Identification of a species-specific aminotransferase in *Pediococcus acidilactici* capable of forming α-aminobutyrate

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**Abstract**

During cheese ripening, the bacterial strain *Pediococcus acidilactici* FAM18098 produces the non-proteinogenic amino acid, α-aminobutyrate (AABA). The metabolic processes that lead to the biosynthesis of this compound are unknown. In this study, 10 *P. acidilactici* strains, including FAM18098 and nine *Pediococcus pentosaceus* strains, were screened for their ability to produce AABA. All *P. acidilactici* strains produced AABA, whereas the *P. pentosaceus* strains did not. The genomes of the pediococcal strains were sequenced and searched for genes encoding aminotransferases to test the hypothesis that AABA could result from the transamination of α-ketobutyrate. A GenBank and KEGG database search revealed the presence of a species-specific aminotransferase in *P. acidilactici*. The gene was cloned and its gene product was produced as a His-tagged fusion protein in *Escherichia coli* to determine the substrate specificity of this enzyme. The purified recombinant protein showed aminotransferase activity at pH 5.5. It catalyzed the transfer of an amino group from leucine, methionine, AABA, alanine, cysteine, and phenylalanine to the amino group acceptor α-ketoglutarate. Alpha-ketobutyrate could replace α-ketoglutarate as an amino group acceptor. In this case, AABA was produced at significantly higher levels than glutamate. The results of this study show that *P. acidilactici* possesses a novel aminotransferase that might play a role in cheese biochemistry and has the potential to be used in biotechnological processes for the production of AABA.

**Keywords:** *Pediococcus acidilactici*, α-Aminobutyrate, α-Ketobutyrate, Aminotransferase

**Key points**

- *Pediococcus acidilactici* possesses a species-specific aminotransferase.
- The aminotransferase catalyzes the reversible transfer of an amino group to α-ketobutyrate to form AABA.
- The aminotransferase uses a variety of amino acids as amino group donors.
- The aminotransferase is active at pH 5.5, a pH encountered in cheese.

**Introduction**

During cheese ripening, proteolysis of the caseins occurs. In a series of degradation steps, the proteins are broken down into amino acids. Thereby, lactic acid bacteria (LAB) play an important role as they use the free amino acids for energy production, regulation of the internal pH, regeneration of co-substrates, and protein biosynthesis (Ardö 2006). Recently, it was shown that *Pediococcus acidilactici* FAM18098 catabolized serine and threonine and formed AABA in vitro and in cheese (Irmler et al. 2013; Eugster et al. 2019). Especially, the formation of the non-proteinogenic amino acid, AABA, is quite unusual, and the enzymes involved in the anabolism of this substance are unknown. It was shown
that the compound could be synthesized in metabolically engineered bacteria by using enzymes that either transaminate or reduce α-ketobutyrate (AKB) (Fotheringham et al. 1999; Zhang et al. 2010).

However, AABA formation also occurs naturally, for example in *P. acidilactici*. It has been suggested that in *P. acidilactici*, threonine is the precursor (Irmler et al. 2013; Eugster et al. 2019). Threonine could be deaminated to AKB, which is then aminated or transaminated to AABA. Likewise, serine could be converted via pyruvate to alanine. These reaction steps are similar to the ones used in the engineered bacteria, where either a transaminase or a dehydrogenase catalyzes the final step of AABA synthesis (Fotheringham et al. 2001; Thage et al. 2004a, and Lactobacillus *paracasei* FAM18098 and *P. pentosaceus* FAM19132 were sequenced using PacBio sequencing. The genomes of the other nine *P. acidilactici* strains and eight *P. pentosaceus* strains were determined using Illumina technology. The protocol for genome sequencing and assembly was performed as described previously (Wuethrich et al. 2017).

The annotated genome sequence of strain FAM18098 (Genbank acc. no. GCA_009808095.1) was retrieved from the FTP directory for GenBank assemblies (https://ftp.ncbi.nlm.nih.gov/genomes/all/GCA/009/808/095/GCA_009808095.1_ASM980809v1/) and searched for genes encoding aminotransferases. Similarity searches with genes of interest were then performed by running BLAST searches (Altschul et al. 1990) against the GenBank (https://blast.ncbi.nlm.nih.gov/Blast.cgi) and the KEGG GENES database (https://www.genome.jp/tools/blast/). Furthermore, BLAST searches were carried out against a custom BLAST database containing the newly generated genome assemblies of the *Pediococcus* strains whose accession numbers are listed in Table 1. Clustal Omega was used for multiple sequence alignments (Sievers et al. 2011).

