Role of the Skp1 prolyl-hydroxylation/ glycosylation pathway in oxygen dependent submerged development of Dictyostelium

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Role of the Skp1 prolyl-hydroxylation/glycosylation pathway in oxygen dependent submerged development of Dictyostelium

Yuechi Xu1, Zhuo A Wang2, Rebekah S Green1 and Christopher M West1*

Abstract

Background: Oxygen sensing is a near universal signaling modality that, in eukaryotes ranging from protists such as Dictyostelium and Toxoplasma to humans, involves a cytoplasmic prolyl 4-hydroxylase that utilizes oxygen and α-ketoglutarate as potentially rate-limiting substrates. A divergence between the animal and protist mechanisms is the enzymatic target: the animal transcriptional factor subunit hypoxia inducible factor-α (HIFα) whose hydroxylation results in its poly-ubiquitination and proteosomal degradation, and the protist E3SCF ubiquitin ligase subunit Skp1 whose hydroxylation might control the stability of other proteins. In Dictyostelium, genetic studies show that hydroxylation of Skp1 by PhyA, and subsequent glycosylation of the hydroxyproline, is required for normal oxygen sensing during multicellular development at an air/water interface. Because it has been difficult to detect an effect of hypoxia on Skp1 hydroxylation itself, the role of Skp1 modification was investigated in a submerged model of Dictyostelium development dependent on atmospheric hyperoxia.

Results: In static isotropic conditions beneath 70-100% atmospheric oxygen, amoebae formed radially symmetrical cyst-like aggregates consisting of a core of spores and undifferentiated cells surrounded by a cortex of stalk cells. Analysis of mutants showed that cyst formation was inhibited by high Skp1 levels via a hydroxylation-dependent mechanism, and spore differentiation required core glycosylation of Skp1 by a mechanism that could be bypassed by excess Skp1. Failure of spores to differentiate at lower oxygen correlated qualitatively with reduced Skp1 hydroxylation.

Conclusion: We propose that, in the physiological range, oxygen or downstream metabolic effectors control the timing of developmental progression via activation of newly synthesized Skp1.

Keywords: Prolyl 4-hydroxylase, Glycosyltransferase, Oxygen sensing, Hypoxia, Hydroxyproline, Cellular slime mold

Background

Cells, whether free-living or residing within multicellular organisms, continuously monitor environmental O2 and integrate this information with other cues to regulate their metabolism, growth and development. Cytoplasmic prolyl 4-hydroxylases (P4Hs) are key O2 sensors in animals [1,2], owing to their ability to distribute the atoms of molecular O2 between the target Pro and the metabolite α-ketoglutarate. The transcriptional co-factor hypoxia inducible factor-α (HIFα) is a main target (Figure 1A), and hydroxylated HIFα is subject to polyubiquitination by the VHL (von Hippel-Lindau protein/cullin-2/elongin B/elongin C) type of E3 ubiquitin ligases leading to subsequent degradation in the 26S-proteasome [2]. Thus low O2 is thought to rapidly induce the expression of new genes appropriate to hypoxia. In contrast, a P4H in the social amoeba Dictyostelium and the human parasite Toxoplasma gondii, known as PhyA (previously referred to as P4H1), appears to solely hydroxylate Skp1 (Figure 1B), at Pro143 [3,4]. Hydroxylation does not affect Skp1 stability [5] but may regulate poly-ubiquitination activity of the SCF (Skp1/cullin-1/F-box) class of E3 ubiquitin ligases, of which Skp1 is an adaptor subunit [6,7]. The 4(trans)-hydroxyproline (Hyp) can then be sequentially modified by 5
sugars whose additions are catalyzed by 5 glycosyltransferase activities encoded by 3 genes [5,8,9]. Reverse genetic analyses demonstrated that hydroxylation and glycosylation of Dictyostelium Skp1 are essential for normal O2 regulation of development [10,11], and recent studies showed its importance for optimal growth of Toxoplasma [4].

