Simple Detection Method of Biogenic Amines in Decomposed Fish by Intramolecular Excimer Fluorescence

Hirofumi Nishikawa, Tatsuya Tabata, Seiichi Kitani

Health Service Center, Tokyo University of Marine Science and Technology, Konan, Japan.
Email: *drkitani@kaiyodai.ac.jp

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ABSTRACT

Biogenic amines are known to have various biological functions such as not only neurotransmitter and cell proliferation but also food poisoning. Bacterially-decomposed amines such as histamine, agmatine, putrescine, cadaverine, spermidine and spermine cause allergic symptoms. We developed simple method for measurement of polyamine as indicator of food decomposition with a fluorometer by using 4-(1-Pyrene)butyric acid N-hydroxysuccinimide ester (PSE). PSE reacts with primary and secondary amino moieties of polyamines and produces the intramolecular excimer fluorescence. Excimer fluorescence with broad peak at around 470 nm was clearly detected in linear type biogenic amines such as putrescine, cadaverine, spermidine and spermine at 10 mM. However neither histamine nor trimethylamine altered the fluorescence. Decomposed sardine and mackerel by improper storage showed stronger intensity than fresh ones. Comparing with OPA-method, PSE method was useful for screening biogenic amines present in food, esp. fish since the analysis was simple after one-step purification procedure. An inexpensive system which can rapidly detect biogenic amines from food is necessary in medium and small-sized food business. The technique using excimer has potential to realize the high through-put screening system for evaluation of food freshness and is expected to bring the public interests such as food security and safety of consumer.

Keywords: 4-(1-Pyrene)butyric Acid N-Hydroxysuccinimide Ester; PSE Excimer Fluorescence; Biogenic Amine; Fish Decomposition; Polyamine; Food Poisoning

1. Introduction

Biogenic amines are a group of chemicals synthesized from amino acids. The amines are naturally present in the body, having various biological functions. For instance, monoamines such as catecholamine (adrenaline and dopamine) and tryptamine (serotonin and melatonin) play a role as neurotransmitters [1]. Polyamine, an cyclic or linear organic compound having two or more primary amino groups as shown in Figure 1, are involved with growth and cell proliferation [2-4]. The heterocyclic amine such as histamine and tertiary amine such as trimethylamine also have biological functions.

Biogenic amines such as histamine, putrescine, cadaverine, agmatine, spermidine and spermine are also found in seafood, meat and cheese, which are generated by bacterial enzymatic decarboxylase of free amino acid [2,5,6]. For instance, histamine is synthesized from histidine by histidine decarboxylase (HDC) secreted from microbes belonging to gram-negative bacteria (i.e., *Mor-ganella morganii*, *Enterobacter aerogenes*, *Photobacterium phosphoreum*, *Raoultella planticola*, etc.) and gram-positive bacteria such as lactic acid bacteria [7]. Similarly, other amino acid specific decarboxylases by various spoil- ing bacterium synthesize putrescine from ornithine, cadaverine from lysine, and spermidine and spermine from arginine, respectively. The composition of these biogenic amines depends on quantity of each amino acid in foods.

![Figure 1. Structure of biogenic amines used in this study.](image-url)
In fact the fish such as families Scombridae (e.g. tuna and mackerel) and some of non-scombroid fish (e.g. mahi-mahi, sardines, pilchards, anchovies, herring, marlin and bluefish) are prone to accumulate histamine because of containing high levels of free histidine in these fish, [8]. On the other hand, seafood such as cephalopods (e.g. squid, octopus and cuttlefish), crustaceans (e.g. shrimp and crab) and bivalve mollusks mainly produce putrescine and cadaverine but not histamine during decomposition [9-11].

