Genetically encodable bioluminescent system from fungi

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Abstract

Bioluminescence is a natural phenomenon of light emission resulting from oxidation of a substrate, luciferin, catalyzed by an enzyme, luciferase. A variety of species are bioluminescent in nature (1); for many of them, the ability to emit light is a defining feature of their biology (2–4). Artificial integration of natural bioluminescent reactions into living systems has also become a reporting tool widely used in molecular and cell biology (5, 6). However, natural bioluminescent systems remain poorly characterized at a biochemical level, limiting more widespread application. Only 9 luciferins and 7 luciferase gene families have been described (7, 8) of at least 40 bioluminescent organisms. Here, we describe the function and evolution of fungal luciferase genes and propose a pathway through several closely related stepping stone nonluminescent biochemical reactions with adaptive roles. The availability of a complete eukaryotic luciferin biosynthesis pathway provides several applications in biomedicine and bioengineering.

Biomolecules

Bioluminescence
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This open access article is distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivatives License 4.0 (CC BY-NC-ND). Data deposition: The data reported in this paper have been deposited in the NCBI BioProject (accession no. PRJNA476325). Transcriptomes of N. nambi and M. circitorum, alignments of P. pastoris genome sequencing reads, and alignment of protein sequences of fungal luciferases are available at Figshare (https://figshare.com/articles/A_genetically_encodable_bioluminescent_system_from_fungi/6738953/2). Files used to reconstruct the Agaricales species tree, including raw and trimmed alignments and RAxML resulting files, are available at Figshare (https://figshare.com/articles/Species_trees_files/6072117).

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We present identification of the luciferase and enzymes of the biosynthesis of a eukaryotic luciferin from fungi. Fungi possess a simple bioluminescent system, with luciferin being only two enzymatic steps from well-known metabolic pathways. The expression of genes from the fungal bioluminescent pathway is not toxic to eukaryotic cells, and the luciferase can be easily co-opted to bioimaging applications. With the fungal system being a genetically encodable bioluminescent system from eukaryotes, it is now possible to create artificially bioluminescent eukaryotes by expression of three genes. The fungal bioluminescent system represents an example of molecular evolution of a complex ecological trait and with molecular details reported in the paper, will allow additional research into ecological significance of fungal bioluminescence.

attract spore-distributing insects (13). Fungal bioluminescence was known to utilize at least four components: molecular oxygen; the luciferin, which was recently identified as 3-hydroxyhispidin [a product of oxidation of the simple plant and fungal metabolite hispidin (14)]; and two previously undescribed key enzymes, an NAD(P)H-dependent hydroxylase and a luciferase (15, 16).

To identify enzymes of the fungal bioluminescent pathway, we first focused on isolation of the luciferase gene. By expressing the \textit{N. nambi} cDNA library in \textit{Pichia pastoris} and spraying agar plates with synthetic 3-hydroxyhispidin, we identified and sequenced a luminescent yeast colony expressing the luciferase gene (\textit{SI Appendix}, Figs. S1 and S13). The \textit{N. nambi} luciferase, nnLuz, is a 267-aa protein (\textit{SI Appendix}, Fig. S2) and has no described homologs or pronounced sequence similarity to conserved protein domains, representing a novel protein family.

Genes coding for enzymes that synthesize secondary metabolites are often clustered in fungal genomes (17). We hypothesized that this may be the case for enzymes of the bioluminescent cascade, because it is thought that the cascade is conserved among the bioluminescent fungi (12). We thus looked for genes related to luciferin biosynthesis in the vicinity of the luciferase gene in the \textit{N. nambi} genome. In addition to \textit{N. nambi}, we also sequenced genomes and transcriptomes of bioluminescent fungi \textit{Neonothopanus gardneri}, \textit{Mycena citricolor}, and \textit{Panellus stipticus} and compared them with publicly available genome sequences of bioluminescent and nonbioluminescent fungi (18, 19). We found that the luciferase is a member of a conserved gene cluster, which includes at least two other genes: \textit{h3h} and \textit{hisps} (Fig. 1 B and C and Datasets S1–S3).

The \textit{h3h} gene showed sequence similarity with 3-hydroxybenzoate 6-monoxygenases, enzymes that catalyze oxidation of 3-hydroxybenzoate using NADH and molecular oxygen. This reaction is identical to that which converts hispidin into luciferin (Fig. 1A); thus, we hypothesized that \textit{h3h} gene codes for hispidin-3-hydroxylase (H3H), the enzyme corresponding to the predicted hydroxylase (15). We cloned the gene from \textit{N. nambi} and found that \textit{P. pastoris} colonies expressing both \textit{nnluz} and \textit{nnh3h} emit light when sprayed with luciferin precursor hispidin, unlike control colonies expressing \textit{nnluz} alone (\textit{SI Appendix}, Figs. S14 and S17)—confirming that \textit{nnH3H} converts hispidin into luciferin.

