

Yeast: the soul of beer's aroma—a review of flavour-active esters and higher alcohols produced by the brewing yeast

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Abstract Among the most important factors influencing beer quality is the presence of well-adjusted amounts of higher alcohols and esters. Thus, a heavy body of literature focuses on these substances and on the parameters influencing their production by the brewing yeast. Additionally, the complex metabolic pathways involved in their synthesis require special attention. More than a century of data, mainly in genetic and proteomic fields, has built up enough information to describe in detail each step in the pathway for the synthesis of higher alcohols and their esters, but there is still place for more. Higher alcohols are formed either by anabolism or catabolism (Ehrlich pathway) of amino acids. Esters are formed by enzymatic condensation of organic acids and alcohols. The current paper reviews the up-to-date knowledge in the pathways involving the synthesis of higher alcohols and esters by brewing yeasts. Fermentation parameters affecting yeast response during biosynthesis of these aromatic substances are also fully reviewed.

Keywords Higher alcohols · Fusel alcohols · Esters · Beer aroma · Brewing yeast · *Saccharomyces* spp

Introduction

Beer is one of the most pleasant beverages in the world, the final taste/aroma of which is the resultant sum of several

hundreds of flavour-active compounds produced in the course of every step of brewing. However, the great majority of these substances are produced during the fermentation phase and consist of metabolic intermediates or by-products of the main living character of brewing—the yeast. Higher alcohols, esters and vicinal diketones (VDKs) are the key elements produced by yeast, which will ultimately determine the final quality of the beer. While higher alcohols and esters are desirable volatile constituents of a pleasant beer, VDKs are often considered as off-flavours. Together with these, yeast metabolism contributes with other three groups of chemical compounds: organic acids, sulphur compounds and aldehydes.

All flavour-active components in beer must be kept within certain limits; otherwise, a single compound or group of compounds may predominate and destroy the flavour balance. Furthermore, potent odourants like esters may act in synergy with other components affecting beer flavour in concentrations well below their threshold values (Meilgaard 1975a). However, each type of beer has its own prevailing aroma triggered either by the yeast strain (Nykanen and Nykanen 1977; Peddie 1990; Ramos-Jeunehomme et al. 1991; Rossouw et al. 2008) chosen or by parameters used during fermentation (Lodolo et al. 2008; Saerens et al. 2008a; Bravi et al. 2009; Verbelen et al. 2009; Blasco et al. 2011; Berner and Arneborg 2012; Dekoninck et al. 2012; Hiralal et al. 2013). For example, while only isoamyl acetate (banana-like aroma) concentrations are above the threshold level in most lager beers, ales normally have ethyl acetate (solvent-like aroma) and ethyl hexanoate (apple-like aroma) as additional flavouring compounds with levels above the threshold (Meilgaard 1975b; Alvarez et al. 1994). However, for the vast majority of beers, other compounds like diacetyl (a vicinal diketone) should be found below the threshold values as it contributes negatively with a buttery flavour to the beer. Table 1 shows threshold values of the main esters and higher alcohols present in lager beer. As a full review of diacetyl

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Table 1 Threshold values of most important esters and higher alcohols present in lager beer (Engan 1974; Meilgaard 1975b; Engan 1981)

Compound	Threshold (mg L ⁻¹)	Concentration range (mg L ⁻¹)	Aroma impression
Acetate esters			
Ethyl acetate	25–30	8–32	Fruity, solvent
Isoamyl acetate	1.2–2	0.3–3.8	Banana
Phenylethyl acetate	0.2–3.8	0.1–0.73	Roses, honey
MCFA ethyl esters			
Ethyl hexanoate	0.2–0.23	0.05–0.21	Apple, fruity
Ethyl octanoate	0.9–1.0	0.04–0.53	Apple, aniseed
Higher alcohols			
<i>n</i> -Propanol	600	4–17	Alcohol, sweet
Isobutanol	100	4–57	Solvent
Isoamyl alcohol	50–65	25–123	Alcoholic, banana
Amyl alcohol	50–70	7–34	Alcoholic, solvent
2-Phenylethanol	40	5–102	Roses

formation was recently published (Krogerus and Gibson 2013), this paper will limit the discussion on the desirable odourants produced by the brewing yeast in the course of fermentation—higher alcohols and esters.

Higher alcohols

Also known as fusel alcohols, higher alcohols are the most abundant organoleptic compounds present in beer. The brewing yeast absorbs amino acids present in wort from which they take the amino group so it can be incorporated in its own structures. What is left from the amino acid (α -keto acid) enters in an irreversible chain reaction that will ultimately form a by-product—higher alcohols. This pathway was suggested long ago by Ehrlich (1907), who was intrigued with the structural molecular similarities between the active amyl alcohol with isoleucine and isoamyl alcohol with leucine. This