Dictyostelium development is ultrasensitive to O2 making it a good model for understanding the mechanism of O2 sensing by other organisms that conserve the Skp1 modification pathway. Development is induced by starvation, which signals the normally solitary phagocytic amoebae to form a multicellular fruiting body, which consists of a cellular stalk that aerially supports thousands of spores for potential dispersal to other locations (see Figure 2A in Results) [12-14]. Initially, the amoebae chemotax together to form a multicellular aggregate, which polarizes in response to environmental cues and elongates into a migratory slug consisting of prestalk cells mostly at its anterior end and prespore cells in the remainder. The slug responds to environmental signals that direct its migration and regulate the slug-to-fruit switch – the process of culmination leading to formation of the fruiting body. Signals include light, low NH3, low moisture, higher temperature, and high O2 which, in the native environment of the soil, draw the subterranean slug to above ground where culmination is most productive [11,12,15-20]. In the laboratory, the process takes place over the course of 24 h after deposition of amoebae on moist agar or filter surfaces wetted with low salt buffers. Whereas amoebae grow and form slugs at an air-water interface in the presence of as little as 2.5% O2, ~10% is required for culmination [21], and slugs immersed in mineral oil require atmospheric hyperoxia to culminate [20]. Overexpression of Skp1 or absence of pathway activity drives the O2 requirement up to 18-21% (near ambient level), whereas decreased Skp1 or overexpression of PhyA drives the O2 requirement down to 5% or less [5,10,11]. These genetic manipulations also revealed effects on timing of slug formation and on sporulation. Together with studies on a Skp1 mutant lacking the modifiable Pro143 residue, and double mutants between Skp1 and pathway enzyme genes, the findings suggested that the Skp1 modification pathway mediates at least some O2 responses. However, O2 contingent modification of the steady state pool of Skp1 has not been demonstrated.

To address this issue, and to investigate the generality of O2 regulation of development, we turned to a previously described submerged development model in which terminal cell differentiation depends on high (>70%) atmospheric O2 [22,23]. The wider range of O2 concentrations presented to cells in this setting may facilitate analysis of the dependence of Skp1 hydroxylation on O2, and absence of the morphogenetic movements of culmination might reveal later developmental steps that are dependent on Skp1 and its modifications. In a static adaptation of the previous shaking cultures, we observed that terminal cell differentiation occurs in a novel radially symmetrical fashion in multicellular cyst-like structures. Under these conditions, we find that O2 is apparently rate-limiting for Skp1 hydroxylation, and that cyst formation and terminal spore differentiation that require high O2 also depend on normal levels of Skp1 and...
both its hydroxylation and glycosylation. This expands the role of Skp1 and its modifications in developmental regulation, and supports the model that O$_2$ regulates its modification in cells.

**Methods**

**Dictyostelium cell strains and growth**
The normal *D. discoideum* strain Ax3 and its derivatives with the following genotypes were described previously: phyA$^{-}$ [3], ecmA::PhyA-myc/phyA$^{-}$, cotB::PhyA-myc/phyA$^{-}$ [24], PKA(cat)/phyA$^{-}$ [24], pgtA$^{-}$ [8], PgtA-N/pgtA$^{-}$ [8], agtA$^{-}$ [25], gmd$^{-}$ [26], ecmA::Skp1A.1/Ax3, ecmA::Skp1A.2/Ax3, cotB::Skp1A.1/Ax3, cotB::Skp1A.3/Ax3, cotB::Skp1A.3/H2/Ax3, ecmA::Skp1B.2/phyA$^{-}$, cotB::Skp1A.2/phyA$^{-}$, cotB::Skp1A.3/phyA$^{-}$ [10]. Note that the number before the decimal point represents alleles, and the number after represents clones that may vary in expression level. Cells were grown in shaking HL-5 axenic medium at 22°C [24], and clones that may vary in expression level. Cells were harvested by centrifugation (2000 g × 1 min) at 4°C, resuspended in PDF buffer (33 mM NaH$_2$PO$_4$, 10.6 mM Na$_2$HPO$_4$, 20 mM KCl, 6 mM MgSO$_4$, pH 5.8), re-centrifuged and resuspended in PDF at 10$^6$/ml, and deposited on 0.45 μm pore Millipore cellulose nitrate filters for standard development at an air-water interface [27]. For submerged development, washed cells were resuspended in PDF at 2 × 10$^6$/ml and 1.4 ml was deposited into each well of a 6-well bacteriological or tissue culture plate (3 cm diameter wells). Plates were incubated for up to 72 h in a sealed plastic box, with inlets and outlet ports for gas flow, under room fluorescent lights at 22°C. The inlet valve was connected via a bubbling water humidifier to a compressed gas tank formulated of 1% CO$_2$. Previously it was shown that inclusion of 1% CO$_2$ did not affect the O$_2$ dependence of culmination [24]. The outlet tube was connected to a Pasteur pipette held under water to monitor gas flow. Cultures were kept unstirred to prevent contact of cells or cell aggregates with the buffer surface, which led to polarization and/or floating fruiting bodies (data not shown). Volume and cell density were optimized for maximal spore differentiation at 100% O$_2$ (data not shown). Alternate buffers, including KP (17 mM potassium phosphate, pH 6.5), or Agg buffer (0.01 M NaPO$_4$, pH 6.0, 0.01 M KCl, 0.005 M MgCl$_2$), yielded lower spor numbers.