**Materials and methods**

**Bacterial strains, media, and growth conditions**

The bacterial strains used in this study are listed in Table 1. *Pediococcus* were cultivated in MRS broth (de Man et al. 1960) at 30 °C. The capability to produce AABA was assessed by the following procedure: the bacteria were grown at 30 °C for 3 d in a basal broth (pH 7.0 ± 0.2) that consisted of di-potassium hydrogen phosphate (9 g L⁻¹), yeast extract (5 g L⁻¹), casein hydrolysate (2 g L⁻¹), magnesium sulfate (0.2 g L⁻¹), manganese chloride (0.2 g L⁻¹), D-galactose (2 g L⁻¹), 5 mM L-serine, and 5 mM L-threonine.

*Escherichia coli* strains were grown in LB broth (Sambrook et al. 1989) supplemented with ampicillin (50 µg mL⁻¹) if necessary, with shaking (220 rpm) at 37 °C.

**Determination of free amino acids**

High-performance liquid chromatography (HPLC) was used to determine the free amino acids in culture supernatants and enzyme assays. Therefore, 100 µL of the sample was mixed with 1 mL of 100 mM of HCl, 0.5 mL of 0.5 µM of l-norvalin, 0.5 mL of 0.5 µM of piperidin, and 0.5 mL of 10% (w/v) of trichloroacetic acid and incubated for 30 min at 5 °C. After centrifugation, the amino acids present in the supernatant were determined using HPLC as described previously (Wenzel et al. 2018).

**Genome sequencing and bioinformatic analysis**

Genomic DNA was extracted as described in Berthoud et al. (2017), and DNA concentration was determined using the Qubit dsDNA BR Assay Kit (Thermo Fisher Scientific, Reinach, Switzerland). The genomes of *P. acidilactici* FAM18098 and *P. pentosaceus* FAM19132 were sequenced using PacBio sequencing. The genomes of the other nine *P. acidilactici* strains and eight *P. pentosaceus* strains were determined using Illumina technology. The protocol for genome sequencing and assembly was performed as described previously (Wuethrich et al. 2017).

**Cloning and heterologous gene expression**

The primers aat_Fw (5′-ATGAGTGATAAAGTTAAC GC-3′) and aat_Rv (5′-TGCTCGTATGGCTTGGAG G-3′) were designed to amplify the complete aminotransferase-coding gene presented by the locus.
GBO44_RS03485. The primer aat_Rv did not contain the stop codon for cloning reasons. The amplification reactions were performed using the AmpliTaq Gold DNA Polymerase (Thermo Fisher Scientific) and applying the manufacturer’s protocol. Five nanograms of genomic DNA were used in 25 µL reaction volume. The amplification occurred in a thermal cycler using the following program: 10 min at 95 °C; 30 cycles of 15 s at 95 °C, 30 s at 45 °C, and 1 min at 72 °C; 5 min at 72 °C. The reaction products were analyzed using agarose gel electrophoresis.

The DNA fragment obtained from strain FAM18098 was cloned into the pEXP5-CT/TOPO expression vector (Thermo Fisher Scientific) applying the manufacturer’s protocol. The vector contains a nucleotide sequence encoding a poly-histidine tag that is added to the 3′-end of the DNA fragment. The recombinant vector was first transformed into E. coli TOP10. The plasmids extracted from transformants were verified using Sanger sequencing at Microsynth (Balgach, Switzerland). A plasmid containing the DNA fragment in the proper orientation was named pEXP5-CT/aat and transformed into E. coli One Shot BL21(DE3) (Thermo Fisher Scientific) for protein production.

