CRISPR/Cas system for yeast genome engineering: advances and applications

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CRISPR/Cas system for yeast genome engineering: advances and applications
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Abstract
The methods based on the Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR)/CRISPR-associated (Cas) system have quickly gained popularity for genome editing and transcriptional regulation in many organisms, including yeast. This review aims to provide a comprehensive overview of CRISPR application for different yeast species: from basic principles and genetic design to applications.

Introduction
In 2003, Francisco Mojica and colleagues discovered that the spacer sequences from bacterial Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR) loci match viral and conjugative plasmid sequences and hypothesized that CRISPR must be part of the bacterial immune system (Mojica et al. 2005; Lander 2016). In the following years multiple studies had been performed to unravel the mechanism of CRISPR functionality (Lander 2016) until, in 2012, two research groups managed to re-program the targeting of CRISPR-associated nuclease (Cas9), so Cas9 would introduce double-strand DNA breaks (DSBs) in a sequence-specific manner in vitro (Gasiunas et al. 2012; Jinek et al. 2012). Following this, applications of CRISPR/Cas9 for in vivo genome editing in mammalian cells was published early in 2013 (Cong et al. 2013; Mali et al. 2013), followed by DiCarlo et al. (2013) reporting the usage of the system in the yeast Saccharomyces cerevisiae. Since then the technology has been optimized and adapted for numerous organisms, covering applications from industrial biotechnology (van Erp et al. 2015) to plant breeding (Bortesi and Fischer 2015) and treatment of human diseases (Cai et al. 2016).

In native type II CRISPR/Cas systems, Cas9 is guided to the target DNA region by a two-RNA molecule hybrid consisting of CRISPR RNA (crRNA) and trans-activating crRNA (tracrRNA). Together with the tracrRNA, crRNA forms a secondary structure loop, which recruits Cas9. The crRNA guides the system to a genomic target of ~20 bp through base-pairing with the

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complementary DNA strand. The particular genomic target must be followed by the protospacer adjacent motif (PAM) NGG. The Cas9 nuclease domain HNH then cleaves the DNA strand complementary to the crRNA-guide sequence, while RuvC-like domain cleaves the other DNA strand, thus resulting in a double-strand DNA break. The DNA cleavage is performed three nucleotides upstream of the PAM site (Gasiunas et al. 2012). For easier use in genome editing, the crRNA and tracrRNA can be fused tail-to-head via a linker into a single guiding RNA (gRNA) (Jinek et al. 2012).

This review covers the technical details of the implementation of CRISPR/Cas-mediated genome editing in various yeast species, transcriptional regulation via the enzymatically inactive “dead” dCas9, which binds but does not cut the DNA target (Jinek et al. 2012), and presents examples of applying the technology for engineering of yeast cell factories.

**CRISPR/Cas9 genome editing in yeasts**

When Cas9 protein and gRNA are expressed in yeast cells, Cas9 introduces DSBs that must be repaired by the cells via non-homologous end joining (NHEJ) or homologous recombination (HR) (Liu et al. 2017). By supplying a DNA repair template for use in HR, various DNA modifications can be obtained. In the case of efficient cutting, the generated DSBs serve as a negative selection. Thus there is no need for using a selective marker as in non-CRISPR genome editing methods. Relatively precise and flexible targeting and elimination of the need for positive selection are the two key advantages of the CRISPR/Cas9 technology for yeast genome engineering. The method also allows engineering of diploid and polyploid industrial strains (Ryan et al. 2014; Zhang et al. 2014; Stovicek, Borodina and Forster 2015), which are challenging to manipulate genetically due to the difficulties with modifying multiple alleles and due to the lack of selection markers (Le Borgne 2012). Additionally, by combining several gRNAs, multiple sites can be targeted simultaneously allowing the unprecedented speed of multiple genetic edits (Ryan et al. 2014; Bao et al. 2015; Jakočiūnas et al. 2015a). On the downside of CRISPR/Cas9, there is a considerable variation in efficiency when targeting different loci, perhaps due to a positional effect of the target region (Smith et al. 2016). At the moment, there also seems to be an upper limit for the number of edits (up to six) that can be introduced simultaneously as every additional introduced DSB decreases the overall yield of surviving clones (Jakočiūnas et al. 2015a; Mans et al. 2015). Furthermore, CRISPR/Cas9 multiplexing still represents a significant increase in workload for finding correct clones.
Progress has been made on adapting the type II CRISPR/Cas system, described in *Streptococcus pyogenes* (Chylinski et al. 2014), to various yeast species – *S. cerevisiae* (Jakočiūnas, Jensen and Keasling 2016), *Yarrowia lipolytica* (Schwartz et al. 2016a), *Komagataella phaffii* (formerly *Pichia pastoris*) (Weninger et al. 2016), *Kluyveromyces lactis* (Horwitz et al. 2015), *Schizosaccharomyces pombe* (Jacobs et al. 2014), and the pathogenic yeast species *Candida albicans* (Vyas, Barrasa and Fink 2015) and *Cryptococcus neoformans* (Wang et al. 2016). We first discuss the design of the targeting gRNA sequence as a critical aspect of all CRISPR/Cas9 applications. As the vast majority of the studies describing CRISPR/Cas9 genome editing in yeasts have focused on *S. cerevisiae*, the larger section dedicated to this model organism also details some of the more general issues related to the Cas9-mediated genome engineering. For clarity, the studies focusing on the other yeasts are discussed in a separate section.

**Computational tools for gRNA design in yeast**

Any ~20-bp sequence proximal to the PAM site in the genome can serve as the gRNA targeting sequence. The rationale behind careful gRNA selection is to minimize the risk of Cas9-mediated cleavage at unwanted sites in the genome (off-target effects) and maximize the cutting efficiency at the selected site (on-target activity). Other factors may outweigh the best parameters and put additional constraints on the design, e.g. position of a target proximal to the beginning of the ORF for generating premature STOP codons or requirement of a target location in promoter/5' UTR region in case of gene repression/activation experiments (Mohr et al. 2016). Several web-based tools have been developed to facilitate and automatize the design of gRNA targets (Table 1). Such tools aim mainly at providing guide sequences that minimize the likelihood of off-target effects, matching all possible targets within the given parameters against the reference genome. Some tools provide a list of targets with specified number of mismatches within the entire target sequence or the “seed” sequence (8–12bp adjacent to the PAM site) (CRISPy, Jakočiūnas et al. 2015a; Ronda et al. 2014; CRISPRdirect, Naito et al. 2015), filter out sequences with potential off-target effects (Yeastriction, Mans et al. 2015) or introduce a specificity score based on number of mismatches within the target sequence and rank the targets accordingly (CRISPR-ERA, Liu et al. 2015; Benchling, ATUM gRNA design). CHOPCHOP (Labun et al. 2016) or E-CRISP (Heigwer, Kerr and Boutros 2014) provide the possibility for user-defined parameters of the off-target evaluation. Even though off-target effects are considered unlikely in such a small genome as yeast (Ryan et al. 2014; Jakočiūnas et al. 2015a), it is advisable to double check the design using yet another tool to avoid introduction of any undesired modifications. Although several potential requirements for
gRNA design have been suggested to ensure efficient generation of DSB at the target site, it is still not easy to establish a set of golden rules that would guarantee a success until more experimental data has been acquired. Some of the tools highlight simple features that might influence gRNA efficiency, such as poly T presence in the sequence, GC content (CRISPRdirect) (Naito et al. 2015), AT content or self-complementarity of a gRNA molecule and provide a score based on these parameters (Yeastriction, E-CRISP, CRISPR-ERA) (Heigwer, Kerr and Boutros 2014; Liu et al. 2015; Mans et al. 2015). Other tools such as Benchling have implemented more sophisticated efficiency scores based on an experimental evaluation of a large set of mammalian gRNAs and their sequence features (Doench et al. 2014; Xu et al. 2015; Doench et al. 2016). In some cases, users can even choose from several different algorithms of the on-target evaluation (CHOPCHOP, E-CRISP). A few tools also include information on the presence of a specific restriction site in the target sequence (CHOPCHOP, CRISPRdirect, Yeastriction) that might facilitate downstream validation of the cloned target molecule (Mans et al. 2015). CRISPR-ERA or E-CRISP also facilitate designing of a gRNA molecule for engineering applications other than genome editing, e.g. gene repression or gene activation applications. While some of the tools support only one yeast genome, typically *S. cerevisiae* reference genome, others provide gRNA design option for several yeast species or various strains of *S. cerevisiae* (Table 1). The CRISPy tool web server implementation, CRISPy-web (Blin et al. 2016), allows for user-upload of any GenBank format genome. CRISPRdirect is being frequently updated with new genomes, and CHOPCHOP offers an upload of new genomes on request.

