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To cite this version:
Jagadeesh Bayry, Vishukumar Aimanianda, J. Iñaki Guijarro, Margaret Sunde, Jean-Paul Latgé. Hydrophobins-Unique Fungal Proteins. PLoS Pathogens, Public Library of Science, 2012, 8 (5), pp.e1002700. 10.1371/journal.ppat.1002700. pasteur-01673646

HAL Id: pasteur-01673646
https://hal-pasteur.archives-ouvertes.fr/pasteur-01673646
Submitted on 31 Dec 2017

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Hydrophobins—Unique Fungal Proteins

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Microorganisms are often covered by a proteinaceous surface layer that serves as a sieve for external molecular influx, as a shield to protect microbes from external aggression, or as an aid to help microbial dispersion. In bacteria, the latter is called the S-layer, in Actinomycetes, the rod-like fibrillar layer, and in fungi, the rodlet layer [1]. The self-assembly properties and remarkable structural and physicochemical characteristics of hydrophobin proteins underline the multiple roles played by these unique proteins in fungal biology.

What Are Hydrophobins?

Hydrophobins, low molecular mass (≤20 kDa) secreted proteins of fungi, are characterized by moderate to high levels of hydrophobicity and the presence of eight conserved cysteine (Cys) residues. These proteins are able to assemble spontaneously into amphipathic monolayers at hydrophobic–hydrophilic interfaces. Although functional homologues are reported in Streptomyces (chaplins, SapB, and SapT for aerial morphogenesis; [2]), hydrophobins are unique to the fungal kingdom. Fungal genome analyses have indicated that hydrophobins generally exist as small gene families with two to ten members, although certain species contain more members (e.g., C. rugosus displays 33 members; http://www.broadinstitute.org) [3,4]. Hydrophobins show very little sequence conservation in general, apart from the idiosyncratic pattern of eight Cys residues implicated in the formation of disulfide bridges (Cys1–Cys6, Cys2–Cys5, Cys3–Cys4, Cys7–Cys8) [5] (Figure 1). Based on hydropathy plots, solubility and the type of layer they form, hydrophobins are divided into two classes [6, Reference S1 in Text S1], although recent bioinformatics studies suggest that intermediate/different forms can also exist and that many hydrophobins with distinct physicochemical characteristics may have been overlooked in the past [4,7]. In class I, considerable variation is seen in the inter-Cys spacing; these hydrophobins assemble into highly insoluble polymeric monolayers composed of fibrillar structures known as rodlets. The rodlets are extremely stable, can only be solubilized with harsh acid treatments, and the soluble forms can polymerize back into rodlets under appropriate conditions. Despite the low sequence similarity, class I hydrophobins from different fungal species can partially complement each other, indicating their relatedness [8]. The sequence and the inter-Cys spacing are more conserved in class II; the monolayers formed by class II hydrophobins lack the fibrillar rodlet morphology and can be solubilized with organic solvents and detergents.

Hydrophobins at the Interface in the Fungal Life Cycle

Fungi are heterotrophic terrestrial eukaryotes, showing two types of growth morphologies: unicellular yeast and multicellular filamentous forms. Yeasts are hydrophilic and they lack hydrophobins. The vegetative hyphae of filamentous fungi growing on moist environments are also hydrophilic and do not show the presence of rodlets on their surface. In contrast, the aerial hyphae and the asexual spores (conidia) are hydrophobic, due to the presence of hydrophobins. The functions of hydrophobins are related to their high surfactant activity, which results from their self-assembly at hydrophobic–hydrophilic interfaces to form an amphipathic monolayer. The hydrophobin layer reduces the surface tension of the medium or the substratum in/on which fungi grow, allowing them to breach the air–water interface or preventing water-logging while maintaining permeability to gaseous exchange [9]. Spores produced on the aerial structures of filamentous fungi are covered by a hydrophobin rodlet layer that renders the conidial surface hydrophobic and wet-resistant, thus facilitating spore-dispersal in the air. The rodlet-forming hydrophobins are essential for these fungi to complete their biological cycle. In many “wet” fungi (e.g., Conidiobolus obscurus), the rodlet-layer is covered by a mucilaginous extracellular matrix that helps the conidia to bind to the substrate, and once the spores are bound to the host, the rodlet-layer is unmasked for better resistance to the environment [10]. In the basidiomycete Agaricus bisporus, the hydrophobin HypA, found in the peel tissue of the mushroom cap, is suggested to form a protective layer during fruiting body development [11]. In Crpphonectia parasitica, the deletion of the gene coding the class II hydrophobin cryparin generated a mutant incapable of erupting through the bark of the tree [12]. Hydrophobins are also reported to play a role in the surface interaction during infection-related development of M. grisea [13, Reference S2 in Text S1]. In the symbiotic phenotypes

Citation: Bayry J, Aimanianda V, Guijarro JI, Sunde M, Latgé J-P (2012) Hydrophobins—Unique Fungal Proteins. PLoS Pathog 8(5): e1002700. doi:10.1371/journal.ppat.1002700

Editor: Joseph Heitman, Duke University Medical Center, United States of America

Published: May 31, 2012

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Funding: This work was supported by ANR-10-Blanc SVSE 3-009 HYDROPHOBIN; European Community’s Seventh Framework Programme (FP7/2007-2013) under Grant Agreement No: 260338 ALLFUN; and the Commonwealth of Australia under the International Science Linkages program (French-Australian Science & Technology Program 2011). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

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of lichen-forming ascomycetes *Xanthoria* spp., the continuous rodlet-layer seals the apoplast continuum [14].