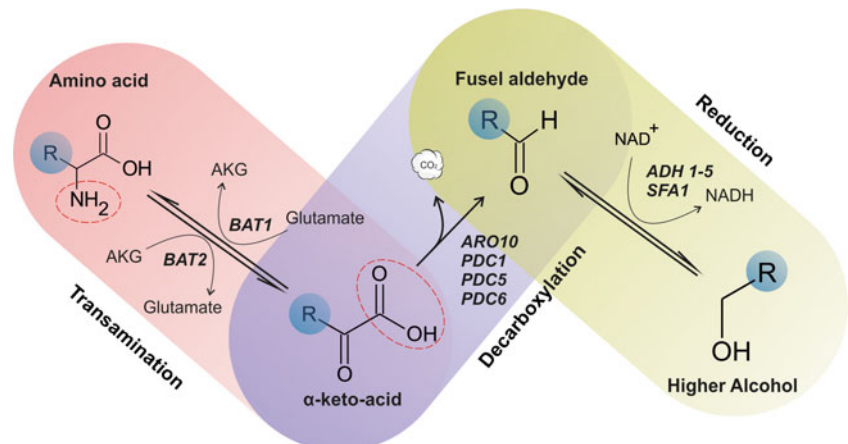
observation has led Ehrlich to investigate whether these amino acids were involved in higher alcohol synthesis or not. When supplementing the fermenting medium with those amino acids, he evidenced an increased production of fusel alcohols. This observation led him to state that amino acids were enzymatically hydrolyzed to form the corresponding fusel alcohols, along with ammonia and carbon dioxide. As ammonia was not detected in the medium, it was assumed to be incorporated into yeast proteins. Few years later, Neubauer and Fromherz (1911) proposed a few intermediate steps to the Ehrlich pathway, completing the metabolic scheme as it is known until today. However, a detailed enzymatic chain reaction was only demonstrated several decades later (Sentheshanmuganathan and Elsdén 1958; Sentheshanmuganathan 1960), establishing the elementary enzymatic sequence for the Ehrlich pathway: transaminase, decarboxylase and alcohol dehydrogenase (Fig. 1). Although this pathway is the most studied and discussed, higher alcohols are also formed during upstream (anabolic pathway) biosynthesis of amino acids (Chen 1978; Dickinson and Norte 1993; Oshita et al. 1995). The most important is the de novo synthesis of branched-chain amino acids (BCAAs) through the isoleucine–leucine–valine (ILV) pathway (Dickinson and Norte 1993).

Biosynthesis of higher alcohols

Transamination

The first step in the Ehrlich pathway involves four enzymes encoded by the genes *Bat1* (*Twt1* or *Eca39*), *Bat2* (*Twt2* or *Eca40*), *Aro8* and *Aro9*. These enzymes are transaminases that catalyse the transfer of amines between amino acids and their respective α -keto acid, using glutamate/ α -ketoglutarate as a donor/acceptor. While *Bat1*- and *Bat2*-encoded enzymes are involved in the BCAA transamination (Eden et al. 1996; Kispal et al. 1996), *Aro8p* and *Aro9p* were first described as being aromatic amino acid aminotransferases I and II,

Fig. 1 The Ehrlich pathway and the main genes involved in the synthesis of enzymes catalyzing each reaction. The reversible transamination reaction uses different BAT-encoded enzymes—while *BAT2* catalyses the transfer of the amino group from the amino acid to α -ketoglutarate (AKG), *BAT1* is usually required on the reverse transamination for amino acid biosynthesis



respectively (Iraqi et al. 1999). Further studies carried out by Urrestarazu et al. (1998) demonstrated that Aro8 and Aro9 encoded enzymes had broader substrate specificity than just for aromatic amino acids. This was confirmed in the work performed by Boer et al. (2007), who cultivated *Saccharomyces cerevisiae* using six independent nitrogen sources followed by transcriptome analysis. All phenylalanine, methionine and leucine activated the transcription of Aro9 and Bat2 genes.

A recent study mapped almost entirely (97 %) the proteome of *S. cerevisiae* (Picotti et al. 2013). The authors organized the proteome into a network of functionally related proteins, which they called as “modules”. Within these modules, they highlighted one comprising of Bat1, Bat2, Rpn11, Hsp60 and Ilv2, which they termed B1B2 module. The core of this module is composed by Bat1p and Bat2p—two paralogous enzymes involved in the metabolism of the BCAAs. While Bat1p is mainly involved in the anabolism of BCAAs (amination of α -keto acids), Bat2p is almost exclusively involved in the catabolism of BCAAs (deamination of BCAAs). Thus, Bat1- and Bat2-encoded proteins catalyse the same metabolic reaction in opposite directions. Strictly related to these two proteins is the Ilv2-encrypted enzyme, which catalyses an early step in the synthesis of BCAAs from pyruvate (Picotti et al. 2013).

The subcellular location of enzymes catalyzing the synthesis of fusel alcohols has been studied in the past (Kispal et al. 1996; Schoondermark-Stolk et al. 2005) and recently emphasized (Avalos et al. 2013). Isobutanol is produced by yeast originally in the cytoplasm via the Ehrlich pathway or by anabolic synthesis inside the mitochondria (Kohlhaw 2003). Avalos et al. (2013) redirected the entire enzymatic biosynthetic pathway of that fusel alcohol to the mitochondrial matrix. Compartmentalization of the Ehrlich pathway within the mitochondria increased isobutanol production by 260 %, whereas overexpression of the same pathway in the cytoplasm only improved yields by 10 %. These results are justified by the more favorable environmental conditions found in the mitochondria matrix, which enhanced enzymatic activity.

