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Potential of Fermented Sausage-Associated Lactic Acid Bacteria to Degrade Biogenic Amines During Storage

Jirasak Kongkiattikajorn

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1. Introduction

Biogenic amines (BAs) are organic bases with aliphatic, aromatic or heterocyclic structures that can be found in several foods, in which they are mainly produced by microbial decarboxylation of amino acids, with the exception of physiological polyamines. BAs may be of endogenous origin at low concentrations in non-fermented food such as fruits, vegetables, meat, milk and fish. High concentrations have been found in fermented foods as a result of a contaminating microflora exhibiting amino acid decarboxylase activity (Sillas-Santos, 1996). However, BAs can also trigger human health problems leading to palpitations, hypertension, vomiting, headaches and flushing if food containing high concentrations are ingested. In fermented foods, some lactic acid bacteria (LAB) are able to convert available amino acid precursors into BAs via decarboxylase or deiminase activities during or following ripening processes. For this reason, amino acid catabolism by LAB can affect both the quality and safety of fermented foods (Verges et al., 1999). The amount and type of BAs formed depends on the nature of food and particularly on the kind of microorganisms present. Enterobacteriaceae and certain LAB are particularly active in the production of BA (Beutling, 1996). These amine-producing microorganisms either may form part of the food associated population or may be introduced by contamination before, during or after processing of the food product. Therefore, microorganisms naturally present in raw materials, introduced throughout the processing or added as starter culture can critically influence BA production during the manufacture of fermented products (Bover-Cid et al., 2000).

Nham is a Thai-style fermented pork sausage. Nham ripening generally takes 3-5 days and relies mainly on adventitious microorganisms, which are normally found in raw materials.
LAB produce organic acids from carbohydrates and cause the pH drop, which contribute to Nham formation. *Micrococcus* and *Staphylococcus* are capable of reducing nitrate to nitrite, which is important in producing the characteristic pigmentation. Also, as a source of lipolytic and proteolytic enzymes, they may contribute to flavor production. Therefore, the acidification and the proteolytic process occurring during Nham ripening make the environment particularly favorable for BAs production.

During meat ripening, microbial growth, acidification and proteolysis provide favourable conditions for BA production. The species of lactobacilli most commonly found in meat and meat products are *Lactobacillus sake* and *Lactobacillus curvatus*, which together with *Lactobacillus bavaricus* and *Lactobacillus plantarum* constitute the main microbial flora isolated from fermented sausages. Other bacteria that can be found in relatively high numbers include enterococci (*E. faecalis* and *E. faecium*), which also contribute to the ripening process. However, the presence of enterococci might also reflect a given level of contamination or a poor curing process. Salt-tolerant, nitrate-reducing coagulase-negative staphylococci are also detected in relatively high numbers in ripened meat products. *Staphylococcus xylosus* is the main species found in Spanish fermented sausages, although *S. carnosus* can also be used as a starter culture. BAs can be degraded through oxidative deamination catalyzed by amines oxidase (AO) with the production of aldehyde, ammonia and hydrogen peroxide. Monoamine oxidases (MAOs) and diamine oxidases (DAOs) had been described from some genus of the family Enterobacteriaceae (Yamashita et al., 1993). The potential role of microorganisms with AO activity had become a particular interest in the last few years to prevent or reduce BA accumulation in food products, especially fermented foods. Mah and Hwang (2009) investigated the effect of *Staphylococcus xylosus* to inhibit BA formation in a salted and fermented anchovy. Reduction of tyramine during ripening of fermented sausages was achieved when *Micrococcus varians* was applied as starter culture (Leuschner and Hammes, 1998). Inoculation of *L. plantarum* in sauerkraut effectively suppressed the production of tyramine, putrescine and cadaverine (Kalac et al., 2000).

BAs are physiologically inactivated by AO, which are enzymes found in bacteria, fungi, plant and animal cells able to catalyse the oxidative deamination of amines with production of aldehydes, hydrogen peroxide and ammonia (Cooper, 1997). The sequential action (in the presence of an electron acceptor, such as O₂) of an AO and an aldehyde dehydrogenase leads to the production of an acid and ammonia, which can be used to support microbial growth (Parrot et al., 1987). MAO and DAO activity has been described in higher organisms as well as in bacteria (Murooka et al., 1976, 1979; Ishizuka et al., 1993). There are relevant differences between microbial AO in terms of substrate specificity and location, as stated by Cooper (1997). DAOs can oxidase several BA, such as putrescine and histamine, and their activity can be affected by substrate inhibition; aminoguanidine, antihistaminic drugs and foodborne inhibitors, such as ethanol, carnosine, thiamine, cadaverine and tyramine, reduce their activity (Lehane and Olley, 2000). The potential role of microorganisms involved in food ripenings with AO activity has been investigated with the aim to prevent or reduce the accumulation of BA in foods. Leuschner et al. (1998) tested in vitro the potential amine degradation by many bacteria isolated from foods and, in particular, in strains belonging to the genera *Lactobacillus*, *Lactococcus* and *Staphylococcus*.
*Pediococcus*, *Micrococcus*, as well as to the species *S. carnosus* and *Brevibacterium linens*. They found that this enzymatic activity can be present at very different quantitative levels. Tyramine oxidase activity of several microbial strains was strictly dependent on pH (with an optimum at 7.0), temperature and NaCl, as well as glucose and hydralazine concentration. Moreover, this enzyme was characterised by a higher potential activity under aerobic conditions. Temperature has also an important effect on histamine degradation (Dapkevicius et al., 2000). The highest degradation rate of this amine was observed at 37 °C, but at 22°C and 15 °C, degradation was still considerable. The AO responsible for this degradation has its optimum temperature at 37°C and retains about 50% of its maximum activity at 20 °C (Schomburg and Stephan, 1993). Many *S. xylosus* strains isolated from artisanal fermented sausages in southern Italy showed the ability to degrade BA in vitro (Martuscelli et al., 2000). Among the strains tested, *S. xylosus* S81 completely oxidised histamine, but it degraded, under the adopted conditions, also a part of tyramine. Even if the AO activity in vitro of microorganisms is not quantitatively reproducible in vivo (due to the more severe conditions and, in particular, to the low O₂ tension, pH and salt concentration), reduction of histamine in dry sausages has been observed in the presence of AO-positive staphylococcal starter cultures (Leuschner and Hammes, 1998). In addition, important reduction of the concentration of tyramine and putrescine in the presence of AO positive *S. xylosus* starter cultures have been observed by Gardini et al. (2002). In other words, BA presence in foods is the consequence of a complex equilibrium between the composition of the food and the enzymatic activities of the microbial population. Together with the decarboxylating aptitude of the starter cultures, the presence and relative activity of AO should be considered as an important characteristic in the selection of starter cultures used in the production of fermented foods.

Since Nham is normally consumed without cooking, proper acid production is important to determine the quality and safety of Nham for consumption. Depending on the initial number of contamination, the occurrence of pathogens such as *Salmonella* spp., *Staphylococcus aureus*, and *Listeria monocytogenes* was found specially in Nham with pH higher than 4.6. Due to inconsistency of product quality and ambiguous product safety, improved process of Nham ripening has been developed by using a starter culture technology. Starter cultures are applied to improve and stabilize the quality of the final product and to shorten the ripening period of Nham production. Meanwhile, only little information is available on the effect of starter culture on BA reduction in Nham. Therefore, the objective of this study was to investigate the effectiveness of AOs activity of LAB in inhibiting BA accumulation during Nham ripening. In addition, the change of chemical and microbial properties of Nham during ripening and subsequently during 28 days stored at different temperature was investigated.

