The Ehrlich Pathway for Fusel Alcohol Production: a Century of Research on *Saccharomyces cerevisiae* Metabolism


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*Saccharomyces cerevisiae* has been used for at least eight millennia in the production of alcoholic beverages (41). Along with ethanol and carbon dioxide, fermenting cultures of this yeast produce many low-molecular-weight flavor compounds. These alcohols, aldehydes, organic acids, esters, organic sulfides, and carbonyl compounds have a strong impact on product quality. Indeed, the subtle aroma balance of these compounds in fermented foods and beverages is often used as an organoleptic fingerprint for specific products and brands (42). Food fermentation by yeast and lactic acid bacteria is accompanied by the formation of the aliphatic and aromatic alcohols known as fusel alcohols. Fusel oil, which derives its name from the German word *fusel* (bad liquor), is obtained during the distillation of spirits and is enriched with these higher alcohols. While fusel alcohols at high concentrations impart off-flavors, low concentrations of these compounds and their esters make an essential contribution to the flavors and aromas of fermented foods and beverages. Fusel alcohols are derived from amino acid catabolism via a pathway that was first proposed a century ago by Ehrlich (13). Amino acids represent the major source of the assimilable nitrogen in wort and grape must, and these amino acids are taken up by yeast in a sequential manner (valine, leucine, isoleucine, methionine, and phenylalanine) are taken up slowly throughout the fermentation time (32). After the initial transamination reaction (Fig. 1), the resulting α-keto acid cannot be redirected into central carbon metabolism. Before α-keto acids are excreted into the growth medium, yeast cells convert them into fusel alcohols or acids via the Ehrlich pathway.

Current scientific interest in the Ehrlich pathway is supported by increased demands for natural flavor compounds such as isoamyl alcohol and 2-phenylethanol, which can be produced from amino acids in yeast-based bioconversion processes (14), as well as by the need to control flavor profiles of fermented food products. The goal of this paper is to present a concise centenary overview of the biochemistry, molecular biology, and physiology of this important pathway in *S. cerevisiae*.

**EARLY HISTORY OF THE EHRLICH PATHWAY**

The turn of the 20th century saw the establishment of many biochemical principles. In 1877 von Ewald and Kühne proposed the notion of enzymes (62), and in 1893 Ostwald proved enzymes to be catalysts (46, 47). The following year (1894), Fischer (15) demonstrated the “lock-and-key” relationship between enzymes and substrates. Alcoholic fermentation in yeast cell extracts was described by Büchner in 1897 (3). In 1902 Fischer (16) and Hofmeister (24) demonstrated that proteins are polypeptides. It was in this background of accomplished chemistry and a growing knowledge of biochemistry that the German biochemist Felix Ehrlich (1877 to 1942) worked.

In 1904, after isolating and characterizing isoleucine, Ehrlich noted the structural similarities between this amino acid and active amyl alcohol and between leucine and isoamyl alcohol. These observations led him to investigate whether these fusel alcohols were derived from the amino acids (13). Indeed, the supplementation of yeast fermentation mixtures with either leucine or isoleucine led to increased production of fusel alcohols. Ehrlich then proposed (13) that the amino acids were split by a “hydrating” enzyme activity to form the corresponding fusel alcohols, along with carbon dioxide and ammonia. Ammonia, which was not detected in these experiments, was assumed to be incorporated into yeast protein. In 1911, Neubauer and Fromherz proposed a modified metabolic scheme (44) that constitutes the Ehrlich pathway as it is still viewed today (Fig. 1). As the first intermediate in the pathway, they proposed an α-keto acid, which can subsequently be decarboxylated into an aldehyde, which is then reduced into the fusel alcohol. The acceptor of the amino group from the starting amino acid was not known and neither were the exact natures of any of the individual enzyme-catalyzed steps. Subsequent work by Lampitt (36), Yamada (67–69), and Thorne (58, 59) confirmed that all fusel alcohols produced by yeast can be derived from amino acid catabolism. An overview of the Ehrlich pathway intermediates and products is presented in Table 1.