Production and purification of recombinant protein
An overnight culture of E. coli BL21(DE3) harboring the plasmid pEXP5-CT/aat was used to inoculate 200 mL of LB broth supplemented with ampicillin. The culture was incubated with shaking (220 rpm) at 37 °C. When the culture reached an optical density at 600 nm of 0.4, 0.5 mM of isopropyl-beta-D-1-thiogalactoside (IPTG) was added to induce gene expression. The culture was then incubated with shaking at 30 °C for 4 h. The bacteria were harvested by centrifugation (3000×g, 10 min, RT), washed with 20 mM of sodium phosphate (pH 7.4), and frozen at −20 °C.

The His-tagged recombinant protein was purified using Protino Ni-TED 1000 Packed columns (Machery-Nagel,
Enzymatic assays

The ability to transfer the amino group from an amino acid to AKG was studied using a coupled-enzyme assay. In the first reaction, the purified His-tagged protein was incubated with either L-aspartate, L-methionine, L-cysteine, L-leucine, L-isoleucine, L-valine, L-phenylalanine, L-serine, L-threonine, L-alanine, or AABA as amino group donor and AKG as amino group acceptor. In the second reaction, the amount of glutamate, which resulted from the transamination of AKG, was measured using glutamate dehydrogenases. The first reaction (200 µL) consisted of 50 mM of potassium phosphate (pH 5.5) using illustra NAP columns (VWR International GmbH, Dietikon, Switzerland). The protein concentration was determined using the Qubit Protein Assay kit (Thermo Fisher Scientific, Waltham, USA). The purity of the purified protein was evaluated using denaturing gel electrophoresis. The proteins within the gel were visualized using QuickBlue Protein Stain (LuBioScience GmbH, Lucerne, Switzerland).

Results

Pediococcus acidilactici produces the non-proteinogenic amino acids, AABA and ornithine

Pediococcus acidilactici FAM18098 and nine other P. acidilactici strains (Table 1) were incubated in a broth in which AABA formation can be detected to test if AABA formation was strain- or species-specific. Additionally, nine strains of P. pentosaceus were included in this analysis. The analysis of the free amino acids after 3 days of incubation showed that AABA and alanine was formed in all of the samples with P. acidilactici (Fig. 1). Concomitantly, serine and threonine were degraded. In addition, the concentration of phenylalanine was on average 0.2 mM lower than in the non-inoculated medium (data now shown). The strains FAM13881 and FAM18987 produced less alanine and AABA in comparison to the other P. acidilactici strains. Accordingly, the levels for serine and threonine were higher in both strains compared to the other P. acidilactici strains. These amino acid changes were not observed in the samples with P. pentosaceus (Fig. 1).

Furthermore, the culture supernatant of all pediococci contained ornithine whereas arginine was not detected any more (Additional file 1: Table S1).

In silico analysis of aminotransferases

The genome data of strain FAM18098 was searched for genes encoding aminotransferases to follow this hypothesis. Six genes represented by GBO44_RS00380, GBO44_RS02360, GBO44_RS03135, GBO44_RS03485, GBO44_RS04865, and GBO44_RS09300 were annotated to encode amino acid aminotransferases (Fig. 2). BLAST searches were performed to obtain more clarity about the function of these aminotransferases. First, it was found that the two genes GBO44_RS00380 and GBO44_RS09300 were not present in all 10 P. acidilactici strains. In addition, orthologs of GBO44_RS00380 were found in four P. pentosaceus genomes. Second, one of the remaining four aminotransferases, GBO44_RS02360, was
predicted to be a glycine hydroxymethyltransferase, an enzyme that catalyzes the interconversion of glycine and 5,10-methylenetetrahydrofolate to serine and tetrahydrofolate. Third, the two aminotransferases encoded by GBO44_RS03135 and GBO44_RS04865 were predicted to be involved in the metabolism of sulfur-containing amino acids. Finally, no function could be predicted from the sequence alignments of the last aminotransferase, GBO44_RS04865. The nearest protein homolog in the phylogenetically closely related \textit{P. pentosaceus} showed only 46% identity (data not shown), confirming that this enzyme was specific to \textit{P. acidilactici}. The gene was cloned and heterologously expressed in \textit{E. coli} to obtain enzymatic data on this aminotransferase.