2. Materials and methods

2.1. Microbiological analysis

Nham sausages (25 g) were aseptically transferred into a stomacher bag, with 225 mL of peptone (0.85% of sodium chloride added) and then homogenized for two minutes. Further decimal dilutions were made and then 100 μL of each dilution was spread onto agar plates.
Aerobic plate count agar was used to determine total aerobic. BA producing bacteria were counted using differential media supplemented with amino acids as precursor of BAs (Joosten and Northolt, 1989). The media contained of tryptone (0.5%), yeast extract (0.5%), sodium chloride (0.5%), glucose (0.1%), Tween 80 (0.05%), MgSO$_4$•7 H$_2$O (0.02%), CaCO$_3$ (0.01%), MnSO$_4$•4H$_2$O (0.005%), FeSO$_4$•7H$_2$O (0.004%), bromoresol purple (0.006%), amino acid (2%) and agar (2%). The medium contained the precursor amino acids (0.5% tyrosine di-sodium salt and 0.25% L-histidine monohydrochoride, L-ornithine monohydrochoride, L-lysine monohydrochoride, L- phenylalanine, and L-tryptophan), pyridoxal-5-phosphate as a co-decarboxylase factor, growing factors and buffer compounds. All plates were then incubated for 48 h at 37 °C. Bacterial colonies which developed on each agar were then enumerated and expressed as log colony forming unit (CFU)/mL. Only bacterial colonies with purple halo in the differential media were counted as BAs producing bacteria.

2.2. Bacterial strains and growth conditions

Bacterial strains isolated from different fermented sausages were tested. LAB were grown in MRS broth.

2.3. Determination of amine degradation

An overnight culture was harvested, washed with 0.05 M phosphate buffer (pH 7) and the cell pellet resuspended in 0.05 M phosphate buffer supplemented with tyramine, histamine, tryptamine, phenylethylamine, putrescine, and cadaverine. The cell concentration was adjusted to $10^6$, $10^7$ and $10^8$ CFU/mL. The cell suspensions (20 mL) were incubated in a 100 ml flask and shaken at 200 rpm. Samples were taken and added to an equal amount of 1 M HCl. The mixture was boiled for 10 min and centrifugated at 9000 g. The supernatant was frozen at -15°C until HPLC analysis.

2.4. Preparation of starter culture

Starter cultures used in this study were *L. plantarum* + *L. sake*, which were isolated from sausage. A loop from a slant tryptic soy agar culture of each culture was inoculated in 10 mL of tryptic soy broth and incubated at 37 °C for 24 h. Five milliliters of the culture was then transferred to 100 mL of tryptic soy broth and incubated at 37 °C for another 24 h. The culture was centrifuged at 10,000 g for 10 min at 4°C and then washed with broth. Broth was prepared by homogenizing 1 part with 9 part of distilled water, filtered, adjusted to pH 7.0 and then autoclaved at 121°C for 15 min. After centrifugation, the cell pellet was resuspended in sterile fish broth, adjusted to approximately $10^7$ cell/g and used as starter culture in sauce ripening.

2.5. Nham preparation

Minced pork (56%), pieced cooked pork skin (37%), garlic (3.2%), cooked rice (2%), sodium polyphosphate (0.15%), sodium chloride (1.5%) and sodium erythrobate (0.15%) chili (1%)
were mixed thoroughly, packed into a plastic casing and sealed before incubation. Two separated batches of fermented sausage were prepared without starter culture and with different starter cultures (\(L. \text{plantarum} + L. \text{sake}\)) of approximately \(10^7\) cell/g. After incubation the fermented sausages were homogenized for analysis.

### 2.6. Physical and chemical analyses

The pH was measured directly from samples using a microcomputerized pH meter, inserting the electrode into the middle of the sausage. Moisture was determined by drying the sample at 100–105°C until a constant weight was achieved. The color of Nham was determined by Minolta Model DP-301 colorimeter. Color values (L, a, and b) were measured. A white standard tile was used to calibrate the colorimeter (L= 100.01, a= -0.01, b= -0.02) before measurements. Therefore L measures lightness (luminosity) and varies from white to black. The chromatically (a and b values) gives designations of color as follows; a-value measures redness when positive, gray when zero, and greenness when negative, b-value measures yellowness when positive, gray when zero, and blueness when negative. The titratable acidity (TA) determined as total acid was estimated according to AOAC (2000) and expressed as g/100 g dry matter. TCA (trichloroacetic acid)-soluble peptide of the fermented sausages was measured by the method of Greene and Babbitt (1990). The oligopeptide content in the supernatant was determined according to the method of Lowry et al (1951). Results were expressed as \(\mu\)mol/g (dry matter). Free \(\alpha\)-amino acid was measured using TNBS according to Benjakul and Morrissey (1997) Results were expressed as \(\mu\)mol/g (dry matter).

### 2.7. Extraction of amino acids and BAs

10 ML of 10% (w/v) trichloroacetic acid (TCA) were added to 3 g-samples, and homogenization of the mixture was effected via shaking for 1 h. The extract was then filtered through Whatman No. 1 filter paper. To remove any fat, the samples were kept at -20°C for 1 d, and then centrifuged at 7000 g for 15 min. The supernatants were collected and filtered through a 0.25 \(\mu\)m membrane filter.

### 2.8. Determination of BAs

Amines were determined by the high-performance liquid chromatography (HPLC) method described by Hernández-Jover et al. (1996). The method is based on the formation of ion pairs between amines extracted with 0.6 M perchloric acid from 5 to 10 g of sample, and octanesulphonic acid present in the mobile phase. Separation is preformed using a reversed phase column, then a postcolumn derivatization with \(o\)-phthalaldehyde (OPA) is followed by spectrofluorimetric detection. The method allows one to quantify, by an external standard procedure, 6 BAs, i.e., tyramine, histamine, tryptamine, phenylethylamine, putrescine, cadaverine. Samples for BA determination were stored at -15°C until required.
2.9. Determination of amino acids

Free amino acids (FAAs) in samples were determined using HPLC according to the method proposed by Rozan et al. (2000). A 20 μL aliquot of amino acid standard and digested sauce samples were transferred into vials and dried under vacuum. Then 20 μL of drying reagent containing methanol, water and triethylamine (ratio 2:2:1 v/v) was added. Then 20 μL of derivatizing reagent containing methanol, triethylamine, water and phenylisothiocyanate (PITC) (ratio 7:1:1:1 v/v) was added. The derivatized samples were then dissolved in 100 mL of buffer A that was used as mobile phase for HPLC. A Purospher® STAR RP-18e, 5 μm column was used with buffer A (0.1 M ammonium acetate, pH 6.5) and buffer B (0.1 M ammonium acetate containing acetonitrile and methanol, 44:46:10 v/v, pH 6.5) as mobile phase set for linear gradient at the flow rate of 1 mL/min. The injected sample volume was 20 μL and monitored at 254 nm of wavelength.

2.10. Statistical analysis

Data was analysed by one-way ANOVA and differences among treatment means were determined by Duncan’s new multiple-range test.

3. Results and discussion

The effect of starter cultures of LAB on BAs and FAAs content was examined during the ripening process of Nham sausages. Microbial counts, pH and proteolysis-related parameters were also studied. The occurrence of amino acid-decarboxylase activity in 7 strains of LAB isolated from Nham sausages was investigated.

