Structural evidence of diffuse growth and parenchymatous cell division in the cortex of the umbilicate lichen Lasallia pustulata

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Abstract: How growth is distributed within the morphologically diverse thalli of lichens is still poorly known and the anatomical mechanisms involved are not well understood. This work applies electron microscopy (SEM and TEM) to examine cell- and tissue-level events in the umbilicate thallus of Lasallia pustulata, whose pattern of expansion was the subject of a previous field study. Stacks of epinecral tissue accumulating at the thallus surface showed broadening bases and recurring rupture attributable to diffuse expansion of the living tissue below. Cortical cells, dividing anticlinally, adjoined septa to previous septa, indicating parenchymatous divisions. These observations are all consistent with previous contentions that mature, organized tissues within the thallus are capable of continued diffuse growth. They provide a developmental explanation for the morphology of the epinecral layer and suggest that anatomical characteristics may be helpful in recognizing diffuse growth patterns. Parenchymatous cell divisions, believed until recently to never occur in lichen thallus tissues, are shown to play a developmental role in the diffuse growth of the umbilicate lichen thallus.

Key words: anatomy, epinecral layer, histogenetic processes, morphogenetic processes, septa

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Introduction

Our understanding of where and how growth occurs in lichen thalli is limited. Previously, lichen growth was assumed to be restricted to the margins and apices of thalli (Hill 1981; Hale 1983), consistent with the notion that the component fungal hyphae grow exclusively at their tips (but see Voisey 2010). In crustose lichens, and foliose taxa that are closely attached to the substratum, expansion may indeed be limited mainly to a marginal zone (Hale 1970; Armstrong 1974; Fisher & Proctor 1978). In foliose lichens with less restrictive attachment, however, much of the thallus may contribute to expansion, as sequential images (Larsson & Gauslaa 2009) and growth rates proportional to size suggest (e.g. Rhoades 1977), at least during early stages of development. In the few fruticose lichens studied, all of which are long pendulous taxa, growth occurs diffusely throughout the thallus, while morphogenetic and histogenetic processes are localized at branch apices (Sanders 1989, 1992; Rolstad & Rolstad 2008; Sanders & de los Ríos 2012; Sanders & Tokamov 2015). However, because lichen thalli are so diverse, many more growth studies are needed before we can assess what generalizations, if any, are warranted.

particularly intriguing in this regard are the umbilicate lichens, whose laminar, dorsi-ventral thalli attach to their rock substratum by a central holdfast. In the opinion of Honegger (2008), their growth patterns are the least understood. Based on carbon fixation rates that varied irregularly throughout individual thalli of Umbilicaria, Larson (1983) inferred that thallus growth is most likely diffuse. Anatomical study of the umbilicate genus Lasallia led Valladares & Ascaso (1994) to postulate that growth occurs mainly in a broad intermediate zone between...
the umbilicus and the margin. Hestmark (1997a) measured the displacement of thallus pustules in *L. pustulata* over a four-year period, reporting a diffuse pattern of growth consistent with the inferences of Larson (1983) and Valladares & Ascaso (1994). His description of pustules arising near the central umbilicus and moving radially outwards over time suggested a pattern of morphogenesis quite different from any proposed previously in macrolichens, where morphogenetic processes are typically localized at apices and margins (e.g. Sanders 1992, 1993). However, no sequential photographs were provided. While geometric increases in length over time were observed in the fruticose *Usnea longissima* (Rolstad & Rolstad 2008), Hestmark (1997b) found no correlation between thallus size and diameter increments over a four-year period in *L. pustulata*. To better understand the structural basis of growth in this lichen, we explore features of tissue behaviour and cell proliferation that would be expected to occur under conditions of diffuse expansion as reported by Hestmark (1997a).