**Enzymatic properties of the recombinant aminotransferase**

The primers were designed based on the nucleotide sequence GBO44_RS04865, which is named \textit{aat} in the following. When the primers were tested on genomic DNA of 10 \textit{P. acidilactici} strains, a DNA fragment with a size of approximately 1200 bp was obtained from all \textit{P. acidilactici} strains except for FAM18987 (data not shown). An alignment of the \textit{aat} gene sequences showed that the binding site for the primer \textit{aat} \textit{Rv} displayed multiple mismatches in strain FAM18987 (Additional file 2: Figure S1), explaining why the gene could not be amplified from this strain.

The \textit{aat} gene of FAM18098 was cloned, and the encoded gene product was produced as a His-tagged fusion protein in \textit{E. coli}. The protein could be purified to apparent homogeneity using nickel affinity chromatography (Fig. 3). First, the purified protein was tested for aminotransferase activity using AKG as amino group acceptor. It was found that the enzyme transferred the amino group from leucine, methionine, AABA, alanine, cysteine, and phenylalanine to AKG (Fig. 4).

When the influence of pH and salt concentration was studied, activity was only found at pH 5.5 but not at pH 7.4 or pH 9.1. An increase of the sodium chloride concentration to 4% (w/v) led to a reduction in activity of approximately 22% (data not shown).

Then, various \textalpha{}-keto acids were used as amino group acceptors. As mentioned before, no evidence for the biosynthesis of AKG in \textit{P. acidilactici} can be found based on
annotated genome data. This lack of evidence indicates that the aminotransferases of *P. acidilactici* may utilize other amino group acceptors. The amino group acceptors AKG, AKB, pyruvate, PPA, KMTB, and KIC were compared using leucine, AABA, and alanine as amino group donors. The highest activity was found with leucine, which was set at 100% for the calculation of the relative activities. The column height represents the mean (± SD) of triplicate measurements. The letters above the columns indicate significant differences between amino group donors (P < 0.05).
donors. It was found that AKB, pyruvate, KMTB, KIC, and PPA significantly preferred amino group acceptors to AKG (Fig. 5).

**Discussion**

*Pediococcus acidilactici* FAM18098 was shown previously to produce AABA in broth and cheese (Irmler et al. 2013; Eugster et al. 2019). The present study shows that this phenotype is widely distributed within the species, since other *P. acidilactici* strains also produced the compound and, to our knowledge, has not been reported for other LAB species. Additionally, all *P. acidilactici* strains produced alanine and degraded serine and threonine, suggesting these two latter amino acids are the precursors for alanine and AABA formation. This hypothesis is supported by the observation that the two strains FAM13881 and FAM18987 that produced less alanine and AABA in comparison to the other *P. acidilactici* strains, concomitantly utilized less serine and threonine.

Together with the degradation of serine and threonine, a significant decrease in the concentration of phenylalanine was observed in the samples with *P. acidilactici*. However, since the affected concentrations of phenylalanine were in the micromolar range and since this decrease was not observed in cheese (Eugster et al. 2019), it can be assumed that the degradation of phenylalanine, if at all, contributes only a minimal amount to the biosynthesis of alanine or AABA.

All inoculated samples also contained ornithine after 3 days of incubation. Ornithine results from the arginine deiminase pathway, which involves the sequential action of arginine deiminase and ornithine transcarbamylase that convert arginine via citrulline to ornithine (Liu et al. 1995). Consequently, arginine was not detected anymore at the end of incubation. This finding is in agreement with the observation that the hydrolysis of arginine is a common phenotypic characteristic of *P. acidilactici* and *P. pentosaceus* within the genus *Pediococcus* (Holzapfel et al. 2006).

AABA formation in *E. coli* was achieved by introducing plasmids harboring a threonine deaminase and an aminotransferase into the cells (Fotheringham et al. 1999). Through the concerted action of both enzymes, threonine was converted via AKB to AABA. Analogously, it can be hypothesized that *P. acidilactici* synthesizes AABA from threonine.

When the genomes of the study strains were searched for genes encoding these two enzymes, a gene encoding a bifunctional threonine ammonia-lyase/1-serine ammonia-lyase (data not shown) and genes encoding aminotransferases were found in the genome sequences of *P. acidilactici*. Amongst the aminotransferases, one gene (GB044_RS04865) was present in all *P. acidilactici* strains, of which the function could not predicted using database searches. The present study focused on this enzyme.