**CRISPR/Cas9 and genome editing in *Saccharomyces cerevisiae***

*S. cerevisiae* is an important eukaryotic model organism and also a widely used industrial host for production of fuels, chemicals and recombinant proteins (Borodina and Nielsen 2014; Li and Borodina 2015). Thanks to its excellent HR capability, *S. cerevisiae* is relatively easy to engineer genetically. Below we discuss the ways for delivering Cas9, gRNA and DNA repair templates to *S. cerevisiae* (summarized in Table 2).

**Cas9 expression**

The most commonly used *Cas9* gene variant in *S. cerevisiae* has been *Cas9* from *S. pyogenes*, fused with a nucleolar localization sequence. The DNA sequence of *Cas9* can be either native (Ryan et al. 2014; Bao et al. 2015), human codon-optimized (DiCarlo et al. 2013; Gao and Zhao 2014; Zhang et al. 2014; Jakočiūnas et al. 2015a; Laughery et al. 2015; Mans et al. 2015; Stovicek, Borodina and
Forster 2015) or yeast codon-optimized (Horwitz et al. 2015; Generoso et al. 2016) (Table 2). Only Xu et al. (2015) reported the use of Streptococcus thermophilus CRISPR3 loci-encoded Cas9 (recognizing a different PAM site), albeit with much lower engineering efficiency. The Cas9 gene was most commonly expressed under the control of constitutive promoters of different strengths from self-replicating low-copy centromeric vectors (DiCarlo et al. 2013; Zhang et al. 2014; Jakočiūnas et al. 2015a; Stovicék, Borodina and Forster 2015) or high-copy 2µ vectors (Gao and Zhao 2014; Ryan et al. 2014; Bao et al. 2015; Laughery et al. 2015; Generoso et al. 2016). Horwitz et al. (2015) or integrated into the genome (Mans et al. 2015) (Table 2). Expression of Cas9 on a high-copy vector from a strong constitutive promoter led to a negative influence on the growth of some yeast strains (Ryan et al. 2014; Generoso et al. 2016). However, this problem was not observed in other studies that used the same mode of Cas9 expression (Gao and Zhao 2014; Bao et al. 2015; Laughery et al. 2015). The toxicity of Cas9 nuclease could be avoided by using weaker promoters for Cas9 expression (Ryan et al. 2014; Generoso et al. 2016). Overall, the form of Cas9 expression does not seem to be a critical parameter in CRISPR/Cas9 engineering strategies for S. cerevisiae.

Guide RNA expression

Design, expression, and delivery of the gRNA components are crucial parameters for successful CRISPR/Cas9 engineering. In S. cerevisiae, the most common strategy has been to express a chimeric gRNA molecule from a high-copy vector to ensure its abundant expression (Table 2). Both ends of the gRNA molecule must be precisely defined to create a functional Cas9/gRNA complex. Functional gRNA transcription has been achieved using a) an RNA polymerase III (Pol III) promoter that provides a transcript with a leader sequence cleaved during molecule maturation (DiCarlo et al. 2013; Farzadfard, Perli and Lu 2013), b) Pol III promoters containing cis-regulatory elements within the mature RNA molecule (tRNA) combined with a ribozyme, cleaving the transcript on its 5´ end (Ryan et al. 2014), and c) an RNA polymerase II (Pol II) promoter, if the gRNA molecule is flanked with two ribozymes executing cleavage on both ends of the molecule (Gao and Zhao 2014) (Fig. 1). Besides the chimeric gRNA approach, separate expression of a targeting crRNA array driven by a Pol III promoter, processed by native RNA processing enzymes, and tracrRNA transcribed from another Pol III promoter has been reported (Bao et al. 2015). The expression cassette containing SNR52 promoter and SUP4 terminator, an approach shown to produce prokaryotic tRNA molecules in yeast (Wang and Wang 2008), was successfully used for targeting a single gene in haploid or diploid laboratory strains with engineering efficiencies
reaching 100% (DiCarlo et al. 2013; Horwitz et al. 2015; Jakočiūnas et al. 2015a; Laughery et al. 2015; Mans et al. 2015, 2015; Generoso et al. 2016) (Table 2). It is important to mention that engineering efficiencies discussed in this review are defined as the number of clones with the desired genomic edit per number of clones surviving after the transformation. Such value should not be mistaken with transformation efficiency values used traditionally in non-CRISPR engineering studies as these relate to the number of viable cells in the transformation reaction (Storici et al. 2003; Alexander, Doering and Hittinger 2014). Although some studies also provide transformation efficiency values that reflect the number of cells not surviving the transformation (DiCarlo et al. 2013; Stovicek, Borodina and Forster 2015), many others do not, leaving the engineering efficiency as the only relevant benchmark. The SNR52 promoter/SUP4 terminator setup also allowed for gene deletion in various diploid industrial strains with efficiencies ranging between 65-78% (Stovicek, Borodina and Forster 2015) and even polyploid strains with 15-60% efficiency (Zhang et al. 2014) (Table 2). The expression of a gRNA fused to a Hepatitis delta virus (HDV) ribozyme controlled by a tRNA promoter and SNR52 terminator led to almost 100% gene deletion efficiency in a diploid laboratory strain and more than 90% in a polyploid industrial strain (Ryan et al. 2014). As for the third mentioned approach, a gRNA molecule flanked with Hammerhead (HH) and HDV ribozymes on the 5´ and 3´ end, respectively expressed from ADH1 promoter also enabled efficient gene disruption in a laboratory strain (Gao and Zhao 2014). The crRNA array method achieved efficiencies of 76%-100% in a laboratory strain after several days-outgrowth of the transformed cells (Bao et al. 2015).

When a researcher decides to engineer a targeted genomic locus, only the ~20 bp recognition sequence of a gRNA molecule needs to be modified to redirect the Cas9/gRNA complex to a particular target site. Several ways of obtaining an expression vector with a customized gRNA molecule have been described (Fig. 2). Several studies exchanged the recognition sequence of a gRNA vector using whole vector amplification with primers containing a new target-specific 20 bp-region. Vector circularization was achieved via PCR with a phosphorylated primer, followed by ligation (Jakočiūnas et al. 2015a; Stovicek, Borodina and Forster 2015), in vivo in yeast or in vitro Gibson assembly using two oligos overlapping at the target sequence (Generoso et al. 2016), or via restriction-free cloning (van den Ent and Löwe 2006) with two 60 bp-complementary oligos containing a target sequence (Ryan and Cate 2014). In other studies, two target-specific complementary oligos containing sequences overlapping with the gRNA cassette were cloned into a vector using Gibson assembly (Reider Apel et al. 2016) or restriction sites located between
promoter and the gRNA structural part (Laughery et al. 2015; Lee et al. 2015), or transformed directly into yeast along with the digested expression vector (Mans et al. 2015). Alternatively, the gRNA cassette was amplified using two-step fusion PCR and cloned via Gibson assembly (DiCarlo et al. 2013), standard restriction cloning (Chin et al. 2016) or transformed along with a digested expression vector for in vivo vector gap repair in yeast (Horwitz et al. 2015). To omit the PCR amplification step, customized gRNA cassettes can be synthesized as gene blocks and integrated into a vector via restriction cloning (Zhang et al. 2014) or USER assembly (Jakočiūnas et al. 2015b; Ronda et al. 2015). Lastly, Golden gate cloning of synthetic parts of the crRNA array has also been shown (Bao et al. 2015).

