Minireview

Yeast two- and three-species hybrids and high-sugar fermentation

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Summary
The dominating strains of most sugar-based natural and industrial fermentations either belong to *Saccharomyces cerevisiae* and *Saccharomyces uvarum* or are their chimeric derivatives. Osmotolerance is an essential trait of these strains for industrial applications in which typically high concentrations of sugars are used. As the ability of the cells to cope with the hyperosmotic stress is under polygenic control, significant improvement can be expected from concerted modification of the activity of multiple genes or from creating new genomes harbouring positive alleles of strains of two or more species. In this review, the application of the methods of intergeneric and interspecies hybridization to fitness improvement of strains used under high-sugar fermentation conditions is discussed. By protoplast fusion and heterospecific mating, hybrids can be obtained that outperform the parental strains in certain technological parameters including osmotolerance. Spontaneous postzygotic genome evolution during mitotic propagation (GARMi) and meiosis after the breakdown of the sterility barrier by loss of *MAT* heterozygosity (GARMe) can be exploited for further improvement. Both processes result in derivatives of chimeric genomes, some of which can be superior both to the parental strains and to the hybrid. Three-species hybridization represents further perspectives.

Introduction
Yeasts play an essential role in bread-making, the fermentation of alcoholic beverages and the production of bioethanol. Although a high number of fermentative yeast species have biotechnological relevance, the strains dominating most natural and industrial fermentations either belong to *S. cerevisiae* and *S. uvarum* or are their chimeric derivatives including the genetically highly diverse “hybrid species” *S. pastorianus/carlsbergensis* and *S. bayanus*. During the fermentation of high-sugar and high-gravity substrates, the *Saccharomyces* cells are exposed to hyperosmotic stress. To counterbalance the extracellular solute concentrations, the cells accumulate glycerol. Since the response to the hyperosmotic shock is a polygenic trait, significant improvement can be expected from concerted modification of the activity of multiple genes rather than from the manipulation of individual genes (e.g. Albertyn *et al.*, 1994; Shi *et al.*, 2018). Hybridization brings all alleles of all relevant genes of different strains together and recombine them during segregation/chimerization of the hybrid genomes. The hybrids and/or their chimeric derivatives can outperform the parental strains in certain technologically relevant properties including stress response. In this review, I describe basic knowledge about the hyperosmotic stress response and the exploitation of interspecies hybridization and hybrid evolution in improving the fitness of the strains under high-sugar conditions.

Yeasts face hyperosmotic stress during fermentation
The yeast cells have to cope with high-sugar and/or VHG (very high gravity) conditions in the fermentation of certain types of beverages, VHG brewing and VHG fuel ethanol production technologies, fermented vegetable extracts and high-sugar dough. When preparing dessert wines from late-harvest or botrytized grapes, the sugar concentrations in the must can be as high as 60% (e.g. Donèche, 1993; Sipiczki *et al.*, 2010). In ice-wine grape juice, the sugar content can reach 50% (Erasmus *et al.*, 2004). The total sugar content of sugar cane and sugar beet molasses fermented in traditional alcohol distilleries varies between 40% and 50% (e.g. Doelle and Docile,
1990; Khoja et al., 2018). Sweet dough (high-sugar dough) contains up to approximately 30% sucrose (Sasano et al., 2012), and high-sugar vegetable extracts can have 40–60% sugar (Ok and Hashinaga, 1997). In VHG brewing, the gravity of the wort can exceed 20°P (Huuskonen et al., 2010). In VHG fuel ethanol fermentation, the mashes contain sugar at concentrations higher than 25% to achieve ethanol yields higher than 15%. Certain S. cerevisiae strains can grow and vigorously ferment potato mash containing as much as 40% (2.2 M) glucose (Watanabe et al., 2015). In the shrinking cytoplasm, the microtubular and actin-based cytoskeletal structures become disorganized and deep plasma membrane invaginations are formed which the cells can fill up with amorphous cell wall material (Chowdhury et al., 1992; Slaninova et al., 2000). The overall cell volume is also reduced, and the yeast cells have to adapt their internal osmolarity to the hyperosmotic external conditions to restore the optimal cell volume (e.g. Pratt et al., 2003; Munna et al., 2015; Talemí et al., 2016).