**Structure of Hydrophobins**

Hydrophobins from both classes have been studied *in vitro* and have been shown to be highly surface active and to form amphipathic monolayers on hydrophobic/hydrophilic surfaces. The crystal structures of the class II hydrophobins HFBI and HFBI from *Trichoderma reesei* have been solved [15,16]. In addition, the structure of the class I EAS protein from *Neurospora crassa* has been determined by NMR [5]. These studies indicate that all hydrophobins share a similar small β-structured core that is dictated by the presence of the four disulfide bonds and that the proteins have large exposed hydrophobic surface regions that give rise to their high surface activity. The structures of the class I hydrophobins *DewA (Aspergillus nidulans)* and *Mpgl (M. grisea)* and the class II hydrophobin from *X. crassa*, as well as the secondary structure of the class I hydrophobins *RodA* and *RodB* from *Aspergillus fumigatus* obtained through the analysis of their backbone NMR chemical shifts, are consistent with this (J. I. Guijarro and M. Sunde, unpublished data). Monolayer formation by class II hydrophobins does not appear to be associated with major conformational changes. In contrast, biophysical analysis of SC3 from *S. commune* and EAS indicate that rodlet formation is associated with significant structural rearrangements, in some cases involving helical intermediates, but always to a final rodlet form with high β-sheet content and amyloid characteristics [5,17, Reference S3 in Text S1]. Digestion and hydrogen-deuterium exchange experiments with SC3 [18] indicated that the Cys3–Cys4 loop is important for adhesion to hydrophobic surfaces and may directly participate in the formation of rodlets. However, truncation [19] and systematic site-directed mutagenesis [20] experiments with EAS have shown that the Cys3–Cys4 loop is not involved in rodlet formation and that the Cys7–Cys8 loop region is crucial for auto-assembly, suggesting that the variability of the sequences of class I hydrophobins may translate into different mechanisms of rodlet formation [18]. Nevertheless, the surface tension seems to be the driving force to recruit class I hydrophobins to the air-water interface where the structural changes from the soluble form to the rodlet conformation take place [21].

**Hydrophobins and Fungus–Host Interactions**

The surface rodlet-layer has a critical role in masking the immunogenicity of airborne fungal spores [22]. By covering the spore surface, the rodlet-layer imparts immunological inertness to the spores and ensures that pathogen-associated molecular patterns (PAMPs) are not recognized by innate and adaptive immune cells, thus preventing the activation of host immune system, inflammation, and tissue damage [22,23,24,25, Reference...
S₄ in Text S1]. Several lines of evidence suggest that the rodlet-layer, which covers the spores of both pathogenic and non-pathogenic fungal species, prevents immune recognition [22,23,25] (Figure 1). In opportunistic pathogen A. fumigatus, the rodlet-layer made up of RodA imparts resistance to NETosis (a process associated with disruption of neutrophil-membranes and release of a mixture of nuclear DNA with a granular content that acts as a neutrophil extracellular trap [NET]) and killing by alveolar macrophages [23,26]. However, removal of RodA and RodB did not affect pathogenicity of A. fumigatus [Reference S5 in Text S1].

In plant-/entomo-pathogenic fungi, hydrophobins are also described as pathogenicity factors, but their precise role in fungal virulence remains to be understood. In the rice blast fungus M. grisea, the hydrophobin Mpg1 is suggested to function as a developmental sensor for appresorium formation, since it is deleted in a mutant of MPG1 gene resulted in a mutant of M. grisea with reduced virulence; the deletion of another hydrophobin gene in M. grisea, MHP1, led also to a loss of viability and a reduced capacity to infect and colonize a susceptible rice cultivar [27]. In Beauveria bassiana, the non-specific hydrophobic interaction between the fungal spore coat hydrophobin and the insect epicuticle is involved in establishing the pathogenicity of the fungus [28].

**Prospective Applications of Hydrophobins**

The potential applications of hydrophobins rely on their ability to reverse the hydrophilic-hydrophobic character of a surface and/or their surfactant capacity. Several biotechnological applications of hydrophobins have been proposed [29, Reference S6–S12 in Text S1]. However, the large-scale applications of hydrophobins might be difficult to implement due to the production cost of recombinant proteins and/or the large-scale requirements of the proteins. In contrast, in the pharmaceutical or in the nanotechnology industry, where the returns of investment are high, it is possible to envisage a potential development for these proteins. For example, the foam and air-/oil-filled emulsion-forming capacity of hydrophobins has been exploited in protecting nanoparticles and drug formulations [30, Reference S13–S16 in Text S1] (Figure 1). From a therapeutic point of view, the degradation-resistance and immunologically inert properties of hydrophobins could be used to generate hydrophobin-based nanoparticles with embedded therapeutically active proteins and molecules that have to be slowly released within the host or transported to a specific body location without being recognized by the host immune system.

Many questions, however, remain unsolved in the study of hydrophobins: for instance, how is the 3D rodlet-structure organized? How are hydrophobins transported to the cell surface? How is the rodlet-layer attached to the spore surface? What are the signals that trigger germination of the spores covered by a rodlet layer? Addressing these questions will reveal the mechanism by which hydrophobins accomplish their multiple roles in the fungal life cycle.

**Supporting Information**

**Text S1** Supplementary references S1–S16.

(ROC)

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