In summary, researchers can choose from a number of cloning systems for generation of a target gRNA molecule and can also benefit from online tools facilitating the particular cloning design (Laughery et al. 2015; Mans et al. 2015) or detailed (Ryan, Poddar and Cate 2016) and straightforward protocols (Jakočiūnas et al. 2015a). However, even in its simplest version, the CRISPR/Cas9 engineering relies on a gRNA vector construction, which can be laborious and costly. The gap repair approach developed by Horwitz et al. (2015) skips the cloning step. However, it requires longer DSB repair templates, high efficiency of HR in the strain, and may result in a non-equimolar expression of the gRNAs when multiplexing. A lower efficiency of engineering with vectors based on in vivo assembly has been documented (Mans et al. 2015; Generoso et al. 2016).

One can also choose to express Cas9 and gRNA from a single vector (Ryan et al. 2014; Bao et al. 2015; Laughery et al. 2015; Generoso et al. 2016). However, due to the large size of the Cas9 gene, generation of a gRNA via the whole plasmid PCR amplification (Ryan and Cate 2014) might be difficult. Such a system is also not compatible with the gap repair gRNA generation as this one requires expression of Cas9 prior the transformation with the gRNA vector (Walter, Chandran and Horwitz 2016).

**Multiplexing gRNA expression**

In *S. cerevisiae*, efficient HR system allows creating multiple genomic changes simultaneously using CRISPR/Cas9. For each genome edit, an individual gRNA must be expressed and a repair template delivered into the cells. The multiple gRNA expression has been achieved using a) several vectors with different selection markers containing up to two different gRNA expression cassettes (Mans et al. 2015), b) a single expression vector carrying several gRNA cassettes (Ryan et al. 2014; Jakočiūnas et al. 2015a; Lee et al. 2015), c) an array of different interspaced crRNAs (Bao et al.
2015), or d) different linear gRNA expression cassettes transformed along with a single gapped expression vector (Horwitz et al. 2015). When up to three different vectors, each carrying two gRNA expression cassettes were transformed, 100%, 70% and 65% efficiency of two, four or six gene deletions was achieved, respectively (Mans et al. 2015). The expression of five individual gRNAs from one vector provided target efficiencies ranging between 50-100% (Jakočiūnas et al. 2015a). Ryan et al. (2014) reported successful gene deletion of two or three genes with efficiencies of 86% and 81% in haploid and 43% and 19% in diploid strains using HDV-gRNA expression cassettes in a single expression vector, respectively. Cloning of crRNA arrays targeting three different genes achieved engineering efficiencies ranging between 27-100% (Bao et al. 2015). The gap repair approach using the transformation of three different gRNA cassettes and a single open vector enabled recovery of 64% positive three-gene deletion mutants (Horwitz et al. 2015) (Table 2).

As described above, all setups enabled successful marker-free multiplexed genome editing in S. cerevisiae. However, despite the reported encouraging results, yeast strains can differ in engineering efficiencies given rather by their nature than differences in the described procedures. Diploid or polyploid industrial strains can be especially difficult to engineer (Zhang et al. 2014; Stovicek, Borodina and Forster 2015; Generoso et al. 2016), and multiplexing can create more work on the other end to identify the correct clones. The CRISPR/Cas9 system also greatly facilitates the sequential introduction of multiple genomic edits. For repeated rounds of editing, the strain is cultivated in the absence of selection pressure for gRNA vector, while maintaining selection pressure for Cas9 vector. Then a new gRNA vector can be introduced to accomplish the next round of genetic modifications. In the final strain, both vectors can be removed in the absence of selection pressure to generate a strain free of selection markers (Stovicek, Borodina and Forster 2015; Jessop- Fabre et al. 2016).

**DNA repair templates**

As mentioned above, the dominant mode of DSB repair in S. cerevisiae is HR when a homologous donor template is available. NHEJ response in S. cerevisiae provides unpredictable results at target sites and severely decreases overall yield of surviving cells (DiCarlo et al. 2013; Mans et al. 2015; Stovicek, Borodina and Forster 2015). It has been shown that short single strand (Generoso et al. 2016) or double strand DNA donor oligos (DiCarlo et al. 2013) sharing homology with a target site can serve as the simplest repair template. The donor oligo can be of various lengths, ranging between 80-120 bp, and can introduce various changes such as a premature STOP codon (DiCarlo...
et al. 2013), a heterologous disrupting sequence (Horwitz et al. 2015), a barcode (Ryan et al. 2014) for easier genotyping, or an entire ORF deletion (Mans et al. 2015) (Fig. 1). The PAM site should always be removed from the donor sequence to prevent the cutting by Cas9 (DiCarlo et al. 2013). The repair template can also be delivered as a part of an expression vector (Bao et al. 2015; Garst et al. 2016). Longer gene expression cassettes with at least 40-bp homology to the target site can also be used as repair templates. They enable integration of larger DNA fragments, e.g., carrying gene expression cassettes (DiCarlo et al. 2013; Stovicek, Borodina and Forster 2015). The CRISPR/Cas9 approach has also been combined with in vivo assembly of several overlapping DNA parts (Fig. 1). Ryan et al. (2014) reported 70-85% efficiency for assembly of a gene expression cassette consisting of three parts with 50bp overlaps to a targeted locus in a diploid or polyploid strain. In another study, one transformation event enabled integration of six overlapping gene expression cassettes into a single gene locus while another gene was deleted using a short oligo simultaneously (Mans et al. 2015). A multiplex approach, CasEMBLR demonstrated assembly of five overlapping DNA parts per locus in up to three different loci simultaneously with an efficiency ranging between 30-97% (Jakočiūnas et al. 2015b). A metabolic pathway consisting of eleven genes on six DNA parts flanked with 500bp arms homologous to three independent loci was used as a DSB repair template in the gap-repair gRNA delivery approach resulting in 4% efficiency of the pathway assembly (Horwitz et al. 2015). Tsai et al. (2015) integrated two copies of a multigene pathway consisting of six genes on four DNA parts with 300bp homologous arms into two different gene loci with 25-100% efficiency. As the integration of a gene expression cassette into an ORF may influence expression of the heterologous gene (Stovicek, Borodina and Forster 2015), several toolkits targeting intergenic regions providing reliable level of gene expression have been developed. Three-DNA part gene expression cassettes with 1 kbp homologous arms used as a donor template resulted in 40%-95% integration efficiency depending on the particular site targeted (Reider Apel et al. 2016). Although versatile, in vivo CRISPR/Cas9-mediated assembly requires tedious multiplex genotyping. Thus, preassembly of donor templates with sufficiently long homologous arms might be an alternative option to omit this. Using the system developed by Bao et al. (2015), preassembled large metabolic pathways were integrated into transposable Ty elements in multiple copies with efficiencies more than 80% (Shi et al. 2016). Ronda et al. (2015) targeted multiple validated intergenic loci with preassembled gene expression cassettes reaching efficiencies of 84% with three simultaneous integrations. When using the marker-free variant of previously designed integrative vectors (Jensen et al. 2014; Stovicek et al. 2015) targeting intergenic loci, integration of
up to six heterologous genes was achieved with 70% efficiency (Jessop- Fabre et al. 2016). In summary, due to the high efficiency of HR, large pathways can be assembled directly in vivo omitting in vitro cloning steps. However, the preassembly of donor DNA fragments always leads to higher integration efficiencies and does not require subsequent extensive genotyping.