The effect of hyperosmotic stress on the Saccharomyces cells

Upon being exposed to hyperosmotic conditions, yeast cells cease growing and propagating, rapidly lose intracellular water, thereby resulting in a loss of turgor pressure followed by shrinkage of the cytoplasm (Slaninova et al., 2000; Schaber and Klipp, 2008; Munna et al., 2015). In the shrinking cytoplasm, the microtubular and actin-based cytoskeletal structures become disorganized and deep plasma membrane invaginations are formed which the cells can fill up with amorphous cell wall material (Chowdhury et al., 1992; Slaninova et al., 2000). The overall cell volume is also reduced, and the yeast cells have to adapt their internal osmolarity to the hyperosmotic external conditions to restore the optimal cell volume (e.g. Pratt et al., 2003; Munna et al., 2015; Talemí et al., 2016).

The complexity of the response of the Saccharomyces cells to hyperosmotic stress

To cope with an increased external osmolarity caused by the high sugar concentration in the environment, the Saccharomyces cells initiate a complex adaptive program that includes the synthesis and retention of the compatible osmolyte glycerol, temporary arrest of cell-cycle progression, altered transcription and translation patterns. The genes and processes involved in the response have been reviewed in several recent studies (Saito and Posas, 2012; Hohmann, 2015; Saxena and Sitaraman, 2016; Taymaz-Nikerel et al., 2016; Auesukaree, 2017) and can be summarized as follows (Fig. 1).

The response is basically governed by the high-osmolarity glycerol (HOG) signalling pathway, whose core is the Hog1 MAP kinase (MAPK) cascade. The phosphorylated Hog1 enters the nucleus and activates the expression of several transcription factors, each of which is responsible for controlling the expression of a subset of osmoreponsive genes. Due to the activity of these genes, the intracellular level of glycerol increases which prevents the efflux of water from the cell into the environment. STL1 codes for a H+ symporter that transports glycerol inside the cell. GPD1 encodes a NADH-dependent glycerol-3-phosphate dehydrogenase that reduces dihydroxy-acetone-phosphate to glycerol-3-phosphate. Glycerol-3-phosphate is further converted to glycerol by the glycerol-3-phosphate phosphatase encoded by GPP2. In response to osmostress, the glycerol efflux mediated by the Fps1 aquaglyceroporin closes to keep the glycerol inside the cell, but this effect seems to be independent of Hog1. Hog1 was found to down-regulate the expression of the aquaporin Aqp2 and thereby restrict water loss. Apart from the genes directly involved in the HOG cascade, large numbers of additional genes have been implicated in the response by transcriptomic analyses and testing of deletion mutants (e.g. Erasmus et al., 2003; O’Rourke and Herskowitz, 2004; Ando et al., 2006; Tanaka-Tsuno et al., 2007; Jimenez-Martí et al., 2011). Recent studies reported results demonstrating that mitochondrial activity is also required for proper osmotic stress adaptation (Pastor et al., 2009; Gonzalez et al., 2016). The high number of up- and down-regulated genes indicates that a large proportion of the gene pool of the yeast cell has to be fine-tuned to give an adequate response to the high external osmolarity.