<table>
<thead>
<tr>
<th>Starter culture</th>
<th>Percent degradation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Tryptamine</td>
</tr>
<tr>
<td>Lactobacillus curvatus 1271</td>
<td>0</td>
</tr>
<tr>
<td>Lactobacillus farcininis 1452</td>
<td>0</td>
</tr>
<tr>
<td>Lactobacillus kandleri 2439</td>
<td>0</td>
</tr>
<tr>
<td>Lactobacillus kefir 2045</td>
<td>0</td>
</tr>
<tr>
<td>Lactobacillus plantarum 9825</td>
<td>0</td>
</tr>
<tr>
<td>Leuconostoc malti 7412</td>
<td>0</td>
</tr>
<tr>
<td>Lactobacillus pentosus 7054</td>
<td>0</td>
</tr>
<tr>
<td>Lactobacillus reuteri 7498</td>
<td>0</td>
</tr>
<tr>
<td>Lactobacillus sake 4127</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 1. Strains exhibiting the potential to degrade BAs in a buffer system within 24 h at 30°C

The presence of BAs in a decarboxylase synthetic broth was determined by high performance liquid chromatography with OPA derivatization. Among the 9 LAB strains tested, 5 lactobacilli (in particular, *L. curvatus*) were amine producers and *L. plantarum* and *L.
sake, were non-amine forming strains. The ability of AO exhibiting strains of LAB to degrade amine in vivo during sausage ripening was investigated.

![Graph showing a* values during ripening of Nham control and Nham with starters at different temperatures.](image1)

**Figure 1.** a* Value during ripening of Nham control at 25°C (●), 30°C (●), 37°C (▲) and Nham with starters (L. plantarum + L. sake) at 25°C (○), 30°C (●), 37°C (●).

Fig. 1 showed a* values represent red color of Nham during ripening time and temperature at 25°C, 30°C and 37°C, respectively. The results showed a value increased according to ripening and the a value of Nham control at 72 hours 37°C was higher than the other sample.

![Graph showing b* values during ripening of Nham control and Nham with starters at different temperatures.](image2)

**Figure 2.** b* Value during ripening of Nham control at 25°C (●), 30°C (●), 37°C (▲) and Nham with starters (L. plantarum + L. sake) at 25°C (○), 30°C (●), 37°C (●).
Fig. 2 shows $b^*$ values represent yellow color of Nham during ripening time and temperature at 25°C, 30°C and 37°C, respectively. The results showed $b$ value decreased according to storage and the $b$ value of Nham with starters was lower than that of Nham control.

**Figure 3.** $L^*$ Value during ripening of Nham control at 25°C (●), 30°C (●), 37°C (▲) and Nham with starters ($L. \text{plantarum} + L. \text{sake}$) at 25°C (×), 30°C (●), 37°C (●).

**Figure 4.** pH during ripening of Nham control at 25°C (●), 30°C (●), 37°C (▲) and Nham with starters ($L. \text{plantarum} + L. \text{sake}$) at 25°C (×), 30°C (●), 37°C (●).
Fig. 3 is represent L*values represent white color of Nham during ripening time and temperature at 25°C, 30°C and 37°C, respectively. The results showed L* value increased according to storage during 72 hour of ripening.

Fig. 4 shows that the initial pH of Nham samples ranged from 5.9 to 6.1. It then gradually decreased throughout the ripening process and there was significant difference at each time of sampling (P <0.05). The pH value reached 4.1 to 4.8 at the end of ripening (hour 72). However, there was significant difference (P<0.05) between the pH of Nham control and samples inoculated with starter cultures after 48 hour of ripening.

**Figure 5.** Total acid content during ripening of Nham control at 25°C (●), 30°C (●), 37°C (▲) and Nham with starters (L. plantarum + L. sake) at 25°C (∗), 30°C (∗), 37°C (∗).

Fig. 5 shows that the initial total acid content of Nham samples ranged from 0.5 to 0.55. It then gradually increased throughout the ripening process and there was significant difference at each time of sampling (P <0.05). The total acid content of Nham control and Nham with starters reached 0.95% to 1.57% and 1.04 % to 1.32% at the end of ripening (hour 72). However, there was significant difference (P<0.05) between the total acid content of Nham control and samples inoculated with starter culture after 48 hour of ripening. The results was shown that Nham control fermented at 37°C contained total acid content higher than the other Nham samples.

Fig. 6 shows that TCA-soluble peptide content of Nham samples, the initial content was 9.02 μmol/g dry matter. It then gradually increased throughout the ripening process. The TCA-soluble peptide content of Nham control and Nham with starters reached 23.6 to 87.2 μmol/g dry matter and 24.1 % to 65.2 μmol/g dry matter, respectively, at the end of ripening (hour 72). However, there was not significant difference (P<0.05) between the TCA-soluble peptide content of Nham control and samples inoculated with starter culture after 48 hour
Figure 6. TCA-soluble peptide content during ripening of Nham control at 25°C (●), 30°C (●), 37°C (▲) and Nham with starters (L. plantarum + L. sake) at 25°C (◆), 30°C (○), 37°C (●).

The results showed that Nham control fermented at 37°C contained TCA-soluble peptide content higher than the other Nham samples after ripening for 72 hours.

Figure 7. Free α-amino acid content during ripening of Nham control at 25°C (●), 30°C (●), 37°C (▲) and Nham with starters (L. plantarum + L. sake) at 25°C (◆), 30°C (○), 37°C (●).

Fig. 7 shows that free α-amino acid content of Nham control samples and Nham with starters, the initial content were 216.2 mmol/g dry matter and 203.7 mmol/g dry matter,
respectively. It then gradually increased throughout the ripening process. The free $\alpha$-amino acid content of Nham control and Nham with starters reached 275.3 to 351.6 mmol/g dry matter and 262.4 to 302.2 mmol/g dry matter, respectively, at the end of ripening (hour 72). However, there was not significant difference ($P<0.05$) between the free $\alpha$-amino acid content of Nham control and samples inoculated with starter culture during ripening at 25°C. The results was shown that Nham control fermented at 37°C contained free $\alpha$-amino acid content higher than the other Nham samples throughout the ripening process.

The differences between Nham in counts of LAB during ripening are shown in Fig. 8. LAB in Nham with starters was increase until the 72 h of ripening. Counts of LAB in Nham with starters (8.7 log CFU/g) were higher ($P<0.05$) than in Nham control (7.7 log CFU/g).

Fig. 9 shows that cadaverine content of Nham samples, the initial content was 14.89 mg/kg dry matter. It then gradually increased throughout the ripening process. The cadaverine content of Nham control and Nham with starters reached 86.2 to 98.7 mg/kg dry matter and 42.4 to 51.6 mg/kg dry matter, respectively, at 72 hour of ripening. However, there was not significant difference ($P<0.05$) between the cadaverine content of Nham with starters during ripening at 25°C and 30°C. The results was shown that Nham control fermented at 37°C contained cadaverine content higher than the other Nham samples throughout the ripening process.

Fig. 10 shows that putrescine content of Nham samples, the initial content was 23.7 mg/kg dry matter. It then gradually increased throughout the ripening process and there was significant difference at each time of sampling ($P<0.05$). The putrescine content of Nham control and Nham with starters reached 115.4 to 242.6 mg/kg dry matter and 65.2 to 98.4 mg/kg dry matter, respectively, at 72 hour of ripening. However, there was not significant
Figure 9. Cadaverine content during ripening of Nham control at 25°C (●), 30°C (○), 37°C (▲) and Nham with starters (L. plantarum + L. sake) at 25°C (☆), 30°C (◆), 37°C (★).