Taken together, mainly linear DNA fragments of different length have been successfully used for efficient DSB repair. However, a recent study demonstrated that episomal vectors that contain both a gRNA expression cassette and a DNA repair template could also be used in the yeast S. cerevisiae (Garst et al. 2016). In a proof-of-concept experiment, ADE2 gene was mutated with 95% efficiency in a laboratory strain and with 70% efficiency in a wine strain. A particular advantage of this method is that the combined DNA elements, which contain a gRNA and a corresponding repair template, are small enough (~200 bp) to be synthesized by high-throughput oligomer synthesis on arrays. Combined with a high transformation efficiency of episomal vectors into yeast, this enables generation of large strain libraries.

Direct DNA editing using CRISPR-cytidine deaminase fusion

A method for CRISPR-based targeted DNA mutagenesis was described by taking advantage of an activation-induced cytidine deaminase (AID), which is normally responsible for somatic hypermutation of the variable regions of antibodies (Nishida et al. 2016). When AID was expressed as a fusion with dCas9 in S. cerevisiae, AID deaminated deoxycytidine to deoxyuridine 15-19 bases upstream of the PAM sequence on the non-complementary strand to gRNA, effectively creating C→G/T point mutations. The efficiency of gene inactivation using this approach was 16-47%, depending on the chosen target site. The advantage of the CRISPR-AID method is a reduced toxicity in comparison to the nuclease-based CRISPR approaches (Nishida et al. 2016).

CRISPR/Cas9 genome editing in different yeast species

Kluyveromyces lactis

K. lactis is used industrially for the production of recombinant proteins, fermented dairy products, and some metabolites (Spohner et al. 2016). Horwitz et al. (2015) demonstrated CRISPR/Cas9 genome editing in an industrial strain of K. lactis. The 2µ element in the expression vector was exchanged for the pKD1 vector-stabilizing element. To decrease the NHEJ activity in K. lactis, the authors deleted YKU80 gene. Although with low efficiency (2.3%), the method allowed integration of three six-gene-DNA parts into three individual chromosomal loci (Horwitz et al. 2015).

Yarrowia lipolytica
*Y. lipolytica* is the most studied oleaginous yeast and is applied in the biotechnology industry for the production of lipase, citric acid, lactone fragrances, and recently also ω-3-fatty acids (Thevenieau, Nicaud and Gaillardin 2009; Xue *et al.* 2013). Several recent studies have demonstrated the potential of the CRISPR/Cas9 system in this yeast. Schwartz *et al.* (2016) constructed *Yarrowia* codon-optimized *Cas9* and hybrid SCR1′-tRNA promoter for gRNA expression on a centromeric vector (Schwartz *et al.* 2016a). It enabled efficient NHEJ-generated gene deletions. More than 50% or 90% of the cells acquired a gene deletion after two or four-day outgrowth of the transformed cells, respectively. HR-mediated gene deletions with a donor fragment with 1-kbp homologous arms were also obtained with high efficiency. The HR-mediated repair was pronounced in KU70 mutant, lacking NHEJ-mediated response (Schwartz *et al.* 2016a). A possibility of multiplex gene deletion in *Y. lipolytica* was also demonstrated (Gao *et al.* 2016). Here a vector was designed to carry *Yarrowia* codon-optimized Cas9 gene driven by the strong, endogenous TEF1 promoter, and also gRNAs flanked with the HH and HDV ribozymes expressed from the TEF1 promoter. In the absence of donor DNA, NHEJ-mediated gene nonsense mutations occurred with efficiencies of 85%, 36% or 19% for one, two or three targeted genes, respectively, after four days of outgrowth of the transformed cells. Furthermore, HR-mediated gene disruption was shown when the donor template was delivered on the Cas9/gRNA vector, with higher rates in KU70/80 mutants (Gao *et al.* 2016). CRISPR/Cas9 also allowed the development of a toolkit for integration of donor cassettes which were delivered into the cells by a separate replicative vector requiring an additional selection during the transformation (Schwartz *et al.* 2016b). In an NHEJ-positive strain, five out of 17 tested locations were targeted with integration efficiencies from 48-69%, while three sites showed less than 6% and the remaining nine sites did not show any positive integration. Sequential marker-less integration of a metabolic pathway into the described loci was shown (Schwartz *et al.* 2016b).

*Komagataella phaffii* (formerly *Pichia pastoris*)

*Ko. phaffii* (*P. pastoris*) is an important recombinant protein producer due to its excellent folding and secretion capability. However, it is poor in HR, which makes it very hard to engineer. Weninger *et al.* (2016) extensively tested different modes of expression of the Cas9 gene and gRNA molecules. Use of a low copy ARS element vector with bi-directional native HXT1 promoter driving the expression of human codon-optimized Cas9 and HH/HDV-ribozyme-flanked gRNA transcript resulted in up to 90% of single gene non-sense mutations. When two genes were targeted, non-sense mutations in both ORFs were observed with a frequency of 69%. Although a donor
template with 1-kbp homologous arms was provided, only very low integration efficiency (2%) occurred suggesting that NHEJ remained the dominant way of DSB repair (Weninger et al. 2016).

_Schizosaccharomyces pombe_

The fission yeast _Sch. pombe_ is an important model organism for the study of eukaryotic cellular biology and in particularly cell cycle regulation (Hoffman, Wood and Fantes 2015). Jacobs _et al._ (2014) used _rrk1_ promoter for expression of gRNA molecule as it provides a defined 5′-leader, cleaved during maturation. The 3′-end of the gRNA molecule was fused to the HH ribozyme, as _rrk1_ is a Pol II promoter resulting in polyadenylation of mature RNAs. Expression of gRNA and Cas9 separately on two low-copy ARS-containing vectors (or together on one vector to minimize the observed negative influence of Cas9 expression on cell growth) led to the 85-98% efficiency of the target modification when a PCR-amplified mutated allele was used as donor template (Jacobs _et al._ 2014). A similar system enabled construction of a single gene deletion with 33% efficiency (Fernandez and Berro 2016).

_Pathogenic yeasts_

Targeted gene deletions are necessary for the study of gene functions in virulence models. In the most prevalent yeast pathogen – _C. albicans_ – the absence of haploid state and frequent aneuploidy of clinical isolates makes gene deletions very tedious. In the absence of autonomously replicating vectors, CRISPR/Cas9 was implemented via integrating Cas9 controlled by _ENO1_ promoter and gRNA expressed from _SNR52_ promoter into _C. albicans_ genome (Vyas, Barrasa and Fink 2015). The _Cas9_ gene was codon-optimized for CTG clade yeasts. In “solo” approach, gRNA expression cassette was integrated into a strain already expressing Cas9. In the “duet” approach, both expression cassettes were integrated in a single transformation. Both “solo” and “duet” systems resulted in an acceptable gene deletion efficiency of 60-80% and 20-40%, respectively. The more efficient “solo” system was then used for generation of deletions in several genes or deletion of two homologous genes with a single targeting gRNA molecule. Moreover, successful non-sense mutations in three different loci combining the solo and duet system for delivery of two different gRNA cassettes were documented (Vyas, Barrasa and Fink 2015). A possibility of transient expression of linear cassettes carrying both components was shown. A single gene was replaced with a linear marker gene cassette reaching more than 50% efficiency while the linear gRNA and _Cas9_ cassettes were lost at the same time (Min _et al._ 2016).
Another pathogenic yeast *Cr. neoformans* exhibits a low rate of HR that hampers its manipulation and thus functional gene analysis. Two studies have demonstrated the CRISPR/Cas9 system capacity to generate non-sense mutations and to stimulate HR response in different serotypes of *Cr. neoformans* (Arras et al. 2016; Wang et al. 2016). As circular molecules are not stable in *Cr. neoformans*, linear DNA vectors were used for expression of Cas9 nuclease and gRNAs. gRNAs were expressed from CnU6 promoter and terminated by 6T terminator (Wang et al. 2016). Alternatively, the Cas9 gene was integrated into the genome and a linear vector was used for expression of a gRNA molecule flanked with HH and HDV ribozymes from a Pol II promoter (Arras et al. 2016). The introduction of non-sense mutations was achieved without donor DNA with efficiency above 80%. Mutated allele used as a donor template resulted in HR-mediated allele exchange when selecting for a particular phenotype. Full removal of an ORF occurred with frequencies of 20-90% when a donor marker gene was fused to the Cas9/gRNA cassette followed by spontaneous loss of the Cas9/gRNA part eliminating thus the persistence of the CRISPR/Cas9 system (Wang et al. 2016). Gene deletions were obtained in different serotypes of *Cr. neoformans* by using a marker cassette with homologous arms to the given ORF. Stimulation of HR led to 70% success rate for obtaining the mutants (Arras et al. 2016).