*Lasallia pustulata* has a stratified (heteromeric) thallus with several distinct tissue layers. A cortex of moderately thick-walled, approximately isodiametric cells covers upper and lower surfaces. At its exterior, especially on the upper surface, dead cortical cells accumulate as a thick epinecral layer (Fig. 1), reportedly serving to remutate as a thick epinecral layer (Fig. 1), on the upper surface, dead cortical cells accumulated from below. The anatomical basis of growth may also be evident in the pattern of cortical cell divisions. Diffuse growth would likely require isodiametric cortical cells to proliferate laterally within the plane of expansion (new anticlinal cross walls) to increase the number of cells in the layer, in addition to the vertical divisions (new periclinal cross walls) that supply the overlying epinecral tissue. Such division patterns would directly contradict the long-standing paradigm that vegetative tissues in fungi and lichens are never parenchymatous (Jahns 1973, 1988; Moore 1998). However, a recent study demonstrated that parenchymatous divisions do indeed occur routinely in the cortices of at least some familiar lichens (Sanders & de los Ríos 2017). That study used TEM to distinguish walls of septal origin and show that new septa adjoin them, which is central to the concept of parenchymatous division (Schwendener 1873; Frisch 1935). In the present work, we examine anatomical and ultrastructural features in *L. pustulata* to determine whether their development is consistent with predictions based on the diffuse growth model (Larson 1983; Valladares & Ascaso 1994; Hestmark 1997a).

**Materials and Methods**

Thalli of *Lasallia pustulata* were collected air-dry near El Escorial and Pinilla del Buitrago, Madrid, Spain, in June 2017. Specimens processed for TEM were first hydrated with distilled water using a spray mister and maintained in covered Petri dishes for 24 h. Moist thalli were then hand-sectioned with a razor blade to a thickness of 0.25 mm to facilitate chemical penetration. Tissue was chosen from a pustular region roughly equidistant from the margin and the umbilicus. Sections were placed immediately in fixative solution (2.5% v/v glutaraldehyde in phosphate buffer), followed by postfixation in 1% w/v OsO4 solution, dehydration in an ethanol series, infiltration with Spurr’s low-viscosity resin and subsequent polymerization, according to de los Ríos & Ascaso (2002). Ultrathin sections were cut c. 70 nm thick using a Leica EM UC-6 ultramicrotome (Leica Microsystems GmbH, Wetzlar, Germany). Specimens were oriented so as to section the upper cortex paradermally (i.e. parallel to the surface, the plane in which thallus expansion takes place). Sections were post-stained with lead citrate (de los Ríos & Ascaso 2002) then examined in a JEM-1011 transmission electron microscope (JEOL, Tokyo, Japan). For SEM imaging, air-dried specimen fragments were sputter-coated with gold and observed with an FEIInspect scanning electron microscope (Thermo Fisher Scientific, Waltham, Massachusetts, USA), using the secondary electron imaging mode.
Results

The upper surface of the *L. pustulata* thallus was uniformly covered by adjacent, spine-like, roughly cone-shaped epinecral accumulations with somewhat angular surfaces. They were distinctly tapered from their broad base, which contacts those of adjacent spines, to their narrowing, irregular apex (Fig. 2). Empty
compartments of dead cells were visible with SEM at higher magnification (Figs 2B & 3). One could distinguish the basal outlines of what were formerly individual tufts of epinecral tissue, now subdivided by further tissue rupture (arrows in Fig. 3). On the lower surface, epinecral accumulations were less extensive and cell walls thicker but a
comparable pattern of tissue rupture was observed (Fig. 4).

TEM images of paradermal sections showed anticlinal divisions of cells within the plane of the thallus surface. An evanescent but still recognizable median layer allowed septal walls to be distinguished. New septa were clearly adjoined to previous septa (Fig. 5). At these junctures, the sequence of wall formation was evident: the newer septum adjoined to the outer wall layers of a previous septum, and therefore its median layer did not make contact with that of the older wall (arrowheads in Fig. 5).