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IDENTIFICATION AND CONFIRMATION OF THE KEY ENZYMATIC STEPS IN THE EHRLICH PATHWAY

In the 1950s and 1960s, Sentheshanmuganathan studied the conversion of tyrosine into tyrosol. He demonstrated that cell extracts of *S. cerevisiae* transfer the amino groups from asparagine, isoleucine, leucine, methionine, norleucine, phenylalanine, tryptophan, and tyrosine to \(\Delta^2\)-ketoglutarate and, hence, that the first step of the Ehrlich pathway can be catalyzed by an aminotransferase activity (54, 55). He also demonstrated the decarboxylation of \(\Delta^2\)-hydroxy \(p\)-hydroxy 3-phenylpyruvate into \(\Delta^2\)-hydroxy \(p\)-hydroxy 3-phenylacetaldehyde and the NADH-dependent reduction of the latter compound into tyrosol. This finding established a key reaction sequence for the Ehrlich pathway: transaminase, decarboxylase, and alcohol dehydrogenase.

While, until the late 1990s, most experimental observations were compatible with the existence of the Ehrlich pathway, its in vivo activity was not proven. For example, experiments in which radioactively labeled amino acids were converted into radioactively labeled fusel alcohols did not rule out alternative intermediates and enzymes. Yeast molecular biology raised further questions on the identities of the enzymes involved in the Ehrlich pathway, as many candidate enzymes and genes for each of the reaction steps were identified. This raised the possibility that different isozymes may be involved under different environmental and/or nutritional conditions.

Final experimental verification of the role of the Ehrlich pathway in amino acid catabolism was obtained through \(^{13}\)C labeling studies. After the growth of *S. cerevisiae* under normal physiological conditions in minimal media with individual...
amino acids specifically labeled with $^{13}$C as the sole nitrogen source, $^{13}$C nuclear magnetic resonance spectroscopy was employed to identify the metabolic sequences involved. Since the fate of individual atoms in each intermediate from the amino acid to the end product could be monitored, the Ehrlich pathway was proved for the first time in the catabolism of leucine, valine, isoleucine, phenylalanine, tryptophan, and methionine into the corresponding fusel alcohols (6–8, 10, 50). Further confirmation of the key in vivo involvement of the Ehrlich pathway and the nonparticipation of other routes was obtained by testing specific mutants in which potential alternative metabolic pathways were blocked (6–8, 10).

During studies of the catabolism of [2-$^{13}$C]valine into isobutanol labeled at the expected C-1 position, the formation of isoamyl alcohol labeled at C-2 was observed (7). This result was shown to be due to the entry of $^{13}$C-labeled $\alpha$-ketoisovalerate, formed by valine deamination, into the leucine biosynthetic pathway, thus giving rise to labeled isoamyl alcohol (Fig. 1) (7). This finding indicates a mixing of catabolic and biosynthetic pathways for these branched-chain amino acids via the shared intermediate $\alpha$-ketoisovalerate.

The Ehrlich pathway for methionine catabolism represents a special case. A conventional Ehrlich pathway involves transamination into $\alpha$-keto-γ-(methylthio)butyrate ($\alpha$-KMBA), decarboxylation into β-(methylthio)propionaldehyde (often called methional), and then reduction into methanol. In addition, a demethiolase activity acts on both methionine and $\alpha$-KMBA to produce methanethiol and $\alpha$-ketobutyrate (50). Whether this reaction occurs spontaneously or enzymatically is still unknown.

INITIAL TRANSMINATION REACTION

Four S. cerevisiae proteins have been implicated in the initial transamination step of the Ehrlich pathway (Fig. 1). Twt1p (also known as Bat1p or Eca39p) is the mitochondrial branched-chain amino acid aminotransferase, and Twt2p (Bat2p or Eca40p) is the cytosolic isozyme. The mitochondrial isozyme is highly expressed in batch cultures during exponential growth and is repressed during stationary phase, while the cytosolic isozyme has the opposite expression pattern. The genes encoding both enzymes were cloned by two independent groups in 1996 (11, 34). Eden et al. (12) showed that an eca40 (ntw2) mutation drastically reduces the production of isobutanol. Decreased production of active amylo alcohol and isoamy alcohol was also seen by others (37, 53, 70), although the results also indicated the involvement of a Twt2p-independent transaminase activity in the formation of these alcohols.