Another pathogenic yeast with a dominant NHEJ pathway, *Candida glabrata* was demonstrated to be amenable to the CRISPR/Cas9-mediated engineering (Enkler et al. 2016). Here two centromeric vectors carrying Cas9 and gRNA expression cassette were used. Although adoption of *S. cerevisiae* system (DiCarlo et al. 2013) for expression of a gRNA appeared to be feasible, specific *C. glabrata* adjustments (RNAH promoter, tRNA terminator) led to better performance of the system (Enkler et al. 2016). Besides efficient generation of indels by NHEJ, deletion of a reporter gene using a donor marker cassette with relatively short homologous arms was achieved with increased HR rates (Enkler et al. 2016).

**Transcriptional regulation via CRISPR**

Targeted regulation of gene expression is important both in the context of metabolic engineering and functional genomics. The CRISPR method has been adapted both for activation and repression of gene transcription in *S. cerevisiae*, but so far not in other yeast species.

Qi et al. (2013) generated an enzymatically inactive variant of Cas9 by mutation of both nuclease sites (D10A and H840A) and showed that this null-nuclease dCas9 when targeted to a coding region of a gene, caused transcriptional repression in *E. coli*. In this approach, termed CRISPR interference (CRISPRi), dCas9 sterically blocks the binding and action of RNA polymerase. In a
follow-up study, dCas9 was guided to a promoter region, resulting in efficient gene repression in *S. cerevisiae* (Gilbert *et al.* 2013). The repression could be further enhanced by fusing a repressor domain to dCas9 (Fig. 3). GFP fluorescence was reduced 18-fold when the *TEF1* promoter driving the GFP expression was targeted by dCas9, and the fluorescence decreased 53-fold when the same region was targeted by dCas9 fused to a mammalian transcriptional repressor domain Mxi1 (Gilbert *et al.* 2013).

Farzadfard, Perli, and Lu (2013) fused dCas9 to an activator domain (VP64) instead. The resulting chimeric protein could both repress and activate gene expression depending on the targeting site in the promoter region. When dCas9-VP64 was targeted to the region upstream the TATA box of the minimal *CYC1m* promoter, the promoter was activated. Targeting the sites immediately adjacent to the TATA box or Transcriptional Start Site (TSS) repressed the expression from *P_{CYC1m}*. The obtained activation level was not very high, max 2.5-fold. To achieve a higher level of activation, the authors created a synthetic promoter by arraying multiple operators upstream the *P_{CYC1m}*. The activation level increased proportionally to the number of operators, reaching 70-fold activation for 12 operators (Farzadfard, Perli and Lu 2013).

Fusion of dCas9 to a tripartite activator (VPR) composed of three strong activation domains (VP64, p65, and Rta) resulted in 38 and 78-fold activation of promoters *P_{HED1}* and *P_{GAL7}*, respectively. Fusion of dCas9 with VP64 only gave 9 and 14-fold activation of the same promoters (Chavez *et al.* 2015).

Zalatan *et al.* (2015) undertook a different approach to achieve targeted up- and down-regulation. Instead of fusing activation or repression domains to dCas9, they included effector protein recruitment domains into the guide RNA (Fig. 3). In the same strain, they expressed dCas9 and regulation proteins, fused with RNA-binding domains. They termed the resulting gRNAs with protein recruitment capabilities “scaffold RNA” (scRNA). Gene activation using scRNA binding VP64 activation domain was 20-50-fold, much higher than the activation achieved with dCas9-VP64 fusion. Several hairpins could be combined in a single scRNA, which allowed amplification of activation or combination of repression and activation of different sites (Zalatan *et al.* 2015).

In addition to the studies mentioned above focusing on the on/off states of gene expression, grade modulation of gene expression using dCas9 fused to either an activation or repression domain was shown (Deaner and Alper 2017). This was achieved by changing the gRNA target location and thus recruiting the dCas9-activator/repressor complex to different positions in gene promoters. It resulted in a dynamic range of gene expression from almost silenced gene to its several ten-fold
overexpression related to the proximity of the dCas9-based regulators to the core of the promoter. The graded gene expression enabled tuning of metabolic pathways and optimization of the desired phenotypes in several metabolic engineering applications (Deaner and Alper 2017).

**Application of CRISPR/Cas9 for engineering of yeast cell factories**

CRISPR/Cas genome editing and transcriptional regulation are particularly suitable for developing yeast cell factories. As the strain development usually proceeds through iterative design-build-test cycles, the CRISPR technology facilitates this process because the strains can repeatedly be edited in a flexible multiplex way. As far as transcriptional regulation is concerned, CRISPR also enables relatively easy multiplexing. We will illustrate this with four brief examples (Fig. 4).

Shi *et al.* (2016) applied CRISPR/Cas9 to engineer *S. cerevisiae* towards production of a non-native product \((R,R)\)-2,3-butanediol (BDO) from a non-native substrate xylose in a single transformation step. A 24-kb integration construct consisting of six gene expression cassettes (three for the xylose consumption pathway and three for the BDO biosynthesis pathway) was integrated into the delta sequence of the Ty transposon elements. Introducing Cas9-mediated DSBs at the delta sites allowed the integration of 10 copies of the 24-kb DNA fragment. A higher copy number of the pathways resulted in both higher xylose consumption rate and higher BDO production, where 0.31 g/L of BDO was produced from 20 g/L xylose (Shi *et al.* 2016).

Stovicek *et al.* (2015) engineered diploid industrial *S. cerevisiae* strain Ethanol Red, used in many first generation ethanol plants, to produce lactic acid by replacing both alleles of pyruvate decarboxylase genes *PDC1* and *PDC5* with L-lactate dehydrogenase encoding gene (*ldhL*) from *Lactobacillus plantarum*. The genetic modification was accomplished in a single transformation event, leading to a strain producing 2.5 g/L lactic acid with the yield of 0.49 g of lactic acid/g of glucose (Stovicek, Borodina and Forster 2015).

Transcriptional regulation via CRISPRi was demonstrated for the production of the bacterial pigment violacein in *S. cerevisiae*. Here CRISPR RNA scaffolds were used to recruit transcriptional activators and repressors, alone or simultaneously, to a promoter site, which allowed tight control of transcriptional activation and repression. By simply changing the RNA scaffolds, the same strain could be reprogrammed to produce different ratios of the pathway products, deoxyviolacein, violacein, prodeoxyviolacein, and proviolacein. Combining these RNA-encoded circuits with conditional expression of Cas9, a system for switching from growth to production phase was obtained (Zalatan *et al.* 2015).
Recently, a combination of Cas9 genome editing and dCas9 transcriptional regulation was demonstrated by engineering *S. cerevisiae* for production of flavonoid precursor naringenin. First, Cas9 was used for integration of a multigene pathway into an intergenic locus leading to production of naringenin from phenylalanine. Next, the naringenin production was increased through dCas9-mediated downregulation of an essential gene TSC13 to prevent the formation of by-product phloretic acid (Vanegas, Lehka and Mortensen 2017).