Discussion

A number of published studies have explored the anatomy of the Umbilicariaceae from structural, functional or biosystematic perspectives (Frey 1929; Scott & Larson 1983; Krog & Swinscow 1986; Sancho & Balaguere 1989; Valladares & Ascaso 1994; Schroeter & Scheidegger 1995; Valladares et al. 1993, 1998; Valladares & Sancho 1995; de los Ríos et al. 1999). The “areolate-echinate” shape of the epicortical accumulations in Lasallia (Sancho & Crespo 1989) was noted and the ecophysiological significance of the fissures that delimit them was discussed (Valladares 1994). However, the developmental context of these structural features does not appear to have been considered previously. The broadening bases of the epinecral accumulations and their repeated sub-fragmentation through tissue rupture are predictable results of the underlying tissue’s continued expansion in surface area (Figs 2–4). Evidence of cortical septa adjoining perpendicularly to each other within the plane of the thallus surface (Fig. 5) indicates growth processes at the cellular level that have not been proposed previously for umbilicate lichens. These results are consistent with Hestmark’s (1997a) report of diffuse growth in Lasallia and may also be applicable to other members of the family that show a variety of surface tissue rupture patterns (Frey 1929; Scott & Larson 1983). Other foliose and squamulose lichen thalli with fissured epinecral layers and/or thorn-like tapered stacks of cortical material (e.g. Timdal 1984; Vogel 1955; Heidmarsson 1996) might also show diffuse growth patterns and merit further investigation from a developmental perspective.

The basal broadening and fissuring of epinecral accumulations can be compared in a
general way to the formation of bark by cambial activity in dicotyledonous plants. In both cases, layers of dead tissue are augmented continuously by divisions of living tissue beneath, whose growth in surface area causes repeated ruptures in the mature and moribund tissues accumulating above them. In the fruticose lichen *Usnea longissima*, diffuse growth of thallus axes (Rolstad & Rolstad 2008) involves the continued elongation of the medullary cord tissue which provides the thallus with mechanical support. Here too, diffuse elongation disrupts overlying tissue, in this case the living cortex and photobiont layers, from whose fragments the lateral branches or fibrils emerge (Sanders & de los Ríos...
2012). Tissue rupture may be further augmented by the regular cycles of hydration and dehydration to which the poikilohydric thallus is continually subjected. The other fruticose lichens known to have diffuse growth are long pendulous species of Ramalina (Sanders 1989; Sanders & Tokamov 2015); their expanding structural tissues are cortical, with no epinecral layers produced.

Diffuse growth of lichen tissue challenges our understanding of how fungal growth occurs at a cellular level. In Ramalina menziesii, R. usnea and U. longissima, supportive (mechanical) tissue consists of prosoplectenchyma oriented longitudinally along the expanding thallus axes; hyphal tip growth cannot account for diffuse elongation of such tissue. In micrographs of these lichen tissues, cell walls appear to be continuously disrupted by tissue expansion and replaced by new wall layers from within. These massive accumulations of wall material form a kind of intercellular matrix, but the concentric layers evident in transverse section can be attributed to the individual cells that produced them (Sanders & Ascaso 1995; Sanders & de los Ríos 2012). Lichen cortices with isodiametric cells, by contrast, may accommodate diffuse growth by proliferating via parenchymatous divisions (Sanders & de los Ríos 2017), as is evident here in Lasallia pustulata (Fig. 5).

Our results suggest that portions of lichen thalli might transition to parenchymatous division patterns while other thallus layers presumably remain plectenchymatous. Vuillemin (1912) proposed the term “merenchyma” for tissue arising (secondarily?) via three-dimensional divisions in fungi, as opposed to synenchyma, arising by cell fusion. Significantly, he pointed out that de Bary’s application of the term “pseudo-tissue” to fungal structures of filamentous origin inconveniently restricts the concept of tissue itself to those originating from parenchymatous division. We would strongly argue that the term tissue should maintain its broad sense, free of any developmental implications. Without such a concept there is no way to refer to organized arrangements of cells without making unwarranted assumptions about development when no such data are available. Such implicit assumptions are probably the main reason why parenchymatous divisions in lichen “plectenchyma” have remained unacknowledged for so long.

Whether parenchymatous divisions in the cortex of L. pustulata occur uniformly or in a locally variable pattern remains to be determined. Certainly, the protruding morphology of pustules suggests that tissue expansion in those structures might be greater or faster than that of the interpustular regions, at least at some point in their development. A separate but equally interesting question is how the remaining thallus tissues, including both lax and prosoplectenchymatous medullary layers, are themselves accommodating diffuse growth at the cellular level. The potential diversity of thallus and tissue growth patterns among lichens is of considerable mycological interest as the few studies carried out so far have revealed patterns of cell behaviour strikingly different from those characteristic of typical fungal hyphae.

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