Figure 10. Putrescine content during ripening of Nham control at 25°C (●), 30°C (○), 37°C (▲) and Nham with starters (L. plantarum + L. sake) at 25°C (☆), 30°C (◆), 37°C (★).

difference (P<0.05) between the putrescine content of Nham with starters during ripening at 25°C and 30°C. The results was shown that Nham control fermented at 37°C contained putrescine content higher than the other Nham samples throughout the ripening process.
Fig. 11 shows that tyramine content of Nham samples, the initial content was 5.63 mg/kg dry matter. It then gradually increased throughout the ripening process and there was significant difference at each time of sampling ($P<0.05$). The tyramine content of Nham control and Nham with starters reached 17.6 to 46.4 mg/kg dry matter and 16.3 to 27.8 mg/kg dry matter, respectively, at 72 hour of ripening. However, there was not significant difference ($P>0.05$) between the tyramine content of Nham with starters during ripening at 30°C and 37°C and Nham control and Nham samples inoculated with starter culture during ripening at 25°C. The results was shown that Nham control fermented at 37°C contained tyramine content higher than the other Nham samples after 48 hour of the ripening process.

The effect of temperature on BA content was evaluated (Fig. 6-9). The storage temperature of Nham with starters at 30°C and 37°C were shown higher BA oxidation comparing Nham control, a low content was observed at 25°C. This suggested that at ripening temperature of 30°C and 37°C, a strong oxidation of the AO activity of the starters was evident, whereas at 25°C activity was low for amino acid decarboxylase for lysine (precursor of putrescine) and tyrosine (precursor of tyramine) in Nham control.

Fig. 12 showed $a^*$ values represent red color of Nham during stored at 15°C, 4°C and 25°C. The initial $a^*$ values of Nham control ranged from 7.0 to 8.4. The results showed $a^*$ value increased according to 4 week storage for storage temperature at 15°C and 25°C. However, there was significant decrease ($P<0.05$) between the $a^*$ value of 4°C storage of the initial 1 week storage and after 4 week of storage. For Nham with starters, the initial $a^*$ values ranged from 7.5 to 8.5. The $a^*$ values of Nham with starter decreased according to 4 week storage for storage temperature at 4°C and 15°C. However, there was not significant difference ($P>0.05$) between the $a^*$ value of 25°C storage of the initial 1 week storage and after 4 week of storage.
Figure 12. *a* Value during storage of Nham control at 25°C (●), 30°C (■), 37°C (▲) and Nham with starters (L. plantarum + L. sake) at 25°C (×), 30°C (●), 37°C (▲).

Figure 13. *b* Value during storage of Nham control at 4°C (●), 15°C (●), 25°C (▲) and Nham with starters (L. plantarum + L. sake) at 4°C (×), 15°C (●), 25°C (▲).

Fig. 13 showed *b* values represent yellow color of Nham during stored at 15°C, 4°C and 25°C. The initial *a* values of Nham control ranged from 5.0 to 6.1. The results showed *a* value increased according to 4 week storage for storage temperature at 4°C and 25°C. However, there was significant decrease (*P*<0.05) between the *b* value of 15°C storage of the initial 1 week storage and after 4 week of storage. For Nham with starters, the initial *a*
values ranged from 5.5 to 6.2. The $b^*$ values of Nham with starter increased according to 4 week storage for storage temperature at 4°C and 25°C. However, there was significant decrease ($P<0.05$) between the $b^*$ value of 15°C storage of the initial 1 week storage and after 4 week of storage.

Fig. 14. $L^*$ Value during storage of Nham control at 4°C (●), 15°C (●), 25°C (▲) and Nham with starters ($L.\ plantarum$ $+ L.\ sake$) at 4°C (●), 15°C (●), 25°C (●).

Fig. 14 showed $L^*$ values represent white color of Nham during stored at 15°C, 4°C and 25°C. The initial $a^*$ values of Nham control ranged from 55.1 to 56.4. The results showed $L^*$ value decreased after 2 week storage and then increased after 3 week storage for each storage temperature. However, there was not significant difference ($P<0.05$) between the $L^*$ value of 25°C storage of the initial 1 week storage and after 4 week of storage. For Nham with starters, the initial $L^*$ values ranged from 55.0 to 55.6. The $L^*$ values of Nham with starter decreased after 2 week storage then the $L^*$ values increased after 3 week storage for each storage temperature and after 4 week storage at 15°C and 25°C, the $L^*$ value was significant increased. However, there was no significant difference ($P<0.05$) between the $L^*$ value of 4°C storage of the initial 1 week storage and after 4 week of storage.

Fig. 15 shows that the initial pH of Nham samples ranged from 4.3 to 4.5. It then gradually decreased throughout the storage. The pH value reached 4.1 to 4.5 at 4 week of storage. The pH values at each storage temperature of Nham with starter were higher than Nham control at each time of sampling. The results was shown that pH value of Nham control stored at 25°C was lower than the other Nham samples throughout the storage process.
Fig. 15. pH during storage of Nham control at 4°C (●), 15°C (●), 25°C (▲) and Nham with starters (L. plantarum + L. sake) at 4°C (●), 15°C (●), 25°C (●).

Fig. 16. Total acid content during storage of Nham control at 4°C (●), 15°C (●), 25°C (▲) and Nham with starters (L. plantarum + L. sake) at 4°C (●), 15°C (●), 25°C (●).

Fig. 16 shows that the initial total acid content of Nham samples ranged from 1.1 to 1.7. It then gradually increased throughout the ripening process and there was significant difference at each time of sampling ($P < 0.05$). The total acid content of Nham control and Nham with starters reached 1.14% to 2.72% and 1.04 % to 2.32% at 4 week of storage, respectively. However, there was not significant difference ($P < 0.05$) between the total acid...
content of Nham control stored at 4°C and 15°C and Nham with starters stored at 15°C and from the results, the total acid content of Nham with starters stored at 4°C was not significant difference ($P<0.05$) during storage process. The total acid content of Nham control stored at 25°C was higher than the other Nham samples throughout the storage process.

**Figure 17.** TCA-soluble peptide during storage of Nham control at 4°C ($\bullet$), 15°C ($\bullet$), 25°C ($\triangle$) and Nham with starters ($L.~plantarum + L.~sake$) at 4°C ($\blacklozenge$), 15°C ($\times$), 25°C ($\circ$).

Fig. 17 shows that the TCA-soluble peptide of Nham control and Nham with starters ranged from 45.2 to 98.4 and 46.3 to 79.6 $\mu$mol/g dry matter, respectively. Nham control and Nham with starters stored at 25°C showed gradually increased throughout the storage process and there was significant difference at each time of sampling ($P<0.05$). The TCA-soluble peptide of Nham control and Nham with starters stored at 4°C and 15°C. However, there was not significant difference ($P<0.05$) between the TCA-soluble peptide of Nham control and samples inoculated with starters culture throughout the storage process at 4°C and 15°C. From the results, the TCA-soluble peptide of Nham control stored at 25°C was higher than the other Nham samples throughout the storage process.