**Outlook**

This review summarizes the recent developments of CRISPR-based systems for genome editing and transcriptional regulation in various yeast species. The CRISPR/Cas9 technology has advantages over conventional marker-based genome editing in several aspects. It enables fast strain engineering of prototrophic wild and industrial yeast strains. Further, it allows performing multiple genome edits simultaneously and is independent of marker cassette integration. For transcriptional regulation, the CRISPR offers an advantage of relatively easy design & implementation, the possibility of multiplexing, and orthogonality. However, to enable the wide adaptation of CRISPR, the current limitations need to be addressed. These include: (i) design of efficient and specific targeting for different yeast species, (ii) elimination of cloning necessity, (iii) enabling large-scale multiplexing, and, finally, (iv) resolving the IP issues. The uncertainty about the ownership of the CRISPR technology delays its adaptation for industrial biotechnology and pharmaceutical applications and must be resolved as soon as possible so the technology can unfold its true potential.

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**References**


Ryan OW, Skerker JM, Maurer MJ et al. Selection of chromosomal DNA libraries using a multiplex CRISPR system. *eLife* 2014;3:e03703.


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Figure 2: Generation of specific gRNA expression cassettes. A. Vector can be circularized via ligation (one oligo phosphorylated) (Jakočiūnas et al. 2015a; Stovicek, Borodina and Forster 2015), ligation-free primer extension reaction (Tsai et al. 2015; Ryan, Poddar and Cate 2016), Gibson assembly or recombination in vivo (pair of oligos overlapping at the specific gRNA target sequence) (Generoso et al. 2016). B. Short synthetic oligos are cloned via e.g. Gibson assembly (oligos with overhangs homologous to the ends of the digested vector) (Reider Apel et al. 2016), restriction cloning (oligos with overhangs complementary to a particular restriction site) (Laughery et al. 2015), modular cloning (seamless assembly using type IIS restriction enzymes, oligos with overhangs complementary to a particular restriction site) (Lee et al. 2015; Vyas, Barrasa and Fink 2015) or in vivo in yeast (Mans et al. 2015). C. Cloning of the 2-step PCR generated gRNA cassette
via Gibson assembly (DiCarlo et al. 2013) or restriction cloning (Chin et al. 2016). D. Several single gRNA cassettes cloned via Gibson assembly (Weninger et al. 2016), restriction cloning (Ryan et al. 2014) or modular assembly (Lee et al. 2015). Alternatively, two-gRNA cassette fragments in opposite orientation can be amplified in one reaction and cloned (Mans et al. 2015; Generoso et al. 2016). E. Pool of several single gRNA cassettes transformed to yeast cells with a gapped vector for in vivo recombination (Horwitz et al. 2015). F. crRNA array is cloned via Golden gate assembly of short synthetic fragments with homologous overlaps (Bao et al. 2015).
Figure 3: Overview of transcriptional control via CRISPR/Cas9 in yeast. A. Steric block of transcriptional initiation/elongation by catalytically inactive (“dead”) dCas9 bound in the promoter region. B. Transcriptional activation/repression using dCas9 fused to transcriptional activator/repressor domains. C. Multiple transcriptional regulation action using effector proteins recruited by RNA scaffolds. Pol III – RNA Polymerase III, NLS – nucleolar localization sequence,
L – self-cleaved leader sequence (e.g. SNR52), cr – crRNA, tracr – tracrRNA, TF – transcription factor, scRNA – scaffold RNA, Linker – scaffold RNA-binding linker protein domain.

Figure 4: Application of CRISPR/Cas9 systems for engineering of yeast cell factories. A. Production of (R,R)-2,3-butanediol from xylose. Multi-copy one-step integration of the xylose utilization and (R,R)-2,3-butanediol pathways into Ty-element delta sites in the genome (The figure is reprinted with permission from Elsevier: Shi et al. 2016). B. Production of lactic acid from glucose in an industrial yeast strain, one-step disruption of two genes in diploid strain and simultaneous integration of lactate dehydrogenase genes from L. plantarum (ldhL) (Stovicek, Borodina and Forster 2015). C. Production of deoxyviolacein, violacein, prodeoxyviolacein, and proviolacein from glucose. Transcriptional regulation (activation/repression) of different genes in violacein pathway leads to production of different violacein derivatives (The figure is reprinted with permission from Elsevier: Zalatan et al. 2015); VP64-activator domain, PP7/MS2 – RNA hairpin structures, PCP/MCP – RNA binding proteins. D. Production of naringenin from glucose. Cas9-
mediated one-step integration of the naringenin pathway into an intergenic locus. Downregulation of TSC13 mediated by catalytically inactive (“dead”) dCas9 (CRISPRi) to avoid the formation of by-products (The figure adapted from Vanegas, Lehka and Mortensen 2017).

Table 1. List of selected web-based bioinformatics tools for gRNA design in yeast.

<table>
<thead>
<tr>
<th>Name</th>
<th>Link</th>
<th>Reference</th>
<th>Input</th>
<th>Main features</th>
<th>Yeast species</th>
</tr>
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<tr>
<td>CRISPy-web</td>
<td><a href="http://crispy.secondarymetabolites.org">http://crispy.secondarymetabolites.org</a></td>
<td>Blin et al. 2016</td>
<td>gene name/ID, genomic coordinates</td>
<td>off-target</td>
<td>any user-submitted genome</td>
</tr>
<tr>
<td>CRISPR-ERA</td>
<td><a href="http://crispr-era.stanford.edu/">http://crispr-era.stanford.edu/</a></td>
<td>Liu et al. 2015</td>
<td>gene name, genomic coordinates, sequence</td>
<td>off-target, efficiency score, gene repression/activation</td>
<td>S. cerevisiae reference</td>
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<tr>
<td>CHOPCHOP v2</td>
<td><a href="http://chopchop.cbu.uib.no">http://chopchop.cbu.uib.no</a></td>
<td>Labun et al. 2016</td>
<td>gene name, genomic coordinates, sequence</td>
<td>off-target user defined, on-target algorithm, restriction sites</td>
<td>S. cerevisiae reference, C. albicans, C. tropicalis, C. glabrata, P. pastoris</td>
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<td>CRISPRdirect</td>
<td><a href="https://crispr.dbcls.jp/">https://crispr.dbcls.jp/</a></td>
<td>Naito et al. 2015</td>
<td>gene name, genomic coordinates, sequence</td>
<td>off-target, GC content, poly T, restriction sites</td>
<td>S. cerevisiae, S. pombe, K. lactis, Y. lipolytica, C. albicans, C. glabrata</td>
</tr>
<tr>
<td>Yeastriction</td>
<td><a href="http://yeastriction.tnw.tudelft.nl">http://yeastriction.tnw.tudelft.nl</a></td>
<td>Mans et al. 2015</td>
<td>gene name</td>
<td>off-target, AT content, self-complementarity, restriction sites</td>
<td>S. cerevisiae, several strains</td>
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<td>Benchling</td>
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<td></td>
<td>gene name, coordinates, sequence</td>
<td>off-target, on-target algorithm</td>
<td>S. cerevisiae reference, S. pombe, C. albicans, Y. lipolytica</td>
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<td>ATUM gRNA Design Tool</td>
<td><a href="https://www.atum.bio/eCommerce/cas9/input">https://www.atum.bio/eCommerce/cas9/input</a></td>
<td></td>
<td>gene name, coordinates, sequence</td>
<td>off-target</td>
<td>S. cerevisiae reference</td>
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</table>

