Bioluminescence: Chemical Study on Visible Light Emission from Fungal Mycelium and Fruiting Body

Katsunori Teranishi*
Graduate School of Bioresources, Mie University, Japan

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*Corresponding author: Katsunori Teranishi, Graduate School of Bioresources, Mie University, 1577 Kurimamachiya, Tsu, Mie, Japan, Tel: +81-592-31-9615. Email: teranisi@bio.mie-u.ac.jp

Abstract

Although the visible light emission by living organisms is generally thought to be rare, in fact the bioluminescent phenomenon can be widely observed in nature, for examples insects, fishes, bacteria, and fungi. The phenomenon has intrigued scientific researchers over the years. The chemical mechanisms underlying bioluminescence differ between species. Some mechanisms, such as bioluminescence of firefly, jellyfish, bacteria, and dinoflagellate, have been elucidated and further more their principles have allowed the development of many novel technologies in agriculture, biology, ecology, and medicine. However, many other bioluminescence mechanisms yet remain to be understood. Further research of bioluminescence will lead discoveries of biological significance and novel possibility in science. Here I present studies on chemical mechanism underlying fungal bioluminescence, which are actively in progress.

Keywords: Bioluminescence; Chemiluminescence; Fungus; Luciferin; Luciferase; Mechanism

Mini Review

Bioluminescence is the emission of visible light by living organisms, such as bacteria, fireflies, fishes, ostracods of the genus *Cypridina*, and jellyfish of the genus *Aequorea* [1]. The bioluminescence phenomenon has been arousing fascination in human and curiosity in scientists from a long time ago (Figure 1). Detailed chemical principles in bioluminescence have been disclosed by biological, organic, and physical chemists since 1950’s, and then it has been understood that the bioluminescence phenomenon is caused by energy conversion of chemical energy to light energy via chemical reactions in bioluminescence systems. Moreover, the bioluminescence principles have been applied in many scientific fields such as agriculture, biology, ecology, and medicine [2]. Since instrumentation for measuring light is very sensitive and low background, chemiluminescence methods based on the bioluminescence principles has come into widespread use for quantitative determination of specific substances in many scientific fields. Bioluminescent organisms use their light, which possesses various colors, periodic patterns, and intensities, for self-defense against predators, camouflage, intra specific communication, or attracting mates or prey [3].

Fungal bioluminescence is also widely observed on decaying wood or leaves in night. Thus, the phenomenon is called “foxfire” or “shining wood”. The total number of documented species of luminous fungi by 2016 is more than 80 [4]. Fungal bioluminescence, which varies among species, occurs in the basidiocarp, mycelium, spores, or in...
their combinations [5-7]. Although fungal bioluminescence generally shows lower intensities than other bioluminescent organisms such as fireflies and ostracods, the light emission from fungi is continuously generated for day and night. Thus, total light emission from certain bioluminescent fungi may be comparable with that of firefly. The color of fungal bioluminescence is commonly green. With regard to functions of fungal bioluminescence, Olivera et al. reported that fruiting bodies of bioluminescent fungus Neonothopanus gardneri can attract potential spore dispersing insects by glowing [8]. On the other hand, Weinstein et al. reported that Australian fruiting bodies of bioluminescent fungus Omphalotus nidiformis did not attract the insects [9]. The mycelium is underground or inside a decaying wood. It is possible that fungal bioluminescence is a by-product of metabolism, and possesses no function. Functions of fungal bioluminescence yet remain to be understood.

In spite of the wide popularity of fungal bioluminescence, chemical mechanism underlying fungal bioluminescence has been subjected to far less scientific investigation than the bioluminescence from other sources. In 1668, Boyle reported that air was needed for the light emission of shining wood [10]. Although many researchers studied on chemical mechanism underlying fungal bioluminescence, their efforts were unsuccessful until Airth's group showed positive results using bioluminescent fungi Collybia velutipes and Armillariamellea in 1959 [11]. Airth et al. reported that the bioluminescence was NAD(P)H-dependent luminescence reaction, in which unknown substrate (luciferin), molecular oxygen, water-soluble enzyme, and water-insoluble enzyme (luciferase) were involved [11-13]. However, after that, the luciferase and luciferin were not isolated and their structures and luminescent properties were not determined. Kuwabara and Wassink isolated a luciferin, which displayed chemiluminescence in the presence of $\text{H}_2\text{O}_2$, in a crystal form from the mycelia of Omphalia flavida [14]; however, its chemical structure was not reported.