Fig. 18 shows that free $\alpha$-amino acid content of Nham control samples and Nham with starters, the initial ranged from 342.3 to 603.4 and 346.6 to 507.2 mmol/g dry matter, respectively. It then gradually increased throughout the storage process at 15°C and 25°C and there was significant difference at each time of sampling ($P<0.05$). The free $\alpha$-amino acid content of Nham control and Nham with starters reached 375.2 to 1867.6 mmol/g dry matter and 359.4 to 1252.4 mmol/g dry matter, respectively, at 4 week of storage. However, there was not significant difference ($P<0.05$) between the free $\alpha$-amino acid content of Nham control and samples inoculated with starters during storage at 4°C. The results was shown that Nham control stored at 25°C contained free $\alpha$-amino acid content higher than the other Nham samples after 3 week storage.
The differences between Nham in counts of LAB during ripening are shown in Fig. 19. LAB in Nham with starters was increase until the 4 week of storage. Counts of LAB in Nham with starters stored at 25°C (9.4 log CFU/g) were higher ($P < 0.05$) than in Nham control stored at 25°C (9.1 log CFU/g). LAB counts in Nham increased steadily during storage, the dependence of the LAB counts of Nham control and Nham with starters on ripening at each
Potential of Fermented Sausage-Associated Lactic Acid Bacteria to Degrade Biogenic Amines During Storage

In the present study in Nham are concerned, total LAB counts in Nham with starters on 3 week of storage were higher \((P < 0.05)\) in comparison with the Nham control produced at the same storage temperature. An increase of LAB in Nham with starters until 3 week of storage and consecutive increase till 4 week of storage was significant. LAB of Nham with starters produced increase steadily during ripening and stored at different temperatures, however, at 4°C storage, LAB counts storage was not different significant.

Fig. 20 shows that cadaverine content of Nham samples, the initial ranged from 43.7 to 58.2 mg/kg dry matter. In Nham control stored at 15°C and 25°C, it then gradually increased throughout the storage process. The cadaverine content of Nham control and Nham with starters reached 58.4 to 91.2 mg/kg dry matter and 41.6 to 47.3 mg/kg dry matter, respectively, at 72 hour of storage. However, there was not significant difference \((P<0.05)\) between the cadaverine content of Nham with starters during stored at 4°C. The results was shown that Nham control stored at 25°C contained cadaverine content higher than the other Nham samples throughout the storage process and there was significant decreased \((P<0.05)\) in the cadaverine content of Nham with starters stored at 25°C for 4 week.

Fig. 21 shows that putrescine content of Nham control and Nham with starters, the initial ranged from 124.6 to 176.3 mg/kg dry matter and 126.2 to 98.3 mg/kg dry matter. Nham control stored at 4°C, 15°C and 25°C gradually increased throughout the storage process. The putrescine content of Nham control and Nham with starters reached 175.3 to 339.4 mg/kg dry matter and 122.6 to 129.3 mg/kg dry matter, respectively, at 4 week of storage. However, there were significant increase \((P<0.05)\) between the putrescine content of Nham with starters at each storage temperature for 2 week and then the putrescine content
decreased and there was not significant difference ($P<0.05$) between the putrescine content of Nham with starters after 3 week storage. The results were shown that Nham control stored at 25°C contained putrescine higher than the other Nham samples throughout the storage process.

Figure 21. Putrescine content during storage of Nham control at 4°C (●), 15°C (♦), 25°C (▲) and Nham with starters (L. plantarum + L. sake) at 4°C (◆), 15°C (★), 25°C (★).

Figure 22. Tyramine content during storage of Nham control at 4°C (●), 15°C (♦), 25°C (▲) and Nham with starters (L. plantarum + L. sake) at 4°C (◆), 15°C (★), 25°C (★).
Fig. 22 shows that tyramine content of Nham control and Nham with starters, the initial ranged from 18.2 to 65.3 mg/kg dry matter and 19.2 to 22.4 mg/kg dry matter. Nham control stored at 15°C and 25°C gradually increased during storage process. The tyramine content of Nham control and Nham with starters reached 25.2 to 198.6 mg/kg dry matter and 21.4 to 27.6 mg/kg dry matter, respectively, at 4 week of storage. There was not significant difference (P<0.05) between the tyramine content of Nham control stored at 4°C and Nham with starters at each storage temperature during storage time. The results was shown that Nham control stored at 25°C contained tyramine higher than the other Nham samples throughout the storage process.

One of the most important factors influencing BA formation in Nham is starter culture (Maijala et al., 1995). Increase of LAB starters culture in Nham resulted in overgrowth more than the microflora and LAB producing BAs in Nham control and caused decrease in BA contents in Nham during ripening and storage. A higher amount of BAs was formed in the Nham control than in starters culture-ones. However, strains of the starters showed lower decarboxylase activity (lower total free amino acid content in Nham) in comparison with the Nham control. Moreover, from the fact that BA production increased in Nham control after the ripening was finished and Nham was stored at the 15°C and 25°C, which coincided with the temporary increase of total LAB, the presence of spontaneous decarboxylating microflora can be inferred, and the refrigerated storage should be recommended. Simultaneously, higher concentration of BAs was found in Nham fermented at high 30°C and 37°C as compared to 25°C at the end of ripening. However, as regards to the strongly hypothetical effects of some substances in the Nham spicing mixtures in connection with the BA formation, more research is needed. Decarboxylase activities present in microflora in Nham are influenced by pH, temperature (Gardini et al., 2001; Silla-Santos, 1996; Suzzi and Gardini, 2003). The decarboxylation of FAAs to BAs was found to be inhibited by low pH (Gardini et al., 2001). Though amino acid decarboxylase activities usually have acid pH optimum (Gale, 1946), the pH rise could favour the cell yield and growth (Maijala, 1994) of decarboxylase-positive microflora.

3.1. BAs contents of Nham

Occurrence of toxic compounds such as BAs is favoured by a high concentration of substrates (i.e., free amino acids) together with environmental and technological factors (e.g. NaCl content, chemico-physical variables, hygienic procedure adopted during production) promoting microbial growth and the decarboxylase activity of microorganisms (Silla-Santos, 1996). In this study, a high correlation among total BAs and total FAAs content was observed. Temperature markedly influences the formation of BAs, and at 15°C decarboxylases might be still active (Bover-Cid et al., 2001). During storage, the more temperature exceeds 14–15°C the more decarboxylase activities might release BAs from FAAs. In this respect, processing procedures for Nham based on low salt addition, high ripening temperatures (over 20°C), may favour proteolytic and decarboxylase activities. The high values of cadaverine, putrescine and tyramine detected in some Nham, may be ascribed to inadequate microflora and LAB producing BAs reduction occurring in some Nham control (Fig. 9- Fig. 11 and Fig. 20- Fig. 22).
The toxicological level of BAs depends on the individual characteristics and the presence of other amines (Brink et al., 1990; Halasz et al., 1994). Toxic doses of tyramine in foods were reported in the range 100–800 mg/kg, but average amounts of tyramine detected in analysed samples (Fig. 9–Fig. 11 and Fig. 20–Fig. 22) were below this range, even if in case of a few samples, the 100 mg/kg value was exceeded. Putrescine has been regarded as not toxic by itself, but as a potentiator for the toxic effect of tyramine and histamine if present (Hui and Taylor, 1985). However, it was probable to demonstrate significant relationship between the concentration of a specific FAA and its corresponding BA in meat products (Eerola et al., 1998). Fig. 9–Fig. 11 and Fig. 20–Fig. 22 shows the BAs content of Nham evidence the effect of starters on the decrease of the BAs occurrence in Nham after ripening and storage. Histamine was always below the minimum detectable, in spite of the abundance of their precursors (histidine) released during the process; phenylethylamine was also not detected. The concentration of tyramine was high in Nham control while low concentration, of their precursors (tyrosine) released during the process. Moreover, tryptamine resulted absent in all the investigated samples. The sum of vasoactive BAs, VBA; (tyramine, phenylethylamine, histamine and tryptamine) lower than 200 mg/Kg has been suggested by Eerola et al. (1998) as a quality index (VBA index) for ripened meat products. It is interesting to note that the computed VBA index of Nham with starters with differently processed resulted appreciable samples (3.70 ± 2.46 mg/Kg). These results could be related to the specific characteristics of the product as well as to the process conditions adopted that could, in general, have limited the growth and activity of amino acid decarboxylase positive microorganisms (Suzzi and Gardini, 2003). Cadaverine, putrescine and tyramine were found in high amounts in Nham control. However, the occurrence of BAs in Nham control, and after the storage could be due to the microflora and LAB producing BAs that could have favoured their formation during ripening and storage. During ripening and storage of Nham control, putrescine and cadaverine show a marked increase with high amounts of their precursor, arginine and lysine, respectively, were detected. In fact, arginine may generate putrescine both via arginine deiminase pathway (ADI) leading to ornithine (Montel and Champomier, 1987) and their subsequent decarboxylation to putrescine, and via arginine decarboxylation to agmatine followed by deamination to putrescine and removal of urea (Moreno-Arribas et al., 2003). It seems reasonable to postulate that the large amounts of arginine could be the source of putrescine, which subsequently may be converted in spermine and spermidine by transamination reactions (Lehnninger et al., 1999).