<table>
<thead>
<tr>
<th>Reference</th>
<th>Availability</th>
<th>Organism &amp; Strain (ploidy)</th>
<th>Cas9 expression (vector, selection marker, promoter)</th>
<th>gRNA expression (vector, selection marker, promoter, terminator)</th>
<th>Application &amp; Efficiency</th>
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</thead>
<tbody>
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<td>DiCarlo <em>et al.</em> 2013</td>
<td>addgene</td>
<td><em>S. cerevisiae</em> BY4733 (n)</td>
<td>CEN/ARS, TRP1, <em>P</em> <em>TEF1</em>/<em>GALL</em>-Cas9</td>
<td>2µ, <em>URA3</em>, <em>P</em> <em>SNR52</em>/<em>T</em> <em>SUP4</em></td>
<td>single gene disruption/marker cassette insertion: 99%</td>
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<td>Gao and Zhao 2014</td>
<td>addgene</td>
<td><em>S. cerevisiae</em> LPY16936 (n)</td>
<td>2µ, LEU2, <em>P</em> _ADH1<em>_Cas9</em></td>
<td>2µ, <em>URA3</em>, <em>P</em> _ADH1*-HH ribozyme/HDV ribozyme-<em>T</em> <em>ADH1</em></td>
<td>single gene disruption: 100%</td>
</tr>
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<td>Ryan <em>et al.</em> 2014</td>
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<td><em>S. cerevisiae</em> S288C (n, 2n)</td>
<td>2µ, <em>kanMX</em>, <em>P</em> _RNR2*-Cas9*</td>
<td>tRNA&lt;sup&gt;P&lt;sub&gt;P&lt;/sub&gt;&lt;/sup&gt; -HDV ribozyme/<em>T</em> <em>SNR52</em></td>
<td>single/multiple gene disruption(s): 90-100%/19-85%, three-part marker cassette insertion: 70-85%</td>
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<td>Bao <em>et al.</em> 2015</td>
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<td><em>S. cerevisiae</em> BY4741 (n)</td>
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<td><em>P</em> _SNR52*-crRNA/<em>T</em> _SUP4*, <em>P</em> _RPR1*_tracrRNA/<em>T</em> <em>RPR1</em></td>
<td>single/multiple gene disruption: 27-100%</td>
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<td>Zhang <em>et al.</em> 2014</td>
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<td>2µ, <em>hphMX</em>, <em>P</em> <em>SNR52</em>/<em>T</em> <em>SUP4</em></td>
<td>single gene disruption: 15-60%</td>
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<td>CEN/ARS, TRP1, <em>P</em> <em>TEF1</em>/<em>Cas9</em></td>
<td>2µ, LEU2, <em>P</em> <em>SNR52</em>/<em>T</em> <em>SUP4</em></td>
<td>single/multiple gene disruption(s): 100%/ 50-100%</td>
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<td>euroscarf</td>
<td><em>S. cerevisiae</em> CEN.PK2-1c (n)</td>
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<td>2µ, <em>URA3</em>, <em>amdSYM</em>hphMX/<em>kanMX</em>/<em>LEU2</em>nat MX/<em>HIS3</em>/<em>TRP1</em>, <em>P</em> <em>SNR52</em>/<em>T</em> <em>SUP4</em></td>
<td>single gene deletion: 25-75%, multiple gene deletions/multiple gene cassette insertions: 65-100%</td>
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<td>Stovicek, Borodina and Forster 2015</td>
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<td><em>S. cerevisiae</em> CEN.PK11-7D (n)</td>
<td>CEN/ARS, <em>kanMX</em>, <em>P</em> <em>TEF1</em>/<em>ADH1</em>-Cas9</td>
<td>2µ, <em>natMX</em>, <em>P</em> <em>SNR52</em>/<em>T</em> <em>SUP4</em></td>
<td>single gene disruption &amp; gene cassette insertion: 65-97%</td>
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<td>Horwitz <em>et al.</em> 2015</td>
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<td><em>S. cerevisiae</em> CEN.PK2-1c (n)</td>
<td>integr. <em>can1A</em>: <em>P</em> <em>FBA1</em><em>Cas9</em>&lt;sup&gt;-hphMX&lt;/sup&gt;</td>
<td>2µ, <em>URA3/HIS3</em>/<em>natMX</em>, <em>P</em> <em>SNR52</em>/<em>T</em> <em>SUP4</em></td>
<td>single allele swap: 82-100%, multiple gene disruptions: 65-91%, multiple gene cassette integrations: 4.2%</td>
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<td>Vectors</td>
<td>Transgene</td>
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<td>S. cerevisiae D452-2 (n)</td>
<td>CEN/ARS, natMX, P_{TFE1}-Cas9</td>
<td>2µ, hphMX, P_{SNR52}/T_{SUP4}</td>
<td>two gene cassettes integration into a single gene locus: 25-100%</td>
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<td>addgene</td>
<td>S. cerevisiae BY4741 (n)</td>
<td>2µ, LEU2/URA3, P_{TRP1}-Cas9</td>
<td>P_{SNR52}/T_{SUP4}</td>
<td>single gene disruption: 97-98%</td>
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<td>Lee et al. 2015</td>
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<td>S. cerevisiae S288C (n)</td>
<td>CEN/ARS, URA3, P_{PGK1}-Cas9</td>
<td>tRNA^{Phe}-HDV ribozyme/T_{SNR52}</td>
<td>single/multiple gene disruption(s): 96%/ 21-76%</td>
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<td>Jakočiūnas et al. 2015b</td>
<td>on request</td>
<td>S. cerevisiae CEN.PK111-27B (n)</td>
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<td>multiple part gene cassette integrations into multiple gene loci: 30-97%</td>
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<td>Ronda et al. 2015</td>
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<td>CEN/ARS, TRP1, P_{TFE1}-Cas9</td>
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<td>gene cassette integration into multiple intergenic loci: 84-100%</td>
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<td>on request</td>
<td>S. cerevisiae HZ848 (n), CEN.PK2-1c (n)</td>
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<td>P_{SNR52}-crRNA/ T_{SUP4}, P_{RPR1} tracrRNA/T_{RPR1}</td>
<td>long gene fragment integration into multiple genomic loci: 75-88%</td>
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<td>Generoso et al. 2016</td>
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<td>S. cerevisiae CEN.PK113-7D (n) Ethanol Red (2n)</td>
<td>2µ, kanMX/natMX, P_{BOX5}-Cas9</td>
<td>P_{SNR52}/T_{SUP4}</td>
<td>single &amp; double gene disruptions: 91-98%</td>
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<td>Jessop-Fabre et al. 2016</td>
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<td>S. cerevisiae CEN.PK113-7D (n) Ethanol Red (2n)</td>
<td>CEN/ARS, kanMX, P_{TFE1}-Cas9</td>
<td>2µ, natMX, P_{SNR52}/T_{SUP4}</td>
<td>integration of a long gene fragment into a single locus: 95-100%/multiple loci: 60-70%</td>
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<td>Reider Ape et al. 2016</td>
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<td>S. cerevisiae BY4742 (2n)</td>
<td>2µ, URA3, LEU2, P_{ADH1}-Cas9</td>
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<td>three-part gene cassette integration into multiple intergenic loci: 40-95%</td>
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<td>S. cerevisiae BY4709 (n) RM11-1 (2n)</td>
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<td>P_{SNR52}-crRNA/ T_{SUP4}, P_{RPR1} tracrRNA/T_{RPR1}</td>
<td>Single gene non-sense mutation: 70-95%</td>
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<td>Liu et al. 2016</td>
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<td>S. cerevisiae var. boulardii ATCC MYA-796 (n)</td>
<td>CEN/ARS, natMX, P_{TFE1}-Cas9</td>
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<td>single/double gene disruption(s):100%/N/A</td>
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<td>S. cerevisiae BY4741 (n) YPH501 (2n)</td>
<td>CEN/ARS, LEU2, P_{GAL1}/nCas9 (840A)/nCas9 (D10A/A/n(d)/Cas9 (D10A/A)/n(840A))-P_{mCDA1}</td>
<td>2µ, URA3, P_{SNR52}/T_{SUP4}</td>
<td>Cytidine deaminase mediated single/double gene disruption(s): 16-54%/14-31%</td>
</tr>
<tr>
<td>Vanegas, Lehka and Mortensen 2017</td>
<td>on request</td>
<td>S. cerevisiae S288C (n) PJ69-4 (n)</td>
<td>Integ. intergenic X-3::P_{TFE1}-Cas9-URA3</td>
<td>CEN/ARS, LEU2 P_{SNR52}/T_{SUP4}</td>
<td>Integration of three-part multiple gene fragment into an intergenic site: 100%</td>
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</tbody>
</table>

