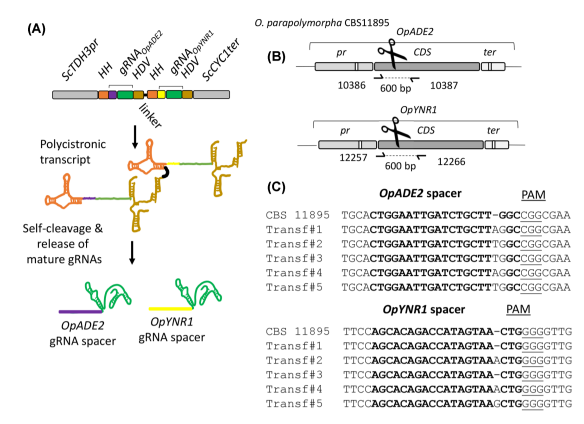


Figure 6. Simultaneous deletion of OpADE2 and OpYNR1 alleles using a single ribozyme flanked gRNA array in O. parapolymorpha CBS11895. (A) Representation of the gRNA array expression cassette in pUDP123. The dual gRNA array was under the control of the RNA polymerase II promoter ScTDH3 and ScCYC1 terminator. Each gRNA was flanked on its 5' by a hammerhead ribozyme (HH represented in orange) and on its 3' by a hepatitis delta virus (HDV represented in bronze) ribozyme which were separated by a 20-bp linker. Upon ribozyme self-cleavage, the mature gRNAs are released. The OpADE2 guiding spacer (in purple), the OpYNR1 guiding spacer (in yellow) and the constant structural gRNA fragment (in green) are indicated. (B) Schematic representation of the OpADE2 and OpYNR1 loci of CBS11895. The primers for the validation of transformants are indicated. (C) Sanger sequencing results of OpADE2 and OpYNR1 editing site of five randomly picked red Ade⁻ mutants that have lost ability to grow on nitrate. Transformant labeled Transf#2 was renamed IMD034.

engineering. However, the low incidence of HR-mediated repair precludes efficient genome editing unless the desired phenotype is easily screenable.

Multiplexed gene targeting in O. parapolymorpha by expression of double ribozyme-flanked gRNA array

As recently demonstrated, HH-HDV ribozyme-flanked gRNAs can be concatenated into polycistronic arrays enabling multiplexing gene editing in *S. pastorianus* (Gorter de Vries et al. 2017) and transcriptional interference in *S. cerevisiae* (Deaner, Mejia and Alper 2017). To explore this possibility in *O. parapolymorpha*, a gRNA expression plasmid pUDP123 carrying spacers targeting *OpADE2* and *OpYNR1* was designed. The additionally targeted YNR1 gene encodes a nitrate reductase, which is involved in utilization of nitrate as N-source (Brito et al. 1996; Navarro et al. 2003). *Ogataea* strains harboring a non-functional nitrate reductase are unable to grow on media with nitrate as sole N-source.

A tandem array of [HH-gRNA-HDV] targeting OpADE2 and OpYNR1 spaced with a 20-bp linker and expressed under control of the ScTDH3 promoter as previously described (Gorter de Vries et al. 2017) was carried by the recombinant plasmid pUDP123 (hph cas9 ScTDH3p-HH-gRNA_{OpADE2}-HDV-HH-gRNA_{OpYNR1}-HDV-ScCYC1t) that was transformed in *O. parapolymorpha* CBS 11895 (Fig. 6A). Following the liquid incubation procedure established for disruption of OpADE2 and OpKU80, a transformant was picked, incubated in selective YPD medium for 96 h and then plated onto selective YPD medium. Of the resulting colonies,

32% were exhibiting a red Ade⁻ phenotype. To verify whether these Ade- mutants were able or not to grow on nitrate, 94 red transformants were randomly picked and grown on nonselective medium (YPD). After full growth, 10 μ l of cell suspension was replicated on non-selective (SM) and selective (SMN) media supplemented with 15 mg L⁻¹ adenine to complement the Ade⁻ phenotype. About 10% of the red-phenotype transformants gradually returned white. While one cannot exclude a reversion of the mutation, the pattern observed would suggest that the replicated red colonies were not pure. Growth of the transformant on SMN was disturbed by the addition of adenine which might also serve as nitrogen source as O. parapolymorpha is equipped with an adenine deaminase. However, since the adenine concentration remained ~300-fold lower than the nitrate concentration, a significant difference between Nit+ and Nit- strains should be noticeable. Indeed out of the 94 red transformants, 17 showed a strongly reduced growth on SMN supplemented with adenine when compared to the control CBS 11895, and 5 of these potential double mutants were further analyzed by Sanger sequencing. While all five were confirmed to harbor a frameshift around the CRISPR cut site in OpADE2, only three transformants were concomitantly exhibiting a frameshift in OpYNR1 (Fig. 6C). Transformant #2 was isolated by restreaking three times on non-selective medium, and renamed IMD034 (Opade2^{C120CA} Opynr1^{G397GT}). Assuming that all other red clones that were also growing slowly on SMN medium shared this genotype, double editing would have occurred at a rate of 2%-5%.