Decarboxylation

After transamination, the remaining α -keto acids can be decarboxylated to form the respective aldehyde and this is a point of no return in the Ehrlich pathway (Dickinson et al. 1997). There are five genes encoding decarboxylases in *S. cerevisiae*: three encoding pyruvate decarboxylases (PDC1, PDC5 and PDC6), ARO10 and THI3 (Dickinson et al. 1997; Romagnoli et al. 2012; Bolat et al. 2013), with all enzymes encrypted depending on the cofactor thiamine diphosphate (TPP). From within these genes, only PDC5 and ARO10 were described to encode decarboxylases with a broad substrate specificity (Vuralhan et al. 2003; Vuralhan

et al. 2005; Romagnoli et al. 2012). Dickinson et al. (1998) have shown that valine is decarboxylated by any of the enzymes encrypted by PDC1, PDC5 or PDC6. In the case of isoleucine, all five decarboxylases of the family are able to produce active amyl alcohol (Dickinson et al. 2000). THI3-encrypted enzyme cannot catalyse the decarboxylation of the aromatic amino acids phenylalanine and tyrosine, while all other four can (Dickinson et al. 2003). The single expression of THI3 in a quadruple deleted (*pdc1 Δ pdc5 Δ pdc6 Δ aro10 Δ*) gene *S. cerevisiae* strain had no α -keto acid decarboxylase activity (Vuralhan et al. 2003, 2005). Further studies involving THI3 suggest that its role in the Ehrlich pathway is rather regulatory than catalytic (Mojzita and Hohmann 2006).

Although the lager-brewing yeast *Saccharomyces pastorianus* is long known to be a natural aneuploid hybrid of *S. cerevisiae* with another *Saccharomyces* sp. (Vaughan and Kurtzman 1985), only recently the missing link was proven to be *Saccharomyces eubayanus* (Libkind et al. 2011). This fact has called the attention of Bolat et al. (2013) upon the contribution of ARO10 gene expression from each of the subgenomes on the production of higher alcohols. The authors amplified by PCR both *S. eubayanus*-like and *S. cerevisiae*-like alleles of ARO10 (*LgSeubARO10* and *LgScARO10*, respectively) from the genomic DNA of *S. pastorianus*. The alleles showed a sequence identity of 80 % at the DNA level and 84 % at the protein level. The results have also shown that *S. cerevisiae* alleles of ARO10 are present in a ratio of 3:1 from those present in *S. eubayanus* subgenome. These authors have equally demonstrated that both *S. eubayanus*-like and *S. cerevisiae*-like ARO10-encoded isoenzymes had similar activity for most of the substrates tested with preferred decarboxylation action against phenylpyruvate. However, the activity of *LgSeubARO10*-encrypted enzyme towards ketoisovalerate (precursor of isobutanol) was twofold higher than that encoded by *LgScARO10*. Moreover, those authors also suggest that *S. eubayanus*-like and *S. cerevisiae*-like ARO10-derived α -oxo acid decarboxylases exert different roles during beer fermentation by *S. pastorianus*. Fusel alcohols produced by the Ehrlich pathway would involve preferentially the *S. cerevisiae*-ARO10 decarboxylase. Conversely, higher alcohols formed by de novo synthesis would rely almost exclusively on the *LgSeubARO10*-encrypted isoenzyme.

Reduction to higher alcohols

After decarboxylation, the fusel aldehydes enter in the last step of the Ehrlich pathway, where they are converted into their respective alcohols by action of alcohol dehydrogenases. Any one of the *S. cerevisiae* alcohol dehydrogenases encoded by Adh1, Adh2, Adh3, Adh4 and Adh5 or the formaldehyde dehydrogenase encrypted by Sfa1 is able to catalyse the conversion of fusel aldehydes into higher alcohols

transamination of amino acids from the growth medium by using a knockout strain (*eca39Δ* and *eca40Δ*). In addition to these deletions, *ilv2Δ* was also investigated, and thus, the activity of acetolactate synthase (ILV2) could be assessed. Without ILV2, the synthesis of isoleucine is hindered, causing an increase of the main precursor (after pyruvate)— α -ketobutyrate. As this α -keto acid is the precursor of propanol, the authors evidenced a significant increase in this fusel alcohol produced by *eca39Δ eca40Δ ilv2Δ* strain (Eden et al. 2001). This strain was also unable to produce isobutanol as α -acetolactate could not be synthesized from pyruvate due to lack of ILV2. Thus, as no external amino acid could be used in the Ehrlich pathway due to *eca39Δ eca40Δ*, the role of ILV2 gene was confirmed in the anabolic pathway of isobutanol. On the other hand, the synthesis of active amyl alcohol and isoamyl alcohol was reduced, but still unexpectedly present (Eden et al. 2001). ILV2 was recently addressed to be integrated to a protein network module of functional similar proteins involved in BCAAs and physically connected to the mitochondria (Picotti et al. 2013).

Esters

Compared to other yeast metabolites, esters are only trace elements. Nevertheless, despite being “a drop in the ocean” of beer’s constituents, esters are the most important aroma elements produced by yeast. That is because esters have a very low odour threshold in beer (Meilgaard 1975b; Saison et al. 2009) and, yet to a large extent, may define its final aroma (Engan 1974; Suomalainen 1981; Nykanen and Suomalainen 1983; Peddie 1990; Meilgaard 1991; Saerens et al. 2008a; Saison et al. 2009; Verbelen et al. 2009; Hiralal et al. 2013). However, if overproduced, they can negatively affect the beer with a bitter, over fruity taste. Thus, it is crucial for the brewer to keep the optimum conditions to obtain a balanced beer in terms of its ester profile.