Correlation between the presence of foreign genes in the genome in ‘natural’ strains and their stress response

In numerous wine and industrial strains isolated from natural fermenting yeast communities, correlation was observed between the presence of genes of two or even more Saccharomyces species and the more adequate response of their cells to certain types of stress (for a review, see Marsit and Dequin, 2015). For example, S. cerevisiae strains harbouring genes from the more cryotolerant species S. uvarum or S. kudriavzevii are better adapted to growth and fermentation at low temperatures than the strains of ‘pure’ S. cerevisiae genomes (e.g. Peris et al., 2018). The proportion of the non-cerevisiae gene pool varies in these chimeric strains from a few genes to an (almost) complete subgenome (e.g. Emri et al., 2012; Peris et al., 2018). The transgressive stress-response phenotypes of certain natural chimeric strains indicate that stress tolerance might be improved synthetically by admixing genes of species. Examples of modification of osmotolerance by interspecies hybridization and posthybridization (postzygotic) genome evolution are shown in Table 1.
Fig. 1. A schematic diagram of the roles of the Hog1 protein kinase phosphorylated by the HOG MAPK (Mitogen-Activated Protein Kinase) pathway in the response of the *Saccharomyces* cell to the osmotic stress. The diagram is based on the review of Hohmann (2015).

Table 1. Examples of modulation of osmotolerance by hybridization and hybrid segregation.

| Species combination (protoplast fusion) | New phenotype | Reference |
|----------------------------------------|---------------|-----------|
| *S. cerevisiae* × *Z. mellis*          | Efficient fermentation at 30% glucose combined with good yields of ethanol production | Legmann and Margalith (1983, 1986) |
| *S. cerevisiae* × *D. hansenii*        | Combination of the higher thermostolerance and growth rate of the *S. cerevisiae* parent with higher osmotolerance from the *D. hansenii* parent | Loray et al. (1995) |
| *S. cerevisiae* × *T. delbrueckii*     | Tolerance to increased glucose concentrations (up to 70%) combined with increased arabinitol and glycerol production | Lucca et al. (1999, 2002) |
| *S. diastaticus* (var. *diastaticus*) × *Z. rouxii* | Fermentation and sugar utilization patterns of *S. diastaticus* combined with increased osmotolerance | Spencer et al. (1985) |
| *S. diastaticus* (var. *diastaticus*) × *S. uvarum* | Fermentation rate superior to those of the parental strain at 30% sugar combined with the ability to ferment at 40 °C | Stewart et al. (1988) |

| Sexual hybridization (mating, conjugation) | | |
|----------------------------------------|---------------|-----------|
| *S. cerevisiae* × *S. carlsbergensis*  | Improved fermentation rates in high-gravity wort (18° Plato) relative to the ale parent | Garcia Sanchez et al. (2012) |
| *S. cerevisiae* × *S. eubayanus*       | Increased fermentation rate in very high-gravity wort (25° Plato) relative to the parent strains | Krogerus et al. (2016) |
| *S. cerevisiae* × *S. kudriavzevii*    | Increased efficiency of fructose and glucose consumption in high-sugar wine fermentations | Lopandic et al. (2016); Gangl et al. (2017) |
| *S. cerevisiae* × *S. mikatae*         | Combination of low-temperature tolerance and osmotolerance (from *S. cerevisiae*) | Bellon et al. (2013) |
| *S. cerevisiae* × *S. uvarum*          | Increased fermentation efficiency, ethanol and glycerol production in most of 32% sugar | Restuccia et al. (2011) |
| *S. cerevisiae* × *S. uvarum*          | Slightly increased osmotolerance in certain hybrids | Pfleigler et al. (2014) |
| *S. cerevisiae* × *S. uvarum*          | Combination of fermentation phenotypes from both parents: robust fermentation in high-sugar juice (25% and 35.5%) and the production of wines with low volatile acidity | Bellon et al. (2015) |

| Hybrid segregation by GARMi | | |
|----------------------------------------|---------------|-----------|
| *S. cerevisiae* × *S. uvarum*          | A segregant becoming dominant after five rounds of fermentation of high-sugar grape must (evolved hybrid) fermented the grape juice of 35% sugar at much faster rate than the original hybrid strain | Bellon et al. (2018) |