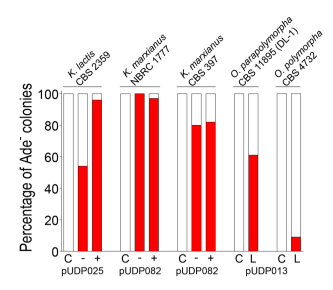


Figure 7. Wide-host-range applicability of the pUDP CRISPR/Cas9 system. The depicted data summarize the ADE2 targeting efficiency (red colonies/total colonies) of the pUDP system in the four yeast species used in this study. The pUDP plasmids differ only in their gRNA target specificity: pUDP025, pUDP082 and pUDP013 harbor gRNA_{KIADE2}, gRNA_{KMADE2} and gRNA_{OpADE2}, respectively. The results shown for Kluyveromyces species were obtained directly on transformation plates either with (+) or without (-) co-transformation of a ca. 960-bp marker-free repair DNA fragment. The results for O. parapolymorpha and O. polymorpha were obtained after 96 and 192 h of prolonged liquid incubation (L), respectively, started from colonies that were transformed without a repair fragment. Control transformations (C) with pUDP002 did not result in the occurrence of Ade $^-$ mutants.

DISCUSSION

The pUDP system, which combines generic DNA parts with expression of self-processing gRNAs driven by RNA polymerase II, was shown to enable genome editing in different yeasts belonging to the Saccharomycotina subphylum. The pUDP002 entry plasmid only required insertion of a functional gRNA to enable successful single deletion or disruption of the ADE2 gene in four different species of *Kluyveromyces* and *Ogataea* yeasts (Fig. 7) but also double editing at two different chromosomal loci in *O. parapolymorpha*. The panARS(OPT)-harboring pUDP plasmids can be cured simply by growth in non-selective medium (Liachko and Dunham 2014), as illustrated by whole-genome sequencing of engineered *O. parapolymorpha* strain IMD001, enabling sequential rounds of genetic modifications with the same system in rapid succession.

In Kluyveromyces species, near-perfect targeting (\geq 96%) and HR-mediated repair DNA integration occurred at practicable rates in K. lactis CBS 2359 (31%) and the haploid K. marxianus NBCR 1777 (24%), while highly efficient gene disruption was achieved in the diploid K. marxianus strain CBS 397. In Ogataea species, mutants could be obtained after prolonged liquid incubation, allowing relatively straightforward and markerfree disruption of genes in the two industrially relevant O. (para)polymorpha strains CBS 4732 and CBS 11895 (DL-1). The observed delay in targeting activity suggests that the heterologous origin of functional parts of the pUDP system might reflect suboptimal expression of gRNA and/or Cas9 in Ogataea species. The expression levels of gRNA and Cas9 may have narrow optima for efficient genome editing, as recently demonstrated in P. pastoris, a related methylotrophic yeast (Weninger et al. 2016). While the A. adeninivorans TEF1 promoter employed for Cas9 expression in this study has been demonstrated to enable strong constitutive expression in *O. polymorpha* (Terentiev *et al.* 2004), optimization of the gRNA expression presently under the control of the *S. cerevisiae* TDH3 promoter might be envisaged to eliminate the observed delay in the occurrence of mutants.

Recently, species-specific CRISPR-Cas9 systems have been published for a range of non-conventional yeasts including K. lactis, K. marxianus and O. polymorpha (Horwitz et al. 2015; Löbs et al. 2017; Nambu-Nishida et al. 2017; Numamoto et al. 2017). To our knowledge, the study by Horwitz et al. (2015) is the only published application of CRISPR-Cas9 in K. lactis. The authors report Cas9-mediated integration at three genomic loci simultaneously, which occurred at a rate of 2% in a NHEJ-deficient strain with an integrated Cas9 cassette using ca. 1000-bp homology flanks. However, data on targeting efficiency were not provided, preventing a meaningful comparison with the results obtained by the pUDP system in this study (96% targeting efficiency, 31% HR-rate, wild-type K. lactis strain, ca. 480-bp homology flanks, single locus). However, the absolute quantification of the HR-mediated repair left no doubt regarding pUDP tool efficacy for gene editing in K. lactis. In K. marxianus NBRC 1777, the pUDP system achieved higher targeting efficiencies (97% vs 65%) and similar efficiencies of marker-free HR (22% vs 38%) compared to the most efficient CRISPR system for this yeast reported to date (Nambu-Nishida et al. 2017). However, in this comparison it should be taken into account that, in this study, longer homology flanks were used (480 bp vs 50 bp), which may have facilitated HR-mediated repair. The published CRISPR-Cas9 tool for O. polymorpha (Numamoto et al. 2017) used strain BY4330, a mutant derived from the O. polymorpha NCYC495 background which is closely related to strain CBS 4732 employed in this study. Compared to this tool, the pUDP system achieved similar efficiencies of NHEJ-mediated gene disruption when compared with O. parapolymorpha CBS 11895 (61% vs 71%), but was less effective when compared with O. polymorpha CBS 4732 (9% vs 71%). In contrast to our findings, the authors were able to obtain gene-disrupted and marker-integrated transformants directly on transformation plates, indicating either a more suitable expression of gRNA and/or Cas9 or a different genetic tractability of the strain lineages. Compared to the pUDP system, all three discussed CRISPR/Cas9 tools (partially) rely on speciesspecific genetic parts. The pKD1 stability element was employed for plasmid replication in K. lactis (Horwitz et al. 2015), while the KmSNR52 promoter was used for gRNA expression in the case of K. marxianus (Nambu-Nishida et al. 2017), and an OptRNA^{Leu} promoter for gRNA expression in O. polymorpha (Numamoto et al. 2017). These RNA polymerase III-dependent promoters are unlikely to function in other species or genera.