Esters are mainly formed during the vigorous phase of primary fermentation by enzymatic chemical condensation of organic acids and alcohols. Volatile esters in beer can be divided in two major groups: the acetate esters and the medium-chain fatty acid (MCFA) ethyl esters. The former group comprises esters synthesized from acetic acid (acetate) with ethanol or a higher alcohol. In the ethyl esters’ family, ethanol will form the alcohol radical and the acid side is a MCFA. Although dozens of different esters can be found in any beer (Engan 1974; Meilgaard 1975b), six of them are of major importance as aromatic constituents: ethyl acetate (solvent-like aroma), isoamyl acetate (banana aroma), isobutyl acetate (fruity aroma), phenyl ethyl acetate (roses and honey aroma), ethyl hexanoate (sweet apple aroma) and ethyl octanoate (sour apple aroma).

Esters are synthesized in the cytoplasm of the brewing yeast, but readily leave the cell as they are lipophilic. However, while small-chain acetate esters easily diffuse through the plasmatic membrane, MCFA ethyl esters may have their passage hindered (Nykanen and Nykanen 1977; Nykiinen et al. 1977; Dufour 1994).

To be synthesized into esters, organic acids must be linked to a coenzyme A to form an acyl-CoA molecule. Acyl-CoAs are highly energetic entities, which in the presence of oxygen can be β -oxidized (“cut”) into smaller units (acetyl-CoA) in the mitochondria. This will happen unless the organic acid involved is the acetic acid itself, which in this case will be turned into acetyl-CoA. However, the great majority of acetyl-CoA produced by the yeast cells comes from the oxidative decarboxylation of pyruvate. Aerobic conditions inside the mitochondria make acetyl-CoA to enter in the Krebs cycle to form ATP (respiration). In the absence of oxygen, acetyl-CoA will be enzymatically esterified with an alcohol to form the acetate esters. Moreover, MCFA ethyl esters are formed from longer chains of acyl-CoA with ethanol. Figure 2 drafts the main metabolic routes of the brewing yeast contributing to higher alcohols and ester synthesis.

Biosynthesis of acetate esters

Acetate esters are the major flavour components of beer as they are present in much higher concentrations than other volatile esters. The enzymatic involvement of synthases on ester production dates from the 1960s (Nordström 1962), but the enzyme was only purified and named as alcohol acetyltransferase (AATases) back in 1981 by Yoshioka and Hashimoto (1981). The most studied and best characterized enzymes responsible for ester synthesis are AATases I and II (EC 2.3.1.84), encoded by the genes ATF1 and ATF2 (Yoshioka and Hashimoto 1981; Malcorps and Dufour 1992; Fujii et al. 1994; Nagasawa et al. 1998; Yoshimoto et al. 1998; Verstrepen et al. 2003b; Molina et al. 2007; Dekoninck et al. 2012; Zhang et al. 2013). It was also found that bottom-fermenting lager yeasts have an extra ATF1 homologous gene (Lg-ATF1) (Fujii et al. 1994), which encodes an AATase very similar to that encoded from the original ATF1 gene (Fujii et al. 1996). This additional gene expression on lager yeast enhances acetate ester production and ultimately beer’s aroma profile.

The best way to understand the role of a gene’s expression is by either overexpressing or deleting it. A heavy body of literature focus on these genetic modifications to better understand the role of ATF1, ATF2 and Lg-ATF1 gene expression on the total acetate ester production (Fujii et al. 1994, 1996; Nagasawa et al. 1998; Yoshimoto et al. 1998; Verstrepen et al. 2003b; Zhang et al. 2013). Very recently, a brewer’s yeast

strain was designed to increase ester/higher alcohol ratio by overexpressing ATF1 and knocking down a gene related to higher alcohol synthesis (Zhang et al. 2013). Ester production by the genetically modified strains was considerably higher than that of parental cells. Verstrepen et al. (2003b) have earlier carried out a more detailed work concerning deletion and overexpression not only of the AFT1 but also of its homologous Lg-ATF1 as well as ATF2. As others in the past (Fujii et al. 1994, 1996; Nagasawa et al. 1998), those authors clearly demonstrated the strong impact exerted by the expression levels of ATF genes on acetate ester production. For example, they have shown that overexpressing ATF1 strains may have up to 180-fold increased isoamyl acetate production and a 30-fold increased ethyl acetate production when compared to wild-type cells. In fact, their analysis also revealed that ATF1-encrypted ATTases seem to be responsible for the great majority of acetate ester production. Through specific deletion of ATF1 and ATF2, no acetate esters originated from alcohols with more than five carbon atoms, like isoamyl acetate and phenyl ethyl acetate, were formed. This means that the most desirable banana aroma (isoamyl acetate) in beer depends exclusively on ATF1- and ATF2-encoded enzymes. Later, Saerens et al. (2008b) confirmed that the maximum expression levels of ATF1 and ATF2 are directly correlated to the final concentration of acetate esters. However, the knockdown (*atf1Δatf2Δ*) executed by Verstrepen et al. (2003b) could only reduce the production of smaller esters such as ethyl acetate by 50 %. Together with other pieces of evidence (Malcorps et al. 1991; Malcorps and Dufour 1992), this result makes clear that there might be more ATTases involved on acetate ester production, but this goes beyond the knowledge present in published data. Figure 3a schematizes the chemical reaction for the production of the main acetate esters and genes involved in these reactions.