**S. cerevisiae gene activation/repression**

<table>
<thead>
<tr>
<th>Authors</th>
<th>Delivery</th>
<th>Strain</th>
<th>Vectors</th>
<th>Transgene</th>
<th>Integration</th>
<th>Disruption</th>
<th>Comments</th>
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</thead>
<tbody>
<tr>
<td>Gilbert et al. 2013</td>
<td>addgene</td>
<td>S. cerevisiae W303</td>
<td>CEN/ARS, LEU2, P_{TRP1}/dCas9(-Mxi1)</td>
<td>CEN/ARS, URA3, P_{SNR52}/T_{SUP4}</td>
<td>several ten-fold reporter gene transcription repression (CRISPRi)</td>
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<tr>
<td>Farzadfar, Perli and Lu 2013</td>
<td>addgene</td>
<td>S. cerevisiae W303 (n)</td>
<td>integr. Atrp:: P_{TRP1}/dCas9(-VP64-TRP1)</td>
<td>2µ, HIS3/LEU2, P_{RPR1}/T_{RPR1}</td>
<td>transcription activation (activator domain)/repression (CRISPRi)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Authors</td>
<td>Type</td>
<td>Organism</td>
<td>Genotype/Description</td>
<td>Integration Method</td>
<td>Gene Silencing Method</td>
<td>Notes</td>
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<td>Zalatan et al. 2015</td>
<td>addgene</td>
<td><em>S. cerevisiae</em> W303 (n)</td>
<td>integr. <em>Aleu2</em>, his3::P_TDH1::Cas9*-CgLEU2/HIS3</td>
<td>CEN/ARS,</td>
<td>multiple gene transcription activation (RNA binding chimeric activators/repression (CRISPRi)</td>
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<td>Chavez et al. 2015</td>
<td>addgene</td>
<td><em>S. cerevisiae</em> W303 (n)</td>
<td>CEN/ARS, TRP1, P_TDH1::Cas9-VPR</td>
<td>2μ, URA3,</td>
<td>transcription activation (multiple activation domains)</td>
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<td>Smith et al. 2016</td>
<td>addgene</td>
<td><em>S. cerevisiae</em> BY4741 (n)</td>
<td>CEN/ARS, TRP1/URA3, P_TEF1::Cas9-Mxi1</td>
<td>P_KRPR (ToR)/TPRPI</td>
<td>transcription repression (repression domain-CRISPRi)</td>
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<tr>
<td>Vanegas, Lehka and Mortensen 2017</td>
<td>on request</td>
<td><em>S. cerevisiae</em> S288C (n)</td>
<td>Integr. Intergenic X-3::P_TEF1::Cas9*-dCas9*-VP64-URA3</td>
<td>CEN/ARS, LEU2</td>
<td>transcription activation (activator domain/repression (CRISPRi)</td>
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<td>Deaner and Alper 2017</td>
<td>on request</td>
<td><em>S. cerevisiae</em> BY4741 (n)</td>
<td>CEN/ARS, LEU2, P_TDH1::Cas9-Mxi1/ P_TDH1::Cas9-VPR</td>
<td>P_SNR52/T_SUP4</td>
<td>graded gene activation/repression (fold transcription activation-gene silencing)</td>
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<td>Horwitz et al. 2015</td>
<td>on request</td>
<td><em>K. lactis</em> ATCC8585 (n)</td>
<td>integr. K legislators::P_FRAI::Cas9*-hphMX</td>
<td>pKD1, natMX,</td>
<td>multiple gene cassette insertion into multiple gene loci: 2.1%</td>
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<td>Schwartz et al. 2016a</td>
<td>addgene</td>
<td><em>Y. lipolytica</em> ATCC MYA-2613 (n)</td>
<td>CEN/ARS, LEU2, P_AS188-TEF-Cas9*</td>
<td>SCR-riRNA_Gi/polYT</td>
<td>single gene disruptions (NHEJ/HR): 90-100%/64-88% (100% in KU mutant)</td>
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<td>Gao et al. 2016</td>
<td>addgene</td>
<td><em>Y. lipolytica</em> ATCC 201249 ATCC MYA-2613 (n)</td>
<td>CEN/ARS, LEU2, P_TEF1::Cas9</td>
<td>P_TEF1::HH</td>
<td>single gene disruption (NHEJ/HR): 62-98%/72% (94% in KU mutant), multiple gene disruptions (NHEJ): 19-37%</td>
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<tr>
<td>Schwartz et al. 2016b</td>
<td>addgene</td>
<td><em>Y. lipolytica</em> ATCC MYA-2613 (n)</td>
<td>CEN/ARS, LEU2, P_AS188-TEF-Cas9*</td>
<td>SCR-riRNA_Gi/polYT</td>
<td>gene cassette integration into an intergenic locus: 48-69%</td>
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<td>Weninger et al. 2016</td>
<td>on request</td>
<td><em>P. pastoris</em> CBS7435 (n)</td>
<td>PARS1*, ZEO, P_HX1::Cas9</td>
<td>P_HX1::HH</td>
<td>single gene disruption: 87-94%, double gene disruptions: 69%</td>
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<td>Jacobs et al. 2014</td>
<td>addgene</td>
<td><em>Sch. pombe</em> (P. pastoris)</td>
<td>tars, ura4, P_adh1::Cas9</td>
<td>P_adh1/HH</td>
<td>single gene disruption (allele swap): 85-90%</td>
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<td>Fernandez and Berro 2016</td>
<td>on request</td>
<td><em>Sch. pombe</em> FY527 (n)</td>
<td>tars, ura4fex1, P_adh1::Cas9</td>
<td>P_adh1/HH</td>
<td>single gene deletion (ORF removal): 33%</td>
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<td>Vyas, Barrasa and Fink 2015</td>
<td>on request</td>
<td><em>C. albicans</em> SC5314 (2n)</td>
<td>tars, ENO1 locus, natMX, P_ENO1::Cas9**</td>
<td>P_SNR52/T_ENO1</td>
<td>single/multiple gene disruption(s): 60-80%/20%</td>
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<td>Min et al. 2016</td>
<td>on request</td>
<td><em>C. albicans</em> SC5314 (2n)</td>
<td>linear cassette, P_ENO1::Cas9**</td>
<td>linear cassette,</td>
<td>single gene deletion (ORF replacement with marker</td>
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<td>P_SNR52/T_ENO1</td>
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<td></td>
<td>SN152 (2n)</td>
<td>cassette: 45-67%</td>
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<td><strong>C. glabrata</strong></td>
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<td>Enkler <em>et al.</em> 2016</td>
<td>on request</td>
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<td><em>C. glabrata</em> CBS138 (n)</td>
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<td>CEN, <em>TRP1</em>, P&lt;sub&gt;CSI&lt;/sub&gt;*&lt;sub&gt;CIS&lt;/sub&gt;-Cas9</td>
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<td>single gene disruption</td>
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<td><strong>Cr. neoformans</strong></td>
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<td>&quot;linear vector, URA5, P&lt;sub&gt;ACT1&lt;/sub&gt;-Cas9&quot;</td>
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<td>URA5, P&lt;sub&gt;Cas9&lt;/sub&gt;/polyT</td>
<td>single gene disruption (NHEJ/HR): 40–90%/20-90%</td>
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<tr>
<td>Arras <em>et al.</em> 2016</td>
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<td>integr. &quot;Safe Heaven&quot;-P&lt;sub&gt;TEF1&lt;/sub&gt;-Cas9</td>
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<td>linear vector, P&lt;sub&gt;ACT1&lt;/sub&gt;-HH ribozyme/HDV ribozyme-T&lt;sub&gt;TRP1&lt;/sub&gt;</td>
<td>single gene deletion (ORF replacement with marker cassette): 65-70%</td>
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</tbody>
</table>

* both components on a single expression element
* native *S. pyogenes Cas9*
** species codon optimized *Cas9*