Shimomura demonstrated a negative result with regard to the Airth's luciferin-luciferase reaction to the fruiting body bioluminescence of the luminous fungus Panellus stipticus; the precursors, panal, PS-A, and PS-B (Figure 2), of light-emitting compounds were isolated from the luminous fruiting bodies that upon activation with ammonium or primary amines, emitted light in the presence of $\text{H}_2\text{O}_2$ and $\text{Fe}^{2+}$ or in the presence of superoxide anion and molecular oxygen [15,16]. Moreover, Shimomura purified the chemiluminescent activation products that were produced by reaction of the precursors with methylamine, and inferred the possible structures of the activation products using the model compounds of the precursors [17]. These activation products or compounds corresponding to the activation products in the P. stipticus fruiting bodies have not yet been found. Subsequently, Shimomura analyzed the interrelationships between the bioluminescence and the superoxide dismutase activities using six species of fungi, including P. stipticus (fruiting body and mycelium), and demonstrated that superoxide dismutase regulated the bioluminescence activity [18]. On the basis of these findings, Shimomura suggested that, unlike Airth's luciferin-luciferase reaction, P. stipticus bioluminescence was not a luciferin-luciferase type. Shimomura also suggested that superoxide anion triggered the bioluminescence reaction in the presence of molecular oxygen for generating light.

In 2012, Oliveira et al. reported the hypothesis that
all known bioluminescent fungal lineages share luciferin-luciferase reaction shown by Airth’s group [19]. In 2015, Yampolsky’s group reported that and trans-3-hydroxyhispidin, which played as luciferin with water-insoluble luciferase, was produced from trans-hispidin with molecular oxygen and NADPH in the presence of water-soluble hydroxylase for mycelia bioluminescence of Neonothopanus nambi (Figure 3) [20]. Then, Yampolsky’s group reported that and trans-3-hydroxyhispidin was oxidized to 2Z,5E-6-(3,4-dihydroxyphenyl)-2-hydroxy-4-oxoexa-2,5-dienoic acid(oxyluciferin) as a light emitter in the presence of a luciferase-enriched protein fraction prepared from N. gardneri and N. nambi mycelium (Figure 3) [21]. Trans-3-hydroxyhispidin is a strong “candidate” of luciferin in N. gardneri and N. nambi mycelia at the present time. Luciferase has been successfully extracted from N. gardneri and N. nambi mycelia and partially purified [21]. Therefore, it is distinct that these mycelia possess luminescence systems involving trans-3-hydroxyhispidin. In the future it is necessary to elucidate whether these luminescence systems generate true bioluminescence in the living N. gardneri and N. nambi mycelia or not, and whether similar bioluminescence systems involving trans-3-hydroxyhispidin exist in other bioluminescent fungi or not.

Although various fungal species have been used for chemical research of fungal bioluminescence over the years, mycelia were subject for the research, except for Shimomura’s experiments above mentioned. Some researchers believe that bioluminescence mechanism for all bioluminescent fungi should be the same, and that chemical mechanism behind bioluminescence in mycelia and fruiting bodies is the same. Because bioluminescent fungus Mycena chlorophos (Figure 1), which was found in Southeast Asia [22] and can be cultivated in laboratory [23,24], produces continuously bright light in the mycelium and pileus gills, in which light is produced in cell membrane [25], this species is a suitable for evaluating those hypotheses. Teranishi has been chemically investigating bioluminescence mechanism of this fungus using living tissue. Generally when fungal tissue is crushed, bioluminescence is immediately stopped. Because the crushed tissue loses the bioluminescence ability, the crushed tissue and extracts from the tissue cannot be used for luminescence assays. Recently Teranishi demonstrated that trans-4-hydroxycinnamic acid and trans-3,4-dihydroxycinnamic acid in gills increased the bioluminescence intensity in living immature gills of M. chlorophos, which emitted weak light [26,27], and that other cinnamic acids shown in Figure 4 did not influence the original light emission. Moreover, the light emission before and after addition of trans-4-hydroxycinnamic acid or trans-3,4-dihydroxycinnamic acid to gills was inhibited by addition of trans-4-aminoacinnamic acid, whereas trans-2- and trans-3-aminoacinnamic acids did not inhibit the light emission [28]. These results indicate that trans-4- and/or 3,4-di-hydroxycinnamic acids contribute to bioluminescence in M. Chlorophos living gills, and that the hydroxy group at position C-4 intrans-cinnamic acid plays an important role for producing the energy for light emission. Although Teranishi reported that the combination of NADPH and hispid in was not essential for the production of bright light in the living gills [29], bioluminescence ability of trans-3-hydroxyhispidin yet remains to be elucidated. Investigation for understanding the chemical mechanism underlying bioluminescence observed in M. chlorophos gills and mycelia is in progress.

![Figure 4: Compounds used in the assay for gill bioluminescence of M. chlorophos (26,27).](image)

**References**


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