Aro8p and Aro9p were initially characterized as the aromatic amino acid aminotransferases I and II, respectively (31). Urrestarazu et al. (60) demonstrated that Aro8p also exhibits in vitro activity with the amino donors methionine, $\alpha$-aminoadipate, and leucine and with phenylpyruvate as the amino acceptor and, reversibly, with their $\alpha$-keto acid analogues as amino acceptors and phenylalanine as the amino donor. Aro9p also has broader substrate specificity than originally described (60). Apparently, Aro8p and Aro9p act as broad-substrate-specificity amino acid transaminases in the Ehrlich pathway. This notion is supported by genome-wide expression profiling; when S. cerevisiae is grown in a glucose-limited chemostat with phenylalanine, methionine, or leucine as the sole nitrogen source, ARO9 and TWT2 are consistently upregulated compared to these genes in cultures grown with ammonia, proline, and asparagine, three nitrogen sources whose catabolism does not involve the Ehrlich pathway (2) (Fig. 2).

DECARBOXYLATION STEP

The irreversible decarboxylation step, which commits 2-oxo acids to the Ehrlich pathway, was initially attributed to pyruvate decarboxylase (5, 55, 65). However, the yeast genome was shown to harbor no fewer than five genes that show sequence similarity with thiamine diphosphate (TPP)-dependent decarboxylase genes. Three of these genes (PDC1, PDC5, and PDC6) encode pyruvate decarboxylases (25–28, 52, 57), while ARO10 and THI3 are alternative candidate genes for Ehrlich pathway decarboxylases (8, 10, 63). Null mutants with defects in the five structural genes encod-
ing (putative) TPP-dependent decarboxylases (pdc1, pdc5, pdc6, aro10, and thi3), both singly and in combination, have been studied to identify whether specific enzymes catalyze the decarboxylation of the individual α-keto acid intermediates. This research led to the initial conclusion that the major decarboxylase involved in leucine catabolism is encoded by THI3 (8). In valine catabolism, any one of the three isozymes Pdc1p, Pdc5p, and Pdc6p will decarboxylate α-ketoisovalerate (7). However, the conclusions from the leucine and valine experiments were drawn while the decarboxylase encoded by the ARO10 (YDR380w) gene was not yet known and consequently could not be quantitatively assessed. In isoleucine catabolism, any one of the family of decarboxylases encoded by PDC1, PDC5, PDC6, ARO10, and THI3 was found to be sufficient for the conversion of isoleucine to active amyl alcohol (6). The THI3 decarboxylase has no role in the catabolism of the aromatic amino acids phenylalanine and tyrosine: in a thi3 null mutant, the decarboxylation of 3-phenylpyruvate and 3-indolepyruvate is apparently accomplished by any one of the remaining four decarboxylases (10), while in methionine catabolism, the decarboxylation of α-KMBA is effected specifically by Aro10p (50, 63).

In chemostat cultures grown with either leucine, methionine, or phenylalanine as the sole nitrogen source, fusel acids and alcohols derived from amino acids other than the nitrogen source are produced in significant amounts (63). An α-keto acid decarboxylase activity for a broad range of substrates was measured in cell extracts of such cultures and suggested the involvement of a common, broad-specificity decarboxylase activity. ARO10 was the only decarboxylase gene whose transcript profile correlated strongly with α-keto acid decarboxylase activity in the chemostat cultures (2, 63, 64) (Fig. 2). A comprehensive characterization of Arop10p using a combination of genetic, physiological, and biochemical approaches (63) confirmed that Aro10p is a broad-substrate-specificity decarboxylase. These experiments also demonstrated the existence of an Aro10p-independent α-keto acid decarboxylase activity (64), which requires the functional alleles of both THI3 and at least one of the pyruvate decarboxylase genes PDC1, PDC5, and PDC6. Transcriptome analyses and decarboxylase activity measurements of an S. cerevisiae aro10Δ strain, a double aro10Δ thi3Δ deletion strain, and a quadruple pdc1Δ pdc5Δ pdc6Δ aro10Δ mutant strain grown in carbon-limited chemostats with phenylalanine as the nitrogen source indicated that (i) PDC5 is strongly upregulated in an aro10Δ background (Fig. 3) and also encodes a broad-substrate α-keto acid decarboxylase (J.-M. Daran, unpublished observation), (ii) PDC5 expression depends on the presence of THI3 (Fig. 3), and (iii) in contrast to the expression of ARO10 only (in a pdc1Δ pdc5Δ pdc6Δ thi3Δ strain), the expression of THI3 only (in a pdc1Δ pdc5Δ pdc6Δ aro10Δ strain) does not result in any α-keto acid decarboxylase activity (63, 64). THI3 has recently been demonstrated to be involved in the regulation of thiamine homeostasis in S. cerevisiae (43, 45), which further suggests that its role in the Ehrlich pathway may be regulatory rather than catalytic. While Pdc1, Pdc5, Pdc6, and Aro10 are cytosolic, Thi3 has been observed to be localized in both the cytosol and the nucleus (43, 45), which would support further its role in regulatory function. A systematic investigation of the catalytic properties of all five (putative) TPP-dependent decarboxylases (Aro10p, Thi3p, Pdc1p, Pdc5p, and Pdc6p) is essential for a final resolution of the substrate specificity of these key enzymes in the Ehrlich pathway.