The presence of acetate esters on alcohol-free beers (AFBs) is imperative. AFBs can be produced either from physical removal of ethanol from the finished beer or by controlling the biological process involved in beer fermentation (Branyik et al. 2012). AFBs produced by membrane processes have usually less body and a low aromatic profile, thermally dealcoholized AFBs may suffer heat damages, while beers obtained by biological methods have often a sweet and worty off-flavour (Montanari et al. 2009). The lack of ethanol itself greatly affects the retention of volatile aroma-active compounds (Perpete and Collin 2000). Very recently, Strejc et al. (2013) isolated a brewing yeast mutant capable of overproducing isoamyl acetate and isoamyl alcohol. The sweet banana odour from isoamyl acetate could then be a solution to overcome the undesirable worty off-flavour of AFB. Sensory analyses showed that the increased level of isoamyl acetate ester had a positive effect on the fruity (banana) palate fullness and aroma intensity of the AFB produced.

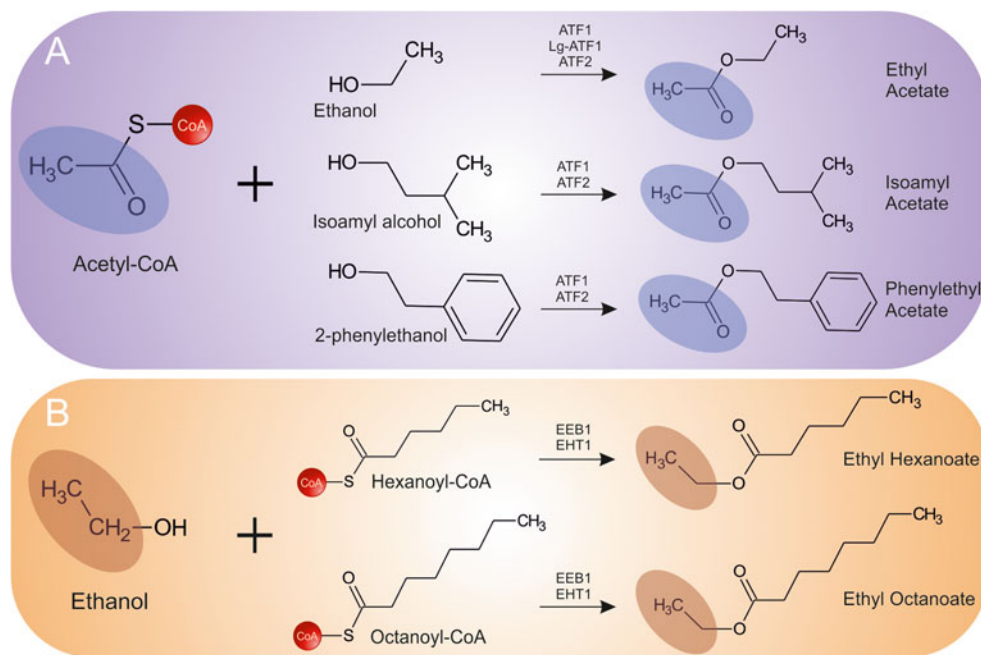
Biosynthesis of ethyl esters

When inserted in all history of volatile ester research, MCFA ethyl esters received less attention from the literature in the past. This happened because they are much less present in beer compared to their acetate counterparts. Nonetheless, works focused on ethyl esters in brewing fermentations have become much more common in the past decade, most of them were carried out by Saerens et al. (2006, 2008a). Before them, with evidence published long ago (Malcorps and Dufour 1992), Mason and Dufour (2000) suggested that apart from ATF1- and ATF2-encoded enzymes, there should be a different enzyme involved on ethyl ester synthesis. This ester-synthesizing enzyme, called ethanol hexanoyl transferase, is responsible for generating ethyl hexanoate from ethanol and hexanoyl-CoA (Mason and Dufour 2000). Later in 2006, Saerens and coworkers proved that MCFA ethyl esters are produced by the brewing yeast through a condensation reaction between an acyl-CoA unity and ethanol (Fig. 3b), catalysed by two acyl-CoA/ethanol *O*-acyltransferases (AEATases) encoded by *Eeb1* and *Eht1* genes (Saerens et al. 2006). Moreover, these authors were highly convinced of the role of each of these genes on the final MCFA ethyl ester content. A single deletion on *EeB1* reduced the formation of ethyl butanoate, ethyl hexanoate, ethyl octanoate and ethyl decanoate in 36, 88, 45 and 40 %, respectively. *Eht1* knockout strain on the other hand only had ethyl hexanoate and ethyl octanoate production affected. Additionally, a double-deletion (*eeb1Δ* and *eht1Δ*) strain produced a similar ethyl ester profile to the *eeb1Δ* single-deletion strain. This means that *EeB1* is the most relevant gene on MCFA ethyl ester synthesis (Saerens et al. 2006). However, even though double deletion caused a pronounced drop in ethyl esters measured, only ethyl hexanoate production was virtually extinguished. Thus, there must be other, yet unknown, AEATases involved on the MCFA ethyl ester synthesis. Also, overexpression of those genes did not increase MCFA ethyl ester production even when more precursors of these esters were added to the fermenting medium. This fact was explained as a consequence of extra esterase (breakdown) activity exerted by *Eeb1*- and *Eht1*-encoded proteins, which was also demonstrated in vitro by the same authors (Saerens et al. 2006).

Ester regulation

The net rate of ester production depends not only on the availability of the precursor's substrates (Saerens et al. 2006; Hiralal et al. 2013) but also, to a significant extent, on the enzymatic balance of synthesis (by AATases) (Yoshimoto et al. 1998; Mason and Dufour 2000; Verstrepen et al. 2003b; Saerens et al. 2006; Zhang et al. 2013) and breakdown (by esterases) of esters (Fukuda et al. 1996; Fukuda et al.