The purified, recombinant aminotransferase was active under conditions encountered in a cheese environment (low pH, elevated salt concentrations). Furthermore, the activity of the aminotransferase was found to be significantly higher with pyruvate, AKB, and KMTB than with AKG using leucine as amino group donor (Fig. 5). This is in line with the previously mentioned fact that a pathway for AKG biosynthesis does not exist in *P. acidilactici*. It can be speculated that amino acid-catabolizing enzymes such as the bifunctional serine/threonine deaminase or cystathionine/methionine lyases provide the amino group acceptors necessary for the aminotransferase activity in *P. acidilactici*.

Data about the substrate specificity of LAB aminotransferases is available from three other species. The branched-chain aminotransferase BcAT from *L. lactis* (Yvon et al. 2000) utilized mainly the amino acids, isoleucine, leucine, valine, and methionine, and clearly preferred AKG as amino group acceptor, used KIC to a lesser extent, and displayed only low activity with pyruvate. A branched-chain aminotransferase with similar activities has been purified from *L. paracasei* (Thage et al. 2004b). This enzyme degraded isoleucine, leucine, and valine at similar rates. In addition, the enzyme used AKG at high rates and only low activities were observed with pyruvate and
AKB as amino group acceptors. The AraT aminotransferase from *L. lactis* (Yvon et al. 1997) showed high activity towards leucine and phenylalanine as amino group donor, and AKG and KIC as amino group acceptors. Furthermore, AraT exhibited no activity with valine, isoleucine, cysteine, and alanine. Unfortunately, neither AABA nor AKB were tested as substrates for AraT in this study.

In contrast to the aforementioned branched-chain aminotransferases, the Aat from *P. acidilactici* showed no activity with valine and isoleucine and significantly preferred AKB, pyruvate, KMB, KIC, and PPA as amino group acceptor. In contrast to AraT, the pediococcal aminotransferase newly described in this study showed activity towards cysteine and alanine (Fig. 4). Therefore, this aminotransferase apparently represents a new aminotransferase class within the LAB with promising new properties.

With regard to the fermentative production of AABA using *P. acidilactici* it has to be considered that the structure and chemical properties of alanine are similar to the ones of AABA. This may introduce difficulties in the separation and purification of AABA. Therefore, *P. acidilactici* probably cannot be used directly for industrial processes as long as serine is present in the starting material for fermentation.

However, the bacterium obviously possesses genes with biosynthetic activities that can be used for metabolic engineering purposes. Based on the capability of the Aat aminotransferase to use alanine as amino group donor for AABA synthesis, the following hypothetical pathway that involves the action of three enzymes could be constructed: the bifunctional threonine-serine deaminase of *P. acidilactici* would degrade threonine and serine to AKB and pyruvate, respectively. Pyruvate could then be converted to alanine in a reductive amination reaction involving an alanine dehydrogenase. Interestingly, two genes encoding alanine dehydrogenases are present in the genomes of *P. acidilactici* (data not shown), which could be used for this approach. Finally, the *P. acidilactici* Aat could use AKB and alanine to produce AABA and pyruvate, respectively. The latter reaction product would be recycled to alanine by the alanine dehydrogenase. In theory, this could shift the equilibrium of the transamination reaction towards the reaction product AABA. To test this hypothetical metabolic pathway, it is planned to clone the corresponding genes from *P. acidilactici* and couple the biosynthetic activities by co-expression of the genes in *E. coli*.

**Supplementary information**

Supplementary information accompanies this paper at https://doi.org/10.1186/s13568-020-01034-2.

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**Authors’ contributions**

AW, SI, LW and EE conceived and designed the research. AW, RP, AR and SI conducted experiments. RSS performed statistical analysis. All authors analyzed and discussed data. AW and SI wrote the manuscript. All authors read and approved the final manuscript.

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Corresponding author could provide all the experimental data on valid request.

**Ethics approval and consent to participate**

This article does not contain any studies with human participants or animals performed by any of the authors.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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