In lactic acid bacteria, the catabolism of branched-chain, aromatic, and sulfur-containing amino acids also involves transamination followed by a decarboxylation reaction. The fate of the aldehyde seems to be species dependent, but in Lactococcus lactis, a gene called kcdA, has been identified and characterized. Similar to the yeast Aro10p gene, kcdA encodes a broad-substrate α-keto acid decarboxylase that exhibits the highest level of activity on branched-chain α-keto acids (56).

REDUCTION OR OXIDATION OF FUSEL ALDEHYDES

In its classical description, the final step of the Ehrlich pathway is the reduction of the fusel aldehyde (Table 1) formed in the decarboxylation step by an alcohol dehydrogenase. Indeed, in glucose-grown batch cultures, the amino acids that can be converted via the Ehrlich pathway are almost entirely converted to fusel alcohol and the formation of fusel acid via the oxidation of the aldehydes plays only a minor role (6–8, 10).

Recent studies have shown that the balance between the oxidation and the reduction of the fusel aldehydes depends strongly on cultivation conditions. In aerobic glucose-limited chemostat cultures grown with various amino acids (leucine, methionine, and phenylalanine) as sole nitrogen sources, the amino acids are converted predominantly to fusel acids and only very low concentrations of fusel alcohols are formed (2, 63, 64). In such cultures, the biomass yield on glucose is 40% lower than that of cultures grown on other nitrogen sources, thus implying that a large energy drain is associated with the
formation of fusel acids (2). Given the relatively high levels of hydrophobicity of these weak organic acids, it seems probable that ATP dissipation occurs through classical (29, 49) weak organic acid uncoupling of the plasma membrane pH gradient.

A key role of the cellular redox status in determining the ratio of fusel acid to fusel alcohol production is consistent with the observations discussed above. In glucose-grown batch cultures of S. cerevisiae, growth is predominantly fermentative, and when phenylalanine is the sole nitrogen source, it is converted into a mixture of 90% fusel alcohol (phenylethanol) and less than 10% fusel acid (phenylacetate). Furthermore, fully fermentative dissimilation of glucose under anaerobic conditions (in batch or chemostat cultures) results in the almost complete conversion of phenylalanine to phenylethanol (63, 64).

Especially under anaerobic conditions, the reduction step of the Ehrlich pathway may have a relevant impact on the overall cellular redox metabolism. In S. cerevisiae, excess NADH formed during anaerobic growth can be oxidized by glycerol-3-phosphate dehydrogenase to yield glyceraldehyde (48, 61). However, glyceraldehyde formation from glucose requires ATP and is thus energetically unfavorable. Amino acid catabolism via the Ehrlich pathway may provide an alternative, energy-efficient means for NADH regeneration. Although this possible role of the Ehrlich pathway has not been systematically investigated, nitrogen-limited, respiratortive cultures of S. cerevisiae produce less glyceraldehyde when valine instead of ammonia is supplied as the nitrogen source (5).

Determining the molecular identities of the oxidoreductases involved in the Ehrlich pathway presents a challenge. The S. cerevisiae genome harbors genes for 16 alcohol dehydrogenases, 6 aldehyde dehydrogenases, and at least 2 other broad-spectrum reductases that catalyze the pyridine nucleotide-dependent interconversion of aldehydes and alcohols (Fig. 1). Known roles of the alcohol dehydrogenases are reviewed in reference 10. Using strains containing all possible combinations of mutations affecting the seven AAD genes (encoding putative aryl alcohol dehydrogenases), five ADH genes, and SFA1, Dickinson et al. showed that the final step of the Ehrlich pathway (fusel alcohol formation) can be catalyzed by any one of the ethanol dehydrogenases (Adh1p, Adh2p, Adh3p, Adh4p, and Adh5p) or by Sfa1p (a formaldehyde dehydrogenase) (10). In addition, the NADPH-utilizing aldehyde reductases encoded by YPR1 and GRE2 have been shown to have activity toward 2-methylbutyraldehyde and isovaleraldehyde, respectively (17, 20).