| Hybrid segregation by GARMMe | | |
|----------------------------------------|---------------|-----------|
| *S. cerevisiae* × *S. kudriavzevii*    | Increased efficiency of fructose and glucose consumption in high sugar wine fermentations in certain F1 spore clones | Lopandic et al. (2016) |
| *S. cerevisiae* × *S. uvarum*          | Slightly increased osmotolerance in certain F1 spore clones | Pfleigler et al. (2014) |
| *S. diastaticus* (var. *diastaticus*) × *Z. rouxii* | Certain spore clones had greater capacity to raise sweet dough than the hybrid | Spencer et al. (1985) |
Improving high-sugar osmotolerance by interspecific and intergeneric hybridization

Somatic hybridization (protoplast fusion)

Osmotolerant somatic hybrids (growing at high sugar or salt concentrations) were obtained by protoplast fusion between *S. cerevisiae* and *Zygosaccharomyces mellis* (Legmann and Margalith, 1983, 1986), *S. diastaticus* (*S. cerevisiae* var. *diastaticus*) and *Z. rouxii* (Spencer et al., 1985), *S. diastaticus* (*S. cerevisiae* var. *diastaticus*) and *S. uvarum* (Stewart et al., 1988), *S. cerevisiae* and *Debaromyces hansenii* (Loray et al., 1995) and *S. cerevisiae* and *Torulaspora delbrueckii* (Lucca et al., 1999). The *S. diastaticus* × *Z. rouxii* hybrid was tolerant of sugar concentrations that were higher than those permitting the growth of the *S. diastaticus* parent. The viable spores of the hybrid formed clones of vegetative cells that had a much greater dough-raising capacity than either the original hybrid or a commercial baker’s yeast (Spencer et al., 1985). Unfortunately, little has been revealed from the genome structures of these hybrids. The few experimental data available indicate that the intergeneric hybrids had *Saccharomyces* genomes supplemented with only small percentages of the genomes of the non-*Saccharomyces* fusion partners (Spencer et al., 1985; Salek, 2002) (Fig. 2A).

**Hybridization by mating (conjugation)**

In the genus, *Saccharomyces* interspecies hybrids can be generated also by natural sexual hybridization because the reproductive isolation of the species is ‘postzygotic’. Cells of different *Saccharomyces* species can form viable hybrids by sexual mating (conjugation) but the hybrids are sterile. The sterility is mainly due to the failure of the (allosyndetic, homeologous) chromosomes to pair in meiosis (no functional gametes are produced) and to MAT heterozygosity (mating-specific genes are repressed) (for a review, see Sipiczki, 2018).

Hybrids of *S. cerevisiae* and *S. uvarum* strains were constructed that displayed combinations of positive phenotypic traits of the parents: robust fermentation in high-sugar grape juice (from *S. cerevisiae*) and the production of wines with low volatile acidity (from *S. uvarum*) (Restuccia et al., 2011; Bellon et al., 2015). A different combination of strains of these species and the mating of *S. cerevisiae* × *S. kudriavzevii* resulted in hybrids showing improved fermentation rate in a must of high sugar content (Lopandic et al., 2016; Gangl et al., 2017). The *S. cerevisiae* × *S. paradoxus* and *S. cerevisiae* × *S. mikatae* hybrids inherited the osmotolerance

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Fig. 2. Interspecies hybridization and the evolution of the hybrid genome.

A. Somatic hybridization (protoplast fusion)

B. Hybridization by sexual conjugation (mating). NS: non-*Saccharomyces* species; S: *Saccharomyces*; S₁: *Saccharomyces* species 1; S₂: *Saccharomyces* species 2. Star marks incomplete subgenome. GARMi: Genome Autoreduction in mitosis. GARMe: Genome Autoreduction in meiosis.
of the *S. cerevisiae* parent (Bellon et al., 2011, 2013). When *S. cerevisiae* was hybridized with *S. eubayanus*, the allotriploids and the allotetraploids, but not the allodiploids, outperformed the parental strains in fermentation of very high-gravity wort (Krogerus et al., 2016).