The \textit{hisps} gene (Fig. 1C) encodes a member of the polyketide synthase family, enzymes that produce secondary metabolites in a variety of organisms across the tree of life (20). Polyketide synthases typically add malonyl moieties to the growing carbon chain; thus, the α-pyrene nature of hispidin suggested that its biosynthesis may be performed by a polyketide synthase from caffeic acid by two cycles of addition of malonyl units followed by lactonization (Fig. 1A and \textit{SI Appendix}, Fig. S3). Large modular polyketide synthases require posttranslational modifications for their activity (21), such as the transfer of a phosphopantetheiny1 group to a conserved serine residue of the acyl carrier protein.

**Fig. 1.** Luciferin biosynthesis pathway in fungal bioluminescence and gene cluster containing key enzymes in the clade of bioluminescent fungi. (A) Proposed pathway of fungal luciferin biosynthesis and recycling. Caffeic acid is converted to hispidin by hispidin synthase (HisPS) and hydroxylated by H3H, yielding 3-hydroxyhispidin (fungal luciferin). The luciferase (Luz) adds molecular oxygen, producing an endoperoxide as a high-energy intermediate with decomposition that yields oxyluciferin (caffeoylpyruvate) and light emission. Oxyluciferin can be recycled to caffeic acid by caffeoylpyruvate hydrolase (CPH). (B) Schematic depiction of the genomic cluster of \textit{N. nambi} containing luciferase, \textit{H3H}, hispidin synthase, and caffeoylpyruvate hydrolase (\textit{cpH}) genes. (C) Gene cluster in the clade of bioluminescent fungi. The species tree in \textit{Left} is based on the comparison of protein-coding genes shared by most of the analyzed species. The red crosses mark the branches of the tree that eventually lost the ability to glow. \textit{Right} shows the structure of the luciferase-containing gene cluster if such a cluster was found in the relevant genome. The genes coding for luciferase (Luz), \textit{h3h}, hispidin synthase (\textit{hisps}), and caffeoylpyruvate hydrolase (\textit{cpH}) are colored. The lighter blue and red colors of \textit{hisps} and \textit{luz} genes indicate that only a partial or truncated gene was found in \textit{Armillaria mellea} and \textit{Guyanagaster necrorhiza}, respectively. Other genes that might belong to the cluster are named from O1 to O4 (colored in gray). Green ticks represent a cytochrome P450-like gene (different shades of green indicate different orthologous groups), and black ticks indicate other genes.
domain. To test whether hisps gene can produce luciferin precursor in a heterologous system, we integrated hisps, nnluz, and nnh3 genes together with the Aspergillus nidulans 4'-phosphopantetheinyl transferase gene npgA into the genome of P. pastoris. When cultured in a medium containing caffeic acid, yeasts expressing all four genes emitted light seen by a naked eye (Fig. 3A), while no significant light production was observed in strains lacking npgA or hisps genes (SI Appendix, Figs. S15 and S17). Therefore, hisps catalyzes synthesis of hispidin from caffeic acid, closing the chain of reactions (the “caffeic acid cycle”) from a common cellular metabolite with known biosynthesis to a eukaryotic luciferin.

In some bioluminescent species of fungi, the genomic cluster accommodates one or two additional genes (Fig. 1C): one belonging to the cytochrome P450 family, and the other belonging to the family of fumarylacetoacetate hydrolases. The latter (cph) likely encodes a caffeoylpyruvate hydrolase (Dataset S4) involved in oxyluciferin recycling, consistent with caffeoylpyruvate, the fungal oxyluciferin, being hydrolyzed to caffeate and pyruvate by a hydrolase present in fungal crude extracts (22).

Conservation of the gene cluster suggests that, in contrast to other groups of bioluminescent organisms (23), bioluminescence evolved in fungi only once, with luc, h3h, and hisps genes emerging through gene duplications. Reconstructed phylogenetic trees for luc, h3h, and hisps genes (SI Appendix, Figs. S4–S6) and the species tree of order Agaricales (Fig. 2) reveal the evolution of the bioluminescence cascade in fungi. The primary luc enzyme of the fungal bioluminescence cascade emerged through a gene duplication at the base of Agaricales followed by the duplication of h3h and hisps a few million years later. Interestingly, many species in a large clade encoding hisps are nonbioluminescent, and the hisps homologs in bioluminescent species lack two domains, the ketoreductase and dehydratase domains (SI Appendix, Fig. S6). It is likely that the loss of these functional domains in the common ancestor of bioluminescent species favored the production of α-pyrones by ancestral hisps, possibly providing the final step for emergence of bioluminescence.