Assigning precise roles to aldo-ketose reductases in many organisms (yeast included) is problematic due to overlapping specificities and the presence of other related enzymes. Transcriptome data for S. cerevisiae grown in glucose-limited chemostats with phenylalanine, methionine, or leucine as the sole nitrogen source were not discriminative, as none of the 16 alcohol dehydrogenase and the 6 aldehyde dehydrogenase transcript profiles could correlate the presence of fusel alcohols and that of fusel acids (2) (Fig. 2).

**EXPORT OF FUSEL ALCOHOLS AND FUSEL ACIDS**

The mechanism by which fusel alcohols are exported from the cells into the culture medium remains unknown. So far, there are no data that link this process to any known membrane transporter in S. cerevisiae, but considering the water-octanol partition coefficients (ranging from 0.76 to 1.36), it is conceivable that export occurs by simple passive diffusion across the lipid bilayer (38). Conversely, the export of fusel acids has recently been shown to involve at least one plasma membrane transporter. The expression of PDR12, encoding an ATP-dependent transporter which is known to be involved in the export of several weak organic acid food preservatives (e.g., benzoate and sorbate) (29), was strongly increased in chemostat cultures utilizing leucine, methionine, or phenylalanine as the sole nitrogen source (2). The phenotypic analysis of a pdr12Δ mutant confirmed the role of Pdr12p in the export of fusel acids formed by the catabolism of leucine, isoleucine, valine, phenylalanine, and tryptophan via the Ehrlich pathway (21). The involvement of an ATP-dependent exporter is fully consistent with the reduced biomass yields observed for chemostat cultures grown on amino acids as the sole nitrogen source that lead to the formation of uncoupling agents (2) (Fig. 2).

**REGULATION OF THE EHRlich PATHWAY**

Although the regulation of many of the individual genes discussed above has already been investigated, these studies did not take into account the context of the Ehrlich pathway. Iraqui et al. identified ARO80 as the transcriptional activator involved in the induction of the ARO9 and ARO10 transaminase and decarboxylase genes by the aromatic amino acid tryptophan, phenylalanine, or tyrosine. A 36-bp upstream activating sequence was shown to be necessary and sufficient to promote transcriptional induction by aromatic amino acids (30). Chromatin immunoprecipitation in chip experiments confirmed the binding of Ara80p to a WRCCGWSATTT RCCG motif uniquely present in the ARO10 and ARO9 promoters (19, 39). It remains unclear whether the strong induction of ARO9 and ARO10 observed in chemostat cultures grown with the Ehrlich pathway precursors leucine and methionine as nitrogen sources (2) is also mediated by Aro80p.

As discussed above, the PDR12-encoded plasma membrane transporter has recently been shown to export branched-chain and aromatic fusel acids (21). The transcriptional induction of PDR12 by the nonphysiological substrates benzoate and sorbate is mediated by War1p, an activator that occupies a cis-acting element in the promoter of PDR12 and becomes active upon phosphorylation (35). A recent genetic screen identified a WAR1 mutant allele carrying mutations corresponding to three amino acid changes. One of these changes proved to be essential for the phosphorylation and binding of War1p to its target sequence. The absence of the phosphorylation of War1p-42 eliminates PDR12 induction and leads to hypersensitivity to sorbate (18). It is as yet unclear whether War1p-mediated induction is also the (sole) mechanism for the transcriptional induction of PDR12 by endogenously produced fusel acids.

In the chemostat-based transcriptome studies by Boer et al. (2), 30 additional genes were shown to follow the transcriptional profile of ARO9, ARO10, and PDR12 (i.e., strong transcriptional upregulation during growth with amino acid precursors of the Ehrlich pathway as the nitrogen source).
However, none of the genes implicated in the Ehrlich pathway (Fig. 1), with the exception of the three mentioned above, were found among the set of 30 genes. This coordinate induction suggests that a general regulatory mechanism participates in the regulation of the Ehrlich pathway. As shown by Boer et al. (2), growth on phenylalanine, leucine, and methionine leads to the relief of nitrogen catabolite repression-mediated regulation. We therefore cannot exclude the involvement of GATA factor-mediated transcriptional regulation of the Ehrlich pathway.