The ingestion of exogenous huge amounts of amines in decomposed food results in food poisoning. Especially, histamine at the concentrations higher than 500 ppm causes food poisoning [12]. The symptom typically occurs within from 10 min to 1 h in case of consumption of poisonous fish, and resembles Type-I allergy such as hives, hot rash, flushing, nausea and facial swelling [13]. Recovery is usually completed within 24 h, but in rare cases can last for days [14]. Rarely are serious cardiac and respiratory complications observed for patients with preexisting disease conditions [15]. Histamine is metabolized by various enzymes such as monoamine-oxidase (MAO) and diamine oxidase (DAO) [16]. Therefore people who are deficient in their enzymatic function owing to genetic causes or through inhibition by taking anti-depression medicines such as monoamine oxidase inhibitors (MAOIs) are more susceptible to histamine toxicity [2,17]. The other aliphatic amines such as putrescine and cadaverine are known to enhance histamine toxicity [18]. The mechanism of amplification is thought as a result of inhibition of DAO [19] and increase of histamine absorption [20]. Release of endogenous histamine from mast cells by scombroid toxin is also suggested to associate with allergy-like symptoms [21]. There are supportive reports that spermine is known to enhance IgE-mediated degranulation [22] and induce the release through G-protein activation in mast cells [23]. Polyamines are present in mast cell secretory granules and associate with granule homeostasis [24], suggesting the synergistic or additive enhancement of allergic symptoms.

From a standpoint of food toxicity and hygiene, there have been developed the various analytic methods of biogenic amines such as thin-layer chromatography, gas chromatography, HPLC as well as capillary electrophoretic methods. Because most amines show neither UV absorption nor fluorescence, most methods require the derivatization, for instance dansyl chloride [25], dansyl chloride [26], O-phthalaldehyde (OPA) [27], 6-aminoquinolyl-N-hydroxysuccinimidyl carbamate [28]. OPA method mentioned above, generally established as simple and high sensitive analysis of histamine, requires troublesome pretreatment to remove impurity. On the other hand, formation of intramolecular excimer by 4-(1-pyrene)butyric acid N-hydroxysuccinimide ester (PSE) is reported to selectively detect biogenic amines with simple pretreatment [29]. PSE reacts with primary and secondary amino moieties of polyamines in weakly alkaline environment, which produces intramolecular excimer-forming fluorescence with wavelength region (450 - 520 nm) longer than the emission from PSE monomer (360 - 420 nm) [30]. 1-Pyrenebutyryl Chloride (PBC) is known as other derivative for analysis polyamine using excimer fluorescence [31]. Marks et al. reported the analysis of putrescine and cadaverine in seafood by HPLC using excimer fluorescence method, but did not monitor the increase of the amines with decomposition [32]. In their reports, spermine and spermidines, mast cell secretagogues, were not analyzed, which is not adequately presented in connection with allergy-like food poisoning.

Analysis method by using HPLC is superior in high resolution and sensitivity, but is not suitable for screening of a mass of samples. Furthermore training of technician and start-up of the system and running cost are so expensive for a medium and small-sized fishery business. Therefore we developed simple method to monitor freshness during storage of fish by using spectrofluorometer, which detects excimer fluorescence of PSE labeled polyamines such as putrescine, cadaverine, spermine and spermidine. We also confirmed the association among fish decomposition and increase of excimer fluorescence.

2. Material and Methods
2.1. Reagents
Histamine, putrescine, spermine, spermidine, agmatine, cadaverine, trimethylamine, O-phthalaldehyde, 4-(1-pyrene)butyric acid N-hydroxysuccinimide ester (PSE) were purchased from Sigma (Sigma Aldrich, St. Louis, MO). 1-Pyrenebutyryl Chloride (PBC) was purchased from Toronto Research Chemicals Inc., (Ontario, Canada). The purity of all reagents was HPLC grade or highest quality.

2.2. Preparation of Solutions
Histamine, putrescine, spermine, spermidine, agmatine, cadaverine and trimethylamine were prepared in 1 M, and diluted to the required concentration by HPLC grade water (Wako pure chemical, Osaka, Japan) before use. OPA solution was prepared by dissolving 20 mg OPA into 10 mL of ethanol. PSE solution was prepared in 5 mM by acetonitrile of highest grade. PBC solution was prepared in 30 mM in DMSO and diluted to 6 mM by acetonitrile. All the solutions were stored in polypropylene micro tubes protected from light and stored at –20°C, and used within 1 week.