3.2. FAAs contents of Nham

FAAs were reported in Table 2-Table 5 as net amounts (mmol/g dry matter) in order to investigate the differences in contents due to starters in Nham during ripening and storage. FAAs were compared to evaluate if the extended storage times gave a similar increase in all of them or different patterns were detectable. Most single FAAs increased during ageing with particular reference to the lipophylic ones; a rise in lipophyllic valine, phenylalanine and tryptophan processed following a traditional prolonged way (Ruiz et al., 1999). In the present study, stored Nham showed a FAA pattern enriched with glutamic acid, alanine,
arginine, cysteine, serine, threonine and glycine, most FAAs displayed a rise during the extended storage. Arginine found in the most stored Nham was increase, due to changing of its content by proteolysis; and rise in arginine in stored Nham control was higher than stored Nham with starters. Arginine hydrolysis could be hydrolysed via the arginine deiminase pathway (ADI) leading to ammonia and ornithine. It seemed reasonable to postulate that ADI pathway enzymes (arginine deiminase and ornithine transcarbamylase) could be still active during storage times. Arginine catabolism, may be regarded as a source of the BA putrescine both via ADI ornithine generation (Montel and Champomier, 1987) and subsequent decarboxylation to putrescine, and via arginine decarboxylation to agmatine followed by deamidation to putrescine and removal of urea (Moreno-Arribas et al., 2003). The presence in Nham of environmental conditions suitable for decarboxylase activities together with large amounts of arginine may be consistent with the increase in putrescine.

The evolution during incubation/storage of the total free amino acid content, in both the Nham control and after inoculation with either of the two Lactobacillus strains selected, is shown in Table 2- Table 5, and encompassed 17 different amino acids. The control Nham showed the highest concentration of total amino acids at a 5% level of significance. The contents of total amino acids in Nham inoculated with L. plantarum and L. sake, increased throughout time, but at lower rates than the control. The contents of free amino acids and BAs in control and experimental Nham increase significantly throughout incubation/storage. However, specific lactic acid strains of the Lactobacillus genus can effectively prevent BAs from building-up excessively, putrescine (for quantitative reasons, owing to its level). This may lead to a favourable contribution to public health, especially in regions where Nham is frequently included in the diet. To have an overall evolution index of the proteases action in the Nham during processing the TCA-soluble peptide was evaluated (Fig. 6 and Fig. 17) (Toldr, 2005). More intense proteolytic activity occurred in the Nham control. The TCA-soluble peptide values of Nham control are quite high compared to those generally observed in other Nham with starters. This could be attributed to the microflora in Nham control slightly higher proteolytic activity during the process, in comparison with those Nham applied with starters. Proteolysis contributes to texture by breakdown of the muscle structure (Monin et al., 1997).

Table 2- Table 5 show the FAAs content of Nham during ripening and storage arginine and glutamic acid were the FAAs most representative; after ripening and storage a marked increase of alanine was observed. Table 6 shows the effect of the starters treatment on the evolution of the FAAs pattern of the Nham investigated during the ripening and storage: a significant increase in the concentration of all FAAs with respect to their initial occurred in Nham control and Nham with starters, resulting from the aminopeptidases activity of meat (Toldr, 2006) as well as microbial proteases (Dur et al., 2004; Molina and Toldrá, 1992; Rodriguez et al., 1998; Scannell et al., 2004). Moreover, starters in Nham seems to affect the production of some amino acids (Table 6). A lower concentration of lysine, threonine, glycine and proline was detected, after storage, in Nham processed. Arginine was the most abundant amino acid in all the final products, and its level was significantly higher in Nham control than in those subjected with starters to Nham. At the end of the ripening step,
cysteine was also present in a relative higher concentration in Nham control, whereas
significant larger amounts of proline, lysine, histidine, serine and threonine were reached in
Nham control samples. The different profile of FAAs observed in Nham control and Nham
with starters may be due to a different evolution of reactions and processes involving both
production and consumption of amino acids that occur simultaneously during the various
steps of the ripening process and storage and whose combined effects could give rise to an
increase or, on the contrary, to a decrease of their concentration. The aminopeptidase
activity is considered the main process implied in the FFA release in meat. Moreover, free
amino acids concentration could be decreased either by chemical and enzymatic reactions
where they act as substrates leading to the formation of secondary products (Ruiz et al.,
1999; Ventanas et al., 1992) and/or by microbial amino acid decarboxylase activity with
consequent BA production (Virgili et al., 2007).

In Nham control, an effect due to higher concentration of decarboxylase than that of Nham
with starters, thus, their reaction with the free amino acids causing an increase of their BA
concentration in these samples.

The ripened taste could be related to lysine and glutamic acid, while isoleucine and aspartic
acid are implied in acid taste and unpleasant aroma (Buscailhon et al., 1994; Flores et al.,
1998). In this study, the increase in concentration of lysine and glutamic acid was observed.
The changes in the contents of free amino acids observed in fermented sausages during
ripening are given in Table 2. The total free amino acid contents of the Nham control and
Nham with starters constituted 212.7–216.4 mmol/g and 197.2–203.4 mmol/g dry matter,
respectively (before ripening) on 0 day. An increase in the content of amino acids of Nham
control and Nham with starters was observed and ranged between 275.2–349.8 mmol/g and
259.8–300.3 mmol/g dry matter during the ripening on day 3, and a further increase up to
the range of 377.6–1851.7 mmol/g and 348.1 nmol/g–1256.0 mmol/g dry matter of total free
amino acids was observed during storage at 4°C-25°C of Nham control and Nham with
starters (4 weeks). The highest total free amino acid concentration of 1867.2 mmol/g was
observed with Nham control stored at 25°C for 4 week, whereas the lowest total free amino
acid concentration of 359.6 mmol/g was observed with Nham with starters stored at 25°C
for 4 week. The hydrolysis of meat proteins generates polypeptides that can be further
degraded to smaller peptides and free amino acids. This degradation can be produced by
endogenous and microbial enzymes (De Masi et al., 1990; Hughes et al., 2002; Molly et al.,
1997). The increase in the total free amino acid concentration was detected in all batches
(Hierro et al., 1999, Bruna et al., 2000, Bolumar et al., 2001 and Hughes et al., 2002).