Transcriptional control may not be the only level for the regulation of the Ehrlich pathway. The overexpression of the ARO10 decarboxylase gene from a strong constitutive promoter does not result in increased 3-phenylpyruvate decarboxylase activity during growth on a medium with glucose and ammonium sulfate (63). However, the replacement of either the ammonium sulfate by phenylalanine or of glucose by ethanol leads to a clear increase of 3-phenylpyruvate decarboxylase activity (63). These observations suggest that the functional expression of the ARO10 gene is regulated at a posttranscriptional level in a carbon and nitrogen source-dependent manner. Further research should address the question of whether this regulation involves posttranslational modification of the Aro10p protein and/or other factors.

**FUSEL ALCOHOLS AND ACIDS AS SIGNAL MOLECULES: ROLE OF THE EHRLICH PATHWAY IN THE CONTROL OF CELLULAR MORPHOLOGY**

Growth in the presence of fusel alcohols such as isoamyl alcohol (derived from leucine) induces pseudohypal growth (9, 33). These pronounced morphological modifications are accompanied by an increase in the specific activity of succinate dehydrogenase and an increase of the chitin content (33). These effects are not restricted to isoamyl alcohol, as other reports have shown that for *S. cerevisiae*, the addition of exogenous 2-phenylethanol (4) or 2-indole acetate (also known as the plant hormone auxin) (51) affects morphogenesis and induces invasive growth. These effects indicate a quorum-signal- ing pathway, which links the environmental sensing of autosignal- naling molecules and morphogenesis. No common regulatory circuit has been identified so far. While 2-phenylethanol stimulates morphological modifications by inducing the expression of *FLO11* by a Tpk2p-dependent mechanism (4), transcriptional regulation after exposure to 2-indole acetate involves Yap1p as a key mediator of the response (51). Similarly, in the human fungal pathogens *Candida albicans* and *Candida dub- liniensis*, phenylethanol and isoamyl alcohol produced during growth stimulate biofilm formation (40).

Based on these reports, it thus seems that one of the functions of the Ehrlich pathway is to form quorum-sensing compounds that induce differentiation and participate in the adaptation of yeast cells to environmental changes. However, knowledge of the thermodynamics of the Ehrlich pathway and, more precisely, of the transaminase reactions may provide more insight and propose a new biological role for the Ehrlich pathway.

**THERMODYNAMICS OF THE EHRLICH PATHWAY: A KEY ROLE DURING NITROGEN-LIMITED GROWTH?**

The equilibrium constants of the transamination reactions are often close to unity, like, for instance, that of the phenylalanine–3-phenylpyruvate α-ketoglutarate–glutamate aminotransferase reaction (estimated from data from reference 22). Under physiological conditions, such as those studied by Wu et al. (66), the concentrations of the substrates of this reaction, in this example, α-ketoglutarate (36 μmol liter⁻¹) and phenylalanine (52 μmol liter⁻¹), are often much lower than that of the desired product, glutamate (5,850 μmol liter⁻¹). Since this transamination reaction has an essential role under conditions in which nitrogen concentrations are low, simply increasing the substrate concentration is often not feasible. To solve this thermodynamic problem, cells may use active transport to accumulate the amino acids intracellularly, but this method also has its limitation given the often low solubility of these compounds in water. The only remaining solution is then to maintain very low concentrations of the fourth compound involved in this reaction: 3-phenylpyruvate. The transaminating reaction is followed by decarboxylation, which is generally characterized by a strongly negative Gibbs free-energy change (22). Consequently, when branched-chain, aromatic, or sulfur-containing amino acids are the nitrogen source, the decarboxylation reaction contributes to low intracellular α-keto acid concentrations, thereby pulling the transaminating reactions toward complete utilization of the nitrogen-donating amino acids.

**OUTLOOK**

Branched-chain, aromatic, and sulfur-containing amino acids that are available in malt wort and grape must are important precursors for (off-)flavor formation. In the past decade, many efforts have contributed to a better understanding of the genes involved in this pathway and how their expression is regulated. A desirable trait in beer and wine fermentation is to achieve high levels of branched-chain and aromatic fusel alcohol production in combination with low levels of off-flavors (e.g., methionol). Our growing understanding of the key components of the Ehrlich pathway and their regulation will aid in the design of strains exhibiting specific flavor profiles in foodstuffs as well as in the metabolic engineering of yeast strains for the production of individual Ehrlich pathway products.

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**REFERENCES**