2.3. Preparation of Extracts
Fresh sardine Sardinops melanostictus (Temminck and Schlegel, 1846) and mackerel Scomber japonicus (Hout-
tuyyn, 1782) were purchased from a supermarket in Tokyo. The fish in polystyrene foam box containing ice were immediately transported to the laboratory. Some fish were stored at 30 °C for 24 h and 72 h to be decomposed by microbial decarboxylation. After peeling, 5.0 g of fish meat was homogenated with 8.0 mL of water and centrifuged at 430 × g for 10 min at 4°C. The supernatant volume was re-adjusted to 8.0 mL and mixed with 2.4 mL of 30% trichloroacetic acid, followed by centrifugation at 13,000 × g for 10 min to remove crude protein. The supernatant was filtered with 0.45 μm syringe filter DIS-MIC (Advantec, Japan), stored at –80 °C until analysis. As control, fresh fish extracts were prepared just after thawing. For OPA-derivatization, the extract was further purified by mixing 3 N NaOH containing 100 mg/mL of NaCl with N-butanol at ratio of 20:25:2 (vol/vol) followed by centrifugation at 10,000 × g for 10 min. The supernatant was added with 0.12 N HCl and N-heptane at 5:3:9 (vol/vol). After shaking, the lower phase was collected and stored at –80°C until analysis.

2.4. Derivatization Method

OPA derivatization was performed as follow. 1200 μL of purified fish extract was mixed with 60 μL of OPA solution and 240 μL of 1 N NaCl. After incubation at 4°C for 40 min, 120 μL of 3 N HCl was added. PSE derivatization was performed as described with slight modification [33]. Briefly, 150 μL of each amine standard solution or fish solution were mixed with 1200 μL of 5 mM PSE solution and 300 μL of 1.5 mM potassium carbonate in 2-mL screw cap polypropylene tubes (Axygen scientific, Unicon city, California). The tubes were tightly sealed and heated for 90 min in boiling water. The reacted solutions were diluted 1000 times with acetonitrile to decrease emission from monomer. PBC-labeling was performed as described, with slight modification [31]. Briefly, sample solutions, 6 mM PBC solution and 0.5 mM potassium carbonate were mixed at ratio (volume) of 10:20:1 (600 μL:1200 μL:60 μL), followed by incubation at 25°C for 5 min. The fluorescence intensity from OPA-, PSE, PBC-derivatized biogenic amines were measured by emission 350 - 600 nm and excitation of 340, 350 and 355 nm, respectively.

2.5. Equipment and Analysis

Fluorescence spectral was measured with RF-5300 spectrofluorometer (Shimadzu, Tokyo, Japan) in 10 × 10 mM quartz cuvette. Spectral band width of 1.5 nm was used for both the excitation and emission. Data were collected and analyzed with Ion Prove Program (version 1.00) (Shimadzu, Tokyo, Japan).

3. Results and Discussion

3.1. Fluorescence from OPA Derivatized Histamine and Decayed Fish

We investigated the fluorescence from OPA labeled histamine at the concentration between 10 pM - 1 M, and found peak fluorescence at 445 nm with dose-dependent increase in a range of 100 nm - 100 μm (Figure 2(a)).

Fluorescence intensity from distilled water, 100 nm and 1 M of histamine were 1.03, 1.56 and 4.31, respectively. The solution of histamine above 1 mM was yellow color, and therefore it is impossible to take proper intensity. We also measured OPA-derivatized fish extracts prepared from fresh sardine, mackerel and their spoiled fishes incubated at 30°C for 3 days. Both spoiled sardine and mackerel obviously increased fluorescence compared with fresh one, suggesting the histamine production through decomposition (Figure 2(b)).