The main differences in the content of total free amino acids among batches were detected
during 72 hour of ripening and during 4 week of storage. The amino acids in which
differences, which were primarily responsible for the increase in total free amino acids
during ripening, were observed were Glu (glutamic acid), Ala (alanine) and Arg (arginine)
in Nham control and Nham with starters. Mateo et al. (1996) reported an increase in the total
free amino acid content during the ripening. The change occurred during ripening and
storage process indicating that the highest enzymatic activity took place during these stages
Table 2. Amino acid content of Nham without and with starter cultures during ripening at different temperatures.

<table>
<thead>
<tr>
<th></th>
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<th></th>
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<tbody>
<tr>
<td>Ala</td>
<td>24 (b)</td>
<td>14.48 (h)</td>
<td>24 (b)</td>
<td>48 (h)</td>
<td>72 (b)</td>
<td>24 (b)</td>
</tr>
<tr>
<td>Arg</td>
<td>24 (b)</td>
<td>90.9 (a)</td>
<td>24 (b)</td>
<td>105.9 (b)</td>
<td>119.5 (c)</td>
<td>24 (b)</td>
</tr>
<tr>
<td>Asp</td>
<td>24 (b)</td>
<td>19.4 (a)</td>
<td>24 (b)</td>
<td>23.6 (b)</td>
<td>25.9 (c)</td>
<td>24 (b)</td>
</tr>
<tr>
<td>Cys</td>
<td>24 (b)</td>
<td>10.9 (a)</td>
<td>24 (b)</td>
<td>13.1 (b)</td>
<td>13.7 (c)</td>
<td>24 (b)</td>
</tr>
<tr>
<td>Glu</td>
<td>24 (b)</td>
<td>54.5 (a)</td>
<td>24 (b)</td>
<td>65.8 (b)</td>
<td>72.9 (c)</td>
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</tr>
<tr>
<td>Gly</td>
<td>24 (b)</td>
<td>6.6 (a)</td>
<td>24 (b)</td>
<td>7.9 (b)</td>
<td>8.7 (c)</td>
<td>24 (b)</td>
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<tr>
<td>His</td>
<td>24 (b)</td>
<td>12.3 (a)</td>
<td>24 (b)</td>
<td>14.3 (b)</td>
<td>16.3 (c)</td>
<td>24 (b)</td>
</tr>
<tr>
<td>Leu</td>
<td>24 (b)</td>
<td>1.8 (a)</td>
<td>24 (b)</td>
<td>2.1 (b)</td>
<td>2.2 (c)</td>
<td>24 (b)</td>
</tr>
<tr>
<td>Lys</td>
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<td>4.3 (a)</td>
<td>24 (b)</td>
<td>5.1 (b)</td>
<td>6.7 (c)</td>
<td>24 (b)</td>
</tr>
<tr>
<td>Ile</td>
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<td>1.2 (a)</td>
<td>24 (b)</td>
<td>1.4 (b)</td>
<td>1.5 (c)</td>
<td>24 (b)</td>
</tr>
<tr>
<td>Met</td>
<td>4.4 (a)</td>
<td>0.4 (a)</td>
<td>4.4 (b)</td>
<td>0.5 (a)</td>
<td>0.6 (b)</td>
<td>4.4 (c)</td>
</tr>
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<td>Phe</td>
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<td>0.9 (a)</td>
<td>4.4 (b)</td>
<td>1.0 (a)</td>
<td>1.2 (b)</td>
<td>4.4 (c)</td>
</tr>
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<td>Pro</td>
<td>4.4 (a)</td>
<td>2.6 (a)</td>
<td>4.4 (b)</td>
<td>2.7 (a)</td>
<td>3.2 (b)</td>
<td>4.4 (c)</td>
</tr>
<tr>
<td>Ser</td>
<td>4.4 (a)</td>
<td>7.6 (a)</td>
<td>4.4 (b)</td>
<td>9.2 (a)</td>
<td>10.4 (b)</td>
<td>4.4 (c)</td>
</tr>
<tr>
<td>Thr</td>
<td>4.4 (a)</td>
<td>8.4 (a)</td>
<td>4.4 (b)</td>
<td>9.3 (a)</td>
<td>10.4 (b)</td>
<td>4.4 (c)</td>
</tr>
<tr>
<td>Tyr</td>
<td>4.4 (a)</td>
<td>1.1 (a)</td>
<td>4.4 (b)</td>
<td>1.3 (a)</td>
<td>1.5 (b)</td>
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<td>Val</td>
<td>4.4 (a)</td>
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<td>4.4 (b)</td>
<td>2.9 (a)</td>
<td>3.2 (b)</td>
<td>4.4 (c)</td>
</tr>
<tr>
<td>Total</td>
<td>214.9 (a)</td>
<td>280.3 (b)</td>
<td>217.1 (a)</td>
<td>275.2 (b)</td>
<td>231.9 (a)</td>
<td>303.2 (b)</td>
</tr>
</tbody>
</table>

Control (without starter culture), means with different letters (a, b, c) along rows are significantly different (P<0.05).
(Verplaetse et al., 1989). A major release of free amino acids at the beginning of the process have been studied in coincidence with the ripening stage (Diaz et al., 1997). This increase has been attributed to the higher temperatures applied during ripening compared to the low temperature. The most significant increases occurred in the content of Arg (arginine) in the sample. The decrease in the content of amino acids may indicate their metabolism by bacteria (Bover-Cid et al., 2000; Ordonez et al., 1999; Sekikawa et al., 2003).

### Table 3. Amino acid content of Nham without and with starter cultures during stored at 4°C.

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>Control</th>
<th>L. plantarum + L. sake</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Storage time (wk)</td>
<td>Storage time (wk)</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Ala</td>
<td>21.4a</td>
<td>22.3a</td>
</tr>
<tr>
<td>Arg</td>
<td>151.9a</td>
<td>154.4a</td>
</tr>
<tr>
<td>Asp</td>
<td>3.2a</td>
<td>3.4a</td>
</tr>
<tr>
<td>Cys</td>
<td>18.2a</td>
<td>18.5a</td>
</tr>
<tr>
<td>Glu</td>
<td>96.0a</td>
<td>98.4a</td>
</tr>
<tr>
<td>Gly</td>
<td>11.2ac</td>
<td>11.4ac</td>
</tr>
<tr>
<td>His</td>
<td>10.5a</td>
<td>10.5a</td>
</tr>
<tr>
<td>Leu</td>
<td>2.9a</td>
<td>3.0a</td>
</tr>
<tr>
<td>Lys</td>
<td>7.7a</td>
<td>7.8a</td>
</tr>
<tr>
<td>Ile</td>
<td>2.0a</td>
<td>2.0a</td>
</tr>
<tr>
<td>Met</td>
<td>0.6a</td>
<td>0.6a</td>
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<td>Phe</td>
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<td>1.5a</td>
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<tr>
<td>Pro</td>
<td>4.6a</td>
<td>4.6a</td>
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<td>Ser</td>
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<td>Tyr</td>
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<tr>
<td>Val</td>
<td>3.7a</td>
<td>3.8a</td>
</tr>
<tr>
<td>Total</td>
<td>352.9a</td>
<td>358.6a</td>
</tr>
</tbody>
</table>

Control (without starter culture).