RNA polymerase III-driven SpCas9-based editing strategies, such as those based on chimeric gRNA expression systems, are not easily compatible with multiplexed genome editing. Hitherto, the maximum number of gRNAs shown to enable simultaneous DSBs at different loci in S. cerevisiae is six. However, this result required an elaborate construction scheme using three plasmids for the individual expression of each gRNA (Mans et al. 2015). In contrast, ribozyme-flanked gRNA expression, as previously reported in S. pastorianus (Gorter de Vries et al. 2017), allowed to achieve the first double gene editing in O. parapolymorpha using a polycistronic gRNA array including two ribozyme-flanked spacers. The RNA polymerase II expression certainly facilitated to increase and to a greater extend modulate the expression of the gRNA. The expression of multiple gRNAs could be used as well to enhance gene deletion, even in absence of efficient HR, by providing the possibility to induce double cuts around the target gene which could result in a complete gene

deletion even when repaired by NHEJ. Exploring and increasing the efficiency of multiplexed editing ability will be instrumental to further unlock genetic tractability of non-conventional yeasts.

In addition to the challenge of introducing DSBs quickly and efficiently, CRISPR-Cas9-mediated genome engineering in nonconventional yeasts is limited by their intrinsic ability for HRmediated DNA repair, as illustrated in this study and by other previously developed CRISPR tools (Nambu-Nishida et al. 2017; Numamoto et al. 2017; Weninger et al. 2016). While low HR activity can often be compensated for by working in a NHEJ-deficient background, such strains may have undesirable properties, e.g. higher stress sensitivity (Nielsen, Nielsen and Mortensen 2008; Takahashi, Masuda and Koyama 2006) and reduction of cellular fitness (Snoek et al. 2009), which might only emerge after prolonged cultivation. For example, the commonly targeted KU70/80 complex is involved in telomere maintenance, and disruption of the complex results in telomere shortening (Boulton and Jackson 1996), altered position of telomeric DNA in the nucleus (Laroche et al. 1998) and promotes telomere degradation and recombination (Polotnianka, Li and Lustig 1998). As an alternative to inactivating NHEJ, Rad52, a highly conserved protein which plays an important role in HR-mediated DNA repair in yeasts and other organisms (Symington 2002), has been exploited to improve the efficiency of HR. Expression (Di Primio et al. 2005) or protein delivery (Kalvala et al. 2010) of S. cerevisiae derived Rad52 as well as the utilization of a recently described Cas9-ScRad52 fusion protein (Shao et al. 2017) increased HR rates in mammalian cells. However, overexpression of KlRad52 in K. lactis did not have a beneficial effect (Kooistra, Hooykaas and Steensma 2004). Another approach to optimize HR could be cell cycle synchronization and transformation of cells in S-phase, during which cells exhibit the highest ratio of HR over NHEJ (Barlow and Rothstein 2010; Chapman, Taylor and Boulton 2012, Karanam et al. 2012). Cell arrest in S-phase with hydroxyurea indeed led to increased rates of HR in various (non-conventional) veasts (Tsakraklides et al. 2015).

To conclude, the pUDP CRISPR/Cas9 system presented here has been shown to work in four different species of yeasts belonging to the Saccharomycotina, limited by host-dependent targeting efficiency and intrinsic HR capability. We expect the pUDP system to be also applicable for Cas9-mediated genome engineering in other industrially relevant Saccharomycotina yeasts. The approach presented in this study demonstrates the potential of wide host-range tools for genome editing. Although highly efficient genome editing is likely to require species-specific characteristics, the pUDP system can be used for rapid introduction of screenable mutations (e.g. auxotrophic markers) and, as demonstrated for O. parapolymorpha, the elimination of NHEJ. Moreover, due to its broad host range, the pUDP plasmid can serve as a starting point for optimization of Cas9-mediated gene targeting in individual yeast species by promoter and terminator replacement studies.

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Conflict of interest. None declared.

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