Fig. 3 A scheme of the chemical reactions involving the biosynthesis of acetate esters (a) and medium-chain fatty acid ethyl esters (b). The main genes involved in each reaction are presented above the reaction arrows



1998a; Lilly et al. 2006). Esterases are a group of hydrolyzing enzymes that catalyse the cleavage and/or avoid the formation of ester bonds. Fukuda et al. (1998b) have chosen another strategy to raise the end net of isoamyl acetate production by a sake strain of *S. cerevisiae*. Instead of enhancing AATases' activity, they avoided isoamyl acetate cleavage by deleting the acetate-hydrolyzing esterase gene (IAH1, previously known as EST2) responsible for encoding the carboxylesterase that hydrolyzes that ester (Fukuda et al. 1996). The mutant-deficient strain produced approximately 19 times higher amounts of isoamyl acetate when compared with the parent strain. Fukuda et al. (1998a) have further proven the important balance activity between AATases and esterases for the net rate of ester accumulation by *S. cerevisiae*. More evidence of the IAH1-encoded esterase on the breakdown of esters was built up recently by Lilly et al. (2006). In addition to isoamyl acetate, the authors also reported decreased production of ethyl acetate, phenyl ethyl acetate and hexyl acetate by the overexpressing IAH1 mutant strain. These findings are in agreement with the recent published data by Ma et al. (2011) whose work determined the crystalline structure of the enzyme encoded by IAH1 gene. They have shown that an additional C-terminus was involved on the substrate-binding region. Furthermore, it was also demonstrated that this C-terminus restricts access to the active site of the enzyme, playing a vital role in determining substrate specificity. Non-modified IAH1-encoded esterase had the highest hydrolytic activity against shorter acetate esters. Moreover, this activity was greatly reduced against ethyl hexanoate and almost null for ethyl decanoate, which suggests that IAH1-encoded enzyme is more specific for shorter-chain esters. This was confirmed by truncating the additional C-terminus present in

the enzyme. This modified variant was now able to hydrolyze longer ethyl ester chains such as decanoate. Those authors concluded that deletion of the C-terminus provides better access to the active site of the enzyme, which allows it to accommodate longer acyl chains (Ma et al. 2011).

Esters on aging beer

The ester profile of a given beer may change drastically during storage either by action of yeast (bottle refermentation) (Vanderhaegen et al. 2003) or by spontaneous chemical condensation of organic acids with ethanol (Vanderhaegen et al. 2006; Saison et al. 2009; Rodrigues et al. 2011). With time, hop-derived components are oxidized to form 3-methylbutyric and 2-methylbutyric acids, which are spontaneously esterified to their respective ethyl esters (3-methylbutyrate and 2-methylbutyrate) (Williams and Wagner 1979) that impart the aged beer a winy aroma (Williams and Wagner 1978). On the other hand, some esters as isoamyl acetate are known to be hydrolyzed during beer storage (Neven et al. 1997). Chemical hydrolysis and esterification are acid-catalysed (Vanderhaegen et al. 2006), but the remaining esterases from yeast autolysis can also play their role in unpasteurized beers (Neven et al. 1997). Other ethyl esters as ethyl nicotinate (medicinal, solvent, anis-like aromas), ethyl pyruvate (peas, freshly cut grass) and ethyl lactate (fruity, buttery) are also formed in aging beers (Saison et al. 2009). For all reasons mentioned in these lines, aging beers tend to lose fresh fruity aromas, giving place to sweeter odours.

Yeast response to fermentation parameters

Yeast strain

As regards the lines written above, it is easy to conclude that the production of most aroma-active compounds is strictly dependent on the yeast strain chosen for the fermentation. The genome associated to each strain is unique and will ultimately define the final aroma profile of the product (Ramos-Jeunehomme et al. 1991; Rossouw et al. 2008). This makes the selection of the right strain the most important task to make good beer. Yet, it is crucial that the brewer keeps his strain safe not only from “wild” yeast contaminations but also from genetic drifts that may occur in the course of serial repitching (Sato et al. 1994; Jenkins et al. 2003; Powell and Diacetis 2007). While the repeated repitching processes will not cause prominent loss of physiological characteristics of the brewing yeast (Powell and Diacetis 2007; Buhligen et al. 2013; Vieira et al. 2013), the accumulation of variants may eventually cause certain characteristics to linger on subsequent generations. That is why brewers must keep frozen stocks of their yeast strains for periodical restart of fresh pitching cultures.

Temperature

It has been reported that rising fermentation temperatures increase BAP2 expression in the brewing yeast *S. cerevisiae* (Yukiko et al. 2001). This gene is responsible for encoding a broad-substrate specificity permease that promotes the transport of the BCAAs valine, leucine and isoleucine into the yeast cell (Didion et al. 1996). The higher availability of amino acids within the cell favor the catalytic Ehrlich pathway, increasing higher alcohol production (Yukiko et al. 2001). Saerens et al. (2008b) obtained increasing levels of propanol, isobutanol, isoamyl alcohol and phenyl ethanol by rising the fermentation temperature using two different brewing yeast strains. Conversely, those authors have shown that although increasing temperatures promote the expression of all BAT1, BAT2 and BAP2, only BAT1 could be strongly correlated with the final concentration of higher alcohols, in particular of propanol (Saerens et al. 2008b).