![Figure 2. Fluorescence from OPA-derivatized histamine. (a) Fluorescence intensity in histamine solution. Number 1 - 3 indicate 100, 10 and 1 μM of histamine stock solutions, respectively; (b) Fluorescence intensity in fish extract. Number 1 - 3 indicate (1) mackerel (72 h incubation), (2) sardine (72 h incubation) and (3) both fresh sample, respectively.](image-url)
3.2. Fluorescence from Biogenic Amines Labeled with PSE

We next analyzed the excimer fluorescence from biogenic amines mainly produced in decayed fish. Each concentration of amine standard solutions (0, 0.1, 1, 10 mM) was reacted with weakly alkaline-PSE solution, and diluted 1000-times with acetonitrile to minimize intermolecular excimer. Neither histamine nor trimethylamine altered the fluorescence (Figures 3(a) and (b)). Agmatine minimally increased fluorescence around 450 nm (Figure 3(c)). Excimer fluorescence with broad peak at around 470 nm was clearly appeared in putrescine, cadaverine, spermidine and spermine at the concentration of 10 mM (Figures 3(d)-(g)). As for PBC-derivatized amines, there is no change of peak and intensity (data not shown). Although the detection of PSE-derivatized histamine from rat brain by using HPLC was reported [33], we could detect the excimer fluorescence from only linear type diamines, namely putrescine, cadaverine, spermidine and spermine (structures shown in Figure 1, right side), but not histamine in our analysis. Trimethylamine, itself tertiary amine, did not emit the fluorescence. In addition, increase of the excimer fluorescence seen in PSE-labeled agmatine seems false-positive, since stokes shift was smaller than that of other polyamine. In case of amine having primary amine at the branched side chain,

![Figure 3. Fluorescence from PSE-derivatized biogenic amines.](image-url)
normally excited dimer may not be composed because of an inadequate tertiary structure. In fact, it is known that two or more fluorophores in a molecule cause quenching of fluorescence, called self-quenching. Therefore, it is supposed that only linear diamines containing primary amines at both ends are detectable with PSE in our method.

3.3. Evaluation of Fish-Freshness by PSE Excimer-Fluorescence

Freshness of fish was evaluated by measurement of excimer fluorescence from PSE-derivatized fish extract after one-step purification with trichloroacetic acid. The fluorescence wave profile in the extracts prepared from fresh fish was the same as water (Figure 4(a), line 4-6). Spoiled mackerel released the significant emission with broad peak at around 470 nm (Figure 4(a), line 2). The fluorescence intensity was also increased in the extract from sardine incubated for 72 h, and was higher than that from that for 24 h (Figure 4(a), line 1 and 3).

Figure 4(b) shows the fluorescence intensity value of each samples subtracted that of blank (HPLC grade water). The fluorescence intensity in fresh sardine and mackerel were 0.65 and 1.43. Similarly, histamine, agmatine and trimethylamine showed the intensity of 0.77, 0.14 and 0.30, respectively. Mackerel incubated for 72 h at 30°C increased the intensity to 8.39. In sardine, the intensity in 24 h and 72 h at 30°C incubation were 3.60 and 21.4, showing polyamine production during improper storage. Therefore, polyamine in fish extract can be estimated as 1 - 10 mM level of putrescine, cadaverine, spermidine and spermine, suggesting significant detection level and practical application for evaluation of freshness (Figure 4(b)).

4. Conclusion

Biogenic amines are involved with food poisoning. In this study, we demonstrated that PSE-derivatized excimer fluorescence was detected in linear polyamines such as putrescine, cadaverine, spermidine and spermine. We also demonstrated here the increase of fluorescence in decomposed sardine and mackerel fish by PSE derivatization just by removing proteins or debris with trichloroacetic acid, which is simpler than OPA method. PSE did not give intramolecular excimer fluorescence from heterocyclic amine histamine. This inexpensive and simple technique will be useful for a medium and small-sized food business, and also brings the public interests in context of food security and safety of consumer.

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