**Postzygotic genome segregation and chimerization broadens phenotypic diversity**

**Postzygotic shaping of the hybrid genome (postzygotic genome evolution)**

The hybrid genomes are usually unstable and prone to ‘postzygotic’ changes either during vegetative (mitotic) propagation of the sterile allopolyploid cells or during sporulation of the allopolyploid cells upon the breakdown of their sterility. As both processes are associated with spontaneous loss of chromosomes, the terms GARMi (Genome AutoReduction in Mitosis) and GARMMe (Genome AutoReduction in Meiosis) were recently proposed to designate them (Sipiczki, 2018) (Fig. 2B).

In GARMi (usually referred to as ‘hybrid stabilization’ or ‘hybrid evolution’), recurrent unequal mitotic divisions (biased segregation) generate alloaneuploid cells lacking chromosomes or arms of chromosomes in one or the other subgenome (for a review, see Sipiczki, 2018). GARMMe can take place after the breakdown of the sterility barrier in allotetraploid meiosis by the loss of *MAT* heterozygosity (misseggregation of the *MAT*-carrying chromosomes in one of the subgenomes) (Pfiegl er et al., 2012; Karanyicz et al., 2017). The alloaneuploid spores establish vegetatively propagating clones of mating-competent cells (F1 spore clones, ‘propagating gametes’) which can conjugate with each other (selfing) to form sporulation-proficient alloaneuploid zygotes producing alloaneuploid cells (F2 generation). As aneuploidy usually destabilizes the genome, the F2 cells can easily lose additional chromosomes when sporulate. The outcomes of both processes are strains of chimeric (mosaic) genomes composed of various combinations and proportions of the parental gene pools.

**Postzygotic genomic changes are associated with novel phenotypic traits**

Several studies have shown that the prolonged cultivation of a hybrid under a specific stress condition (high ethanol or sugar concentration, nutrient-limited conditions, extreme temperature, etc.) gradually improves its fitness (e.g. Piotrowski et al., 2012; Dunn et al., 2013; Lopandic et al., 2016; Smukowski Heil et al., 2017; Krogerus et al., 2018). Where investigated, the improved fitness turned out to be associated with the accumulation of segregants (‘evolved hybrids’ or ‘evolved variants’) of specific types of chimeric genomes in the population.

The genetic changes are most probably due to spontaneous misseggregation and/or structural changes of certain chromosomes at mitotic divisions, and to randomly occurring mutations. The stress factor does not induce these genetic changes; it only differentially affects the growth of the derivatives if they differ in fitness. The less sensitive derivatives will gradually overgrow the rest of the population. Different conditions lead to different genomic outcomes from the same hybrid (e.g. Piotrowski et al., 2012; Lopandic et al., 2016).

The gradual enrichment of the hybrid culture with a specific derived segregant is nicely illustrated by the results of Bellon et al. (2018) who let a *S. cerevisiae × S. uvarum* hybrid propagate vegetatively through a series of successive fermentations of a botrytized Riesling must of high sugar content (350 g l⁻¹ initial sugar) and monitored the changes in the population. After the 5th round of fermentation, 95% of the cells lacked the entire chromosome 14 of *S. uvarum* and the rest lacked its left arm. It was hypothesized that the loss of the *S. uvarum* chromosome 14 accounted for the increased fitness of the population. However, chromosome 14 instability had been observed also in *S. cerevisiae × S. uvarum* hybrids exposed to nitrogen limiting conditions (Dunn et al., 2013). Thus, the loss of this chromosome may not be specific for the high-sugar stress response.