As higher alcohol formation is temperature-dependent (Landaud et al. 2001), changes in temperature may cause changes in the availability of fusel alcohols, which are necessary for ester formation (Calderbank and Hammond 1994). Indeed, a slight change in temperature from 10 to 12 °C can increase ester production of up to 75 % (Engan and Aubert 1977). Saerens et al. (2008b) have shown that the AATase-encoding genes ATF1 and ATF2 are more expressed with increasing temperatures during beer fermentation. Furthermore, the maximum expression of these genes clearly

correlated with the end concentration of ethyl acetate, isoamyl acetate and phenyl ethyl acetate.

Hydrostatic pressure

With increasing market demands, breweries are continuously increasing reactor sizes for beer production. The incredibly high fermenters naturally generate a huge hydrostatic pressure that generally increases the concentration of carbon dioxide dissolved in beer. The excess in dissolved CO₂ inhibits yeast growth by unbalancing decarboxylation reactions (Rice et al. 1977; Knatchbull and Slaughter 1987; Renger et al. 1992; Shanta Kumara et al. 1995; Landaud et al. 2001). As said before, decarboxylation is a fundamental step in either higher alcohol or acetyl-CoA synthesis. As acetyl-CoA is the main precursor of acetate esters, hydrostatic pressure unbalances beer flavour most probably by limiting the substrate availability for ester formation (Landaud et al. 2001).

Wort composition

It is not hard to understand that wort's composition will greatly influence the final beer aroma. After all, fermenting wort is nothing more than a growth medium from which the brewing yeast absorbs nutrients for living and to where it lays its metabolic by-products. Thus, changes in the nutrient content will trigger different yeast responses.

Sugars

High-gravity brewing (HGB) or even very high-gravity (VHG) brewing became a common practice for commercial breweries as it can bring great economic benefits (Yu et al. 2012; Lei et al. 2013b). The use of HGB can not only increase the brewery capacity in up to 20–30 % without any investment in equipment but also improve the haze and smoothness of the beer (Stewart 2007). However, HGB often brings an unbalanced flavour profile to the finished beer, being the most common perturbation the overproduction of acetate esters, impairing the beer with over-fruity and solvent-like aromas (Anderson and Kirsop 1974; Peddie 1990; Saerens et al. 2008b). Anderson and Kirsop (1974) observed up to an eight-fold increase in acetate ester production when the specific gravity of wort was doubled. Saerens et al. (2008b) have tested ale and lager strains upon increasing specific wort gravity. Although all higher alcohols measured showed an increased accumulation, after dilution to reach the normal ethanol content (5.1 %, v/v), only the fermentations performed by the ale strain remained with unbalanced high levels of fusel alcohols. Additionally, all acetate esters were overproduced by both lager and ale strains when increasing the initial specific gravity of the fermenting wort. Yet, not only the amount but also the type of sugars may influence the changes in the

aromatic profile of the final beer. Easily assimilable glucose- and fructose-rich worts normally generate beers with higher contents of esters than those rich in maltose (Younis and Stewart 1998, 1999, 2000; Pidlocke et al. 2009). Both 21 and 24 °P worts enriched with maltose syrup fermentations performed by Pidlocke et al. (2009) produced less acetate esters compared to fermentations carried out with glucose syrup-enriched worts. The reason why an individual assimilable sugar has a different impact on ester production is still unknown. Younis and Stewart (1998) suggested that higher levels of glucose increase acetyl-CoA, which is the main substrate for acetate ester synthesis. In the same way, maltose-rich worts may weakly induce acetyl-CoA formation for acetate ester production (Shindo et al. 1992). Moreover, while glucose rapidly enhances ester synthase activity in carbon-starved cells by directly inducing ATF1 transcription through Ras/cAMP/PKA nutrient pathway, maltose relies on the slow “fermentable growth medium-induced” (FGM) pathway to do so (Verstrepen et al. 2003a). Furthermore, a nitrogen source is needed to maintain the transcription of ATF1 and Lg-ATF1 in the course of the fermentation (Verstrepen et al. 2003a). Increasing levels of maltose as a sole carbon source in a synthetic medium showed an increasing tendency to accumulate acetate esters (Saerens et al. 2008a). Conversely, Dekoninck et al. (2012) have shown that although sucrose had a higher impact on ATF1 expression when compared to maltose, a remarkable decrease in acetate esters was observed during HGB. The high amount of sucrose stimulated yeast’s metabolism and growth which ultimately increased the uptake of amino acids. This discussion leads to another important factor on HGB altering aroma profile of the beer—the carbon-to-nitrogen (C/N) ratio. The addition of sugary syrups is a common practice to increase the specific gravity of the wort in HGB. However, these syrups generally lack nitrogen, which normally reduces the total free amino nitrogen (FAN) content of the wort. Therefore, adjuncts usually increase the C/N ratio, which in turn may lead nitrogen to be a growth-limiting factor (Verstrepen et al. 2003a; Saerens et al. 2008a; Lei et al. 2012, 2013a). Any alteration in sugar or FAN levels affects acetate ester accumulation, but not ethyl esters (Saerens et al. 2008a). Additionally, diluted FAN content observed in HGB leads to abnormal yeast physiology and unbalanced beer flavour (Lei et al. 2012).