Results are expressed as means of three replicates in mmol/g dry matter.

Means with different letters along rows are significantly different (P<0.05).
### Table 4. Amino acid content of Nham without and with starter cultures during stored at 15°C.

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>Control</th>
<th>L. plantarum + L. sake</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Storage time (wk)</td>
<td>Storage time (wk)</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Ala</td>
<td>35.6a</td>
<td>40.6b</td>
</tr>
<tr>
<td>Arg</td>
<td>232.9a</td>
<td>280.6b</td>
</tr>
<tr>
<td>Asp</td>
<td>4.7a</td>
<td>5.8ad</td>
</tr>
<tr>
<td>Cys</td>
<td>28.7a</td>
<td>34.6b</td>
</tr>
<tr>
<td>Glu</td>
<td>140.9a</td>
<td>174.2b</td>
</tr>
<tr>
<td>Gly</td>
<td>16.9a</td>
<td>20.3b</td>
</tr>
<tr>
<td>His</td>
<td>17.8a</td>
<td>21.0b</td>
</tr>
<tr>
<td>Leu</td>
<td>4.8a</td>
<td>5.9a</td>
</tr>
<tr>
<td>Lys</td>
<td>10.9a</td>
<td>13.5b</td>
</tr>
<tr>
<td>Ile</td>
<td>3.0a</td>
<td>3.6a</td>
</tr>
<tr>
<td>Met</td>
<td>0.9a</td>
<td>1.1ad</td>
</tr>
<tr>
<td>Phe</td>
<td>2.3ac</td>
<td>2.8a</td>
</tr>
<tr>
<td>Pro</td>
<td>6.5a</td>
<td>8.0b</td>
</tr>
<tr>
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<td>20.1a</td>
<td>24.2b</td>
</tr>
<tr>
<td>Thr</td>
<td>17.7a</td>
<td>21.9b</td>
</tr>
<tr>
<td>Tyr</td>
<td>2.8a</td>
<td>3.4a</td>
</tr>
<tr>
<td>Val</td>
<td>5.7a</td>
<td>7.0b</td>
</tr>
<tr>
<td>Total</td>
<td>348.8a</td>
<td>367.2a</td>
</tr>
</tbody>
</table>

Control (without starter culture).

Results are expressed as means of three replicates in mmol/g dry matter.

Means with different letters along rows are significantly different (P<0.05).

### Table 5. Amino acid content of Nham without and with starter cultures during stored at 25°C.

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>Control</th>
<th>L. plantarum + L. sake</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Storage time (wk)</td>
<td>Storage time (wk)</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Ala</td>
<td>35.6a</td>
<td>40.6b</td>
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<tr>
<td>Arg</td>
<td>232.9a</td>
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<tr>
<td>Asp</td>
<td>4.7a</td>
<td>5.8ad</td>
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<tr>
<td>Cys</td>
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<tr>
<td>Glu</td>
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<tr>
<td>Gly</td>
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<tr>
<td>His</td>
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<td>21.0b</td>
</tr>
<tr>
<td>Leu</td>
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<td>5.9a</td>
</tr>
<tr>
<td>Lys</td>
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<td>13.5b</td>
</tr>
<tr>
<td>Ile</td>
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</tr>
<tr>
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<td>367.2a</td>
</tr>
</tbody>
</table>

Control (without starter culture).

Results are expressed as means of three replicates in mmol/g dry matter.

Means with different letters along rows are significantly different (P<0.05).
Two types of fermented sausage differing in starter culture were produced in parallel with two different starter cultures (no starter and *L. plantarum* + *L. sake*). The sausages were ripened 3 days and subsequently stored 7, 14, 21 and 28 days at the 4 °C, 15 °C and 25 °C. Concentration of three most abundant amines, cadaverine, putrescine and tyramine increased significantly (*P* < 0.05) in Nham during ripening and also during storage. The dominant BAs in the control were cadaverine – and tyramine and putrescine, to a lesser extent; the cadaverine, putrescine and tyramine content were lower if inoculation had added with *L. plantarum* + *L. sake*; whereas they ranked above 300 mg/kg in the control by 3 d. At the end of ripening, cadaverine (98.7 mg/kg dry matter), putrescine (242.6 mg/kg dry matter) and tyramine (46.4 mg/kg dry matter) content in the A-samples-sausage was higher (*P* < 0.05) than in Nham with starters (51.6, 98.4 and 27.8 mg/kg dry matter, respectively). Starter culture influenced significantly in decrease of (*P* < 0.05) cadaverine, putrescine and tyramine content in the sausage. Due to the significant (*P* < 0.05) increase of total aerobic counts in the Nham control between the end of ripening and during storage, followed by the significant (*P* < 0.05) increase of the sum of total BAs between the 72 hour of ripening (387.7 mg/kg dry matter) and the 4th week of storage at 25°C (629.2 mg/kg dry matter).

The main rate of BAs production was during the first two days, when a sharp pH decrease and the development of LAB occurred. Sausages fermented with starters had lower amounts of cadaverine, putrescine and tyramine than naturally fermented sausages (control) during storage at 15°C and 25°C. However, phenylethylamine, histamine and tryptamine were not detected.

Nham control showed proteolysis that was correlated with pH values higher than those with starters. However, no positive correlation was found between the proteolysis index and BAs production. Since proteolysis was stronger during the second half of the ripening process, the FAAs occurred later than the early amine production. No effect on pH development in the fermented sausage was observed when non-amine forming strain of *L. plantarum* + *L. sake* were present during 4 week of 4°C storage period. A study on the evolution of FAAs and BAs in Nham during 4 week at different temperatures of storage (4°C, 15°C and 25°C) was performed. FAAs and BAs were determined by RP-HPLC. Storage temperature of 15°C and 25°C promoted a significant increase of the contents of arginine, glutamic acid, cadaverine, putrescine and tyramine, expressed as g/kg of dry matter while storage temperature of 4°C decreased a significant of the contents of arginine, glutamic acid, cadaverine, putrescine and tyramine, expressed as g/kg of dry matter. These two amino acids and three BAs may serve as indicators of temperatures changes in stored fermented sausage.

4. Conclusions

The aim of this study was to investigate the effect of non-amine forming LAB as starter culture during ripening and storage time and temperature on the evolution of FAAs of Nham during processing. The correlation between FAAs and BAs content was also investigated. Larger increases of FAAs occurred in Nham without starter in the ripening and storage step. Total FAAs content was highly correlated with total BAs amount. Sausage
ripening was further carried out with non-amine forming strain of \textit{L. plantarum} + \textit{L. sake} after ripening and stored at different temperature. The amount of amine in the product was significantly less than the control. The results obtained for BAs degradation by bacteria in a synthetic medium suggest that AO activity is strain dependent rather than being related to specific species. In all batches, the total amino acid contents increased with time – and the predominant ones were arginine and glutamic acid. However, upon inoculation with non-amine forming strain, the total BAs contents remained considerably lower than those of the control. Hence, an efficient food-grade biological tool was made available that constrains buildup of BAs in fermented sausage during storage.

**Author details**

Jirasak Kongkiattikajorn  
School of Bioresources and Technology, King Mongkut’s University of Technology Thonburi, Thailand

**Acknowledgement**

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**5. References**