In a different study (Lopandic et al., 2016), no karyotype changes (no GARMi) were detected in *S. cerevisiae × S. uvarum* and *S. cerevisiae × S. kudriavzevii* hybrids after the fermentation of a high-sugar grape must (Grüner Veltliner with an initial sugar concentration of 337.5 g l⁻¹) but both hybrids segregated at meiosis (GARMMe). As the spores were viable (produced colonies), the hybrids must have been allotetraploid. The tetrad isolated from the *S. cerevisiae × S. uvarum* hybrid showed the segregation pattern characteristic of the tetrads in which the (second) sterility barrier breaks down (Pfiegl er et al., 2012): two spores lacked the *MAT*-carrying chromosome of the *S. uvarum* subgenome. The clones established by these alloaneuploid spores remained stable during fermentation, whereas the other two, presumably allodiploid spore clones underwent GARMi, resulting in mixed populations of cells lacking various chromosomes mostly in the *S. uvarum* subgenome. In contrast to the finding of the Bellon laboratory (Bellon et al., 2018), no chromosome 14 instability was detected here. The spore clones that retained the *MAT*-carrying chromosomes of both parents showed the phenotype of the hybrid: had higher fermentation rates and richer aroma profile in the high-glucose must, produced more ethanol and less volatile acids than the other spores and the parental strains (Lopandic et al., 2016; Gangl et al., 2017). The karyotype of the
S. cerevisiae × S. kudriavzevii hybrid also showed 2:2 segregation (GARMe), but different chromosomes were involved, and all spore clones performed worse than the hybrid for all parameters examined. The segregation of AFLP (amplified fragment length polymorphism) pattern in these alloaneuploids was much more complex, indicating that smaller-scale GARMe, and GARMi changes took place both within the subgenomes and between them. The observed diversity of spore clones indicates that meiotic segregation poses a risk to the application of hybrids of genomes larger than allodiploid in fermentation technologies.

New perspectives in three-species hybridizations

The gene pools of the two-species Saccharomyces hybrids or their alloaneuploid or chimeric derivatives can be enriched with genes from a third species by mating with a strain of the third species (e.g. Sipiczki et al., 2014; Antunovics et al., 2018a,b). Two strategies have been developed for the construction of three-species synthetic hybrids and/or genomic chimeras (Antunovics et al., 2018b) (Fig. 3). One method is based on the fertility of the alloaneuploid spore clones produced in GARMe after the breakdown of the sterility barrier in tetraploid meiosis. These clones sporulate, and their spores can be mass-mated with spores of a homothallic strain or with cells of a heterothallic strain of the third species. The outcome of the process is a chimeric strain with a complete genome of the third species and incomplete genomes of the parental strains of the two-species alloaneuploid. As spores of diverse alloaneuploid/chimeric genomes are produced during GARMe of the two-species hybrid, the arising zygotes will have diverse genomes composed of various proportions of the genes of the three parental species. The other approach generates true allotriploid hybrids having complete genomes of all three species. Their construction starts with hybridizing two species and selecting a stable (sterile) allodiploid hybrid. This sterile MATa/MATalpha hybrid is then mated with fertile spores or cells of the third species by taking advantage of the phenomenon referred to as ‘rare mating’ (Gunge and Nakatomi, 1972). MAT heterozygosity suppresses the mating-specific genes and makes the allo diploid cells mating-incompetent, but very rarely certain cells escape the block and conjugate with mating-competent cells or spores of a different strain. S. cerevisiae × S. kudriavzevii × S. uvarum (‘cekudvarum’) chimeric and allotriploid hybrid strains have been recently produced with these methods (Antunovics et al., 2018a,b). Both types of three-species strains showed transgressive phenotypic traits. Although their response to high-sugar stress has not been examined, improved osmotolerance can be expected in certain strains or in their GARMI/GARME derivatives because natural isolates of composite genomes consisting of genes from these species had phenotypes suitable to overcome stuck fermentation (Christ et al., 2015).

Conflict of interest

None declared.

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Interspecies hybridization and high-sugar fermentation 1107
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