FANs

Although a wide range of nitrogen-containing compounds are dissolved in the wort, the brewing yeast can only assimilate the smaller molecules, generally called FANs. The discussion of FANs interfering with beer aroma will inevitably lead to the absorption of amino acids to form higher alcohols through the Ehrlich pathway. The type and amount of amino acids will also lead the yeast to different responses and ultimately to final

beer aromatic profile (Äyräpää 1971; Lei et al. 2013a). In fact, treating the wort with proteases increases the final FAN and ultimately increases the higher alcohols and esters by the brewing yeast in either HGB or normal gravity brewing (Lei et al. 2013c). The addition of BCAAs like valine, leucine and isoleucine to the fermenting wort increases the formation of their respective fusel alcohols—*isobutanol*, *isoamyl alcohol* and *amyl alcohol* (Engan 1970; Äyräpää 1971; Procopio et al. 2013). Recently, Procopio et al. (2013) have shown that not only the addition of valine, leucine and isoleucine increased the formation of fusel alcohols but intriguingly also did *proline*. Since *proline* cannot be converted into a higher alcohol via the Ehrlich pathway, its role on fusel alcohol formation induction was attributed to the synthesis of *glutamate* from this amino acid. A recent study showed that the supplementation of wort with *lysine* and *histidine* improved the performance of a lager brewing yeast in HGB (Lei et al. 2013a). Compared to *lysine*, *histidine* greatly affected the aromatic profile by increasing the formation of higher alcohols and ester. Moreover, recent reports confirmed that FAN content of wort can affect the transcription of both ATF1 and BAT1 genes (Saerens et al. 2008b; Lei et al. 2012).

Oxygen and unsaturated fatty acids

Dissolved oxygen and unsaturated fatty acids (UFAs) in wort are remarkably known as negative regulators of ester synthesis by brewing yeast (Anderson and Kirsop 1974; Anderson and Kirsop 1975a, b; Taylor et al. 1979; Thurston et al. 1982; Malcorps et al. 1991; Fujii et al. 1997; Fujiwara et al. 1998). Oxygen was once thought to indirectly reduce ester formation by decreasing acetyl-CoA availability (Anderson and Kirsop 1974). However, when genetic studies came into fashion, oxygen and UFAs were proven to directly inhibit the expression of ATF1 and ATF2 (Fujii et al. 1997). Fujiwara et al. (1998) have further complemented that oxygen and UFAs repress the expression of ATF1 by different regulatory pathways. While the repression of ATF1 by oxygen is mediated by the Rox1–Tup1–Ssn6 hypoxic repressor complex (Fujiwara et al. 1999), UFAs intermediate through the low-oxygen response element (Vasconcelles et al. 2001). In addition to acetate esters, it has been also shown that increasing levels of UFAs in the fermenting medium reduce the production of ethyl esters by the brewing yeast (Saerens et al. 2008a).

Considering what is written above, Moonjai et al. (2002) assessed the potential of rich UFA lipid supplements to decrease the need of wort aeration. The results have shown that the treated yeast with UFAs can be pitched into poor-oxygenated worts without losing fermentation potency or organoleptic quality. A reduced amount of oxygen supplied to the wort may increase flavour stability of the final beer and will limit potential oxidative stress upon the brewing yeast (Gibson et al. 2008). Inspired by this potential, Hull (2008)

assessed the replacement of wort oxygenation by treatment of the pitching yeast with olive oil rich in UFAs. The industrial-scale test succeeded without major effects on the acceptability of the beer produced. Therefore, UFA-treated yeast may be of particular help on HGB, once worts with specific high gravity have limited oxygen solubility (Baker and Morton 1977).

Conclusions

Formation of higher alcohols and esters by brewing yeast involves complex enzymatic and regulatory pathways. Nonetheless, much progress has been made in elucidating not only the genes involved in the transcription of key enzymes during the biosynthesis of these aroma-active substances but also on the importance of the subcellular location of these enzymes inside the yeast cell. Also, important steps have been taken towards substrate specificity as regards the analysis of the crystalline structure of IAH1-encoded esterase for small acetate esters.

Much progress has been also done in proteomics describing the B1B2 module of functionally related proteins. The core of this module is composed by Bat1p and Bat2p—two paralogous enzymes involved in the metabolism of the BCAAs catalyzing the same metabolic reaction in opposite directions. It was also interesting to know the contribution of the subgenomes of *S. pastorianus* to the final concentration of higher alcohols. Fusel alcohols originated from the catabolic pathway involve preferentially the *S. cerevisiae*-ARO10 decarboxylase. Conversely, higher alcohols formed by de novo synthesis rely almost exclusively on the *LgSeubARO10*-encoded isoenzyme.

While acetate ester production is largely dependent on ATF1- and ATF2-encoding enzymes, substrate concentration seems to be the major limiting factor for ethyl ester synthesis.

The bad aspects involving HGB affecting beer aroma are still a reality. However, each day, breweries and scientists head forward to overcome these problems. Not only genetically modified brewing yeasts are being developed but also interesting techniques like pretreatment of pitching yeast with oil-rich UFAs to substitute wort oxygenation are being applied. In addition to those well-known BCAAs valine, leucine and isoleucine increasing the formation of their respective fusel alcohols (isobutanol, isoamyl alcohol and amyl alcohol), proline interestingly can also raise the end net of fusel alcohols through the formation of glutamate—a key compound in yeast metabolism that may be used as an amine donor in amino acid synthesis.

Despite the clear progress describing fusel alcohols and ester synthesis by brewing yeast, there is still much to be found in this field.

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