The gene cluster continued to evolve dynamically after the acquisition of bioluminescence. At least six independent complete or partial gene loss events of genes from the genomic cluster led to secondary loss of bioluminescence (Fig. 2). The cph gene was inserted into the cluster in the nonmycenoid clade, possibly twice (Fig. 1). This mosaic pattern resembles evolutionary history of fluorescent proteins (24), another visually relevant protein family with an unclear biological role, and may indicate that selective advantage provided by bioluminescence in fungi depends on a specific or transient ecological context.

Complex adaptations can be a source of biotechnologically relevant solutions. In addition to revealing the nature of basic photochemical processes and protein evolution, luciferases are among the primary types of reporter genes used in various research pipelines, methods of diagnostics, and environmental applications (5, 6, 25). To determine whether a fungal bioluminescence pathway can provide reporter genes, we characterized nnLuz in vitro and tested its ability to produce light in heterologous systems. We s.c. implanted murine colon carcinoma cells expressing either nnLuz in a whole-body imaging setup of tumor xenografts in mice. We also compared nnLuz qualitatively with the luciferase from the firefly Photinus pyralis in a whole-body imaging setup of tumor xenografts in mice. We s.c. implanted equal amounts of murine colon carcinoma cells expressing either nnluz or firefly luciferase under the same promoter, injected a
mixture of firefly and fungal luciferins i.p., and obtained almost identical signals from the implants (Fig. 3C).

Finally, we aimed to test whether luciferin biosynthesis can be achieved in organisms lacking caffeic acid biosynthesis. Introduction of three additional genes coding for enzymes of caffeic acid biosynthesis from tyrosine, Rhodobacter capsulatus tyrosine ammonia lyase and two E. coli 4-hydroxyphenylacetate 3-monooxygenase components (26), into the genome of P. pastoris strain carrying npgA, hispS, h3h, and luz genes resulted in a strain that was autonomously bioluminescent when grown in a standard yeast medium (SI Appendix, Figs. S12 and S16).

Thus, under all tested conditions, wild-type N. nambi luciferase is functional in heterologous systems, positioning itself as a promising reporter gene, and fungal luciferin can be synthesized from aromatic amino acids in other eukaryotes. In addition, fungal luciferin is a water-soluble and cell-permeable compound, and its light-emitting reaction is not dependent on the availability of ATP, making the fungal bioluminescent system attractive for numerous applications in biomedical imaging. Furthermore, various luciferin analogs can be used to enhance light emission and tune its spectra, improving light penetration in deep tissue imaging applications (22).

In conclusion, we present the enzymatic cascade that leads to light emission in fungi, which is a eukaryotic bioluminescent system with known biosynthesis of luciferin. We have shown that luciferin is synthesized from its precursor hispidin by N. nambi H3H and that hispidin can be directly synthesized by hispidin synthase from caffeic acid, a widespread cellular metabolite with efficient biosynthesis that was achieved in various organisms, including industrially relevant yeast strains (26). Just two enzymatic steps from the mainstream metabolic pathways, the fungal system has a high potential for synthetic biology to create autonomously glowing animals and plants: attempts to develop such organisms have so far been constrained by the poor performance in eukaryotes of the bacterial bioluminescent system, the only system for which luciferin biosynthesis was known (27, 28).

Reconstitution of fungal bioluminescent pathway in eukaryotic organisms might enable applications where tissues or organisms report changes in their physiological state with autonomous light emission. It might also push forward development of the next generation of organic architecture (29), where genetically modified glowing plants will be integrated into buildings and city infrastructure. Apart from that, with its intriguing evolutionary history, a family of luciferases, and overall simplicity, the fungal bioluminescent system presented here is a molecular playground holding numerous opportunities for basic and applied research.

Materials and Methods

SI Appendix includes the details of the materials and methods used in this study, including experiments with Xenopus embryos, mice, yeasts, bacteria, mammalian cells, and bioinformatic analyses. Animal experiments were approved by the local ethical committee of Pirogov Russian National Research Medical University and were carried out in accordance with European Union Directive 2010/63/EU.

Genomes of P. stipitatus, Lentinula edodes, N. gardneri, N. nambi, and M. citricolor and transcriptomes of P. stipitatus, L. edodes, and N. gardneri are available at the National Center for Biotechnology Information Bioproject PRJNA476325. Transcriptomes of N. nambi and M. citricolor, alignments of P. pastoris genome sequencing reads, and alignment of protein sequences of fungal luciferases are available at Figshare (https://figshare.com/articles/ A_genetically_encode_bioluminescent_system_from_fungi/6738953/2). Files used to reconstruct the Agaricale species tree, including raw and trimmed alignments and RAxML resulting files, are available at Figshare.
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