Cell aggregates were visualized in a stereomicroscope using transmitted light, or using phase contrast illumination on an inverted microscope. For detection of cellular losic cell walls, samples were analyzed under epifluorescence illumination in the presence of 0.1% (v/v) Calcofluor White ST (American Cyanamid) in 10 mM potassium phosphate (pH 8.0), using DAPI-filters. Multiphoton confocal microscopy was performed at the OUHSC Imaging Laboratory on a Leica SP2 MP Confocal microscope.

For determining spore numbers, samples were supplemented with 0.2% NP-40, and spores were counted in a hemacytometer. Spores were identified based on their resistance to detergent, shape, refractility, and labeling with Calcofluor White ST or anti-spor coat Abs. Spore plating efficiency was determined by spreading an aliquot of detergent-treated spores on SM agar in association with *Klebsiella aerogenes*, and dividing the number of colonies by the counted number of input spores.

**Immunofluorescence**

Spores were released from cysts by probe sonication in 0.2% NP-40 in KP, centrifuged at 13,000 g × 10 s, and resuspended in KP buffer. Spores were recovered from fruiting bodies on non-nutrient agar by slapping the inverted Petri plate on a counter and washing the spores from the lid, and processed in parallel. An aliquot was treated with 6 M urea, 1% (v/v) 2-mercaptoethanol in TBS (10 mM Tris–HCl, pH 7.4, 150 mM NaCl) for 3 min at 100°C prior to dilution in cold TBS and recovery by centrifugation. Spore suspensions (2 × 10$^6$/50 μl) were deposited on glass slides onto which had been dried a 50-μl volume of 10 μg/ml poly-L-lysine in H$_2$O. After 15 min, non-bound spores were removed by aspiration and washing with TBS. The monolayer was incubated in 4 mg/ml hemoglobin in TBS for 5 min, 1 μg/ml mAb 83.5 [28] in 4 mg/ml hemoglobin in TBS for 1 h, TBS (5 washes), 2 μg/ml Alexa 568-conjugated Rabbit anti-mouse IgG (Molecular Probes/Invitrogen) in 3% (w/v) bovine serum albumin in TBS, TBS (5 washes), and Vectashield mounting medium. Samples were analyzed through a 40× (N.A. 0.75) lens via the TRITC-channel of an Olympus epifluorescence microscope, and images were identically recorded using a SPOT Flex camera (Diagnostic Instruments) and processed using Photoshop CS3.

**Western blotting**

Developing cells were collected by centrifugation at 2000 g × 1.5 min at 4°C and boiled for 2 min in Laemmlli sample buffer containing 50 mM DTT. Low O$_2$ samples were first supplemented with 2 mM sodium dithionite [5] to minimize possible hydroxylation during sample preparation. Whole cell lysates were resolved by SDS-PAGE on a 4-12% gradient gel (NuPAGE Novex, Invitrogen), and transferred to nitrocellulose membrane using an iBlot system (Invitrogen). Blots were probed with primary and fluorescent secondary Abs as described [10]. Blots were
blocked in, and Abs were dissolved in, 5% non-fat dry milk in 20 mM Tris–HCl (pH 7.4), 150 mM NaCl, 0.02% NaN₃, and Alexa 680 fluorescence was imaged using a Li-Cor Odyssey scanner. Prespore cell differentiation was probed using mAbs 5F5 and 83.5 [28], and Skp1 isoforms were detected using pAb UOK87 [5], pAb UOK85 [5], mAb 4H2 [29], mAb 1C9 [29], and mAb 4E1 [3]. Affinity-purified anti-actin was from Sigma Chemical Co.

Images were analyzed densitometrically using NIH Image J. mAb 4E1 was used in its linear response range [10] to obtain the fraction of Skp1 that was not modified. Initially, values for each upper and lower band were corrected for general background by subtraction of a blank intensity value obtained from the vicinity of the band of interest. Studies using pAb UOK87, which selectively recognizes unmodified Skp1, showed that 5% of Skp1 was unmodified at 100% O₂ based on comparison with a phyA⁻ sample (not shown). The remaining density in the lower band of the 100% O₂ sample is of uncertain identity but, since its level was observed to be proportionate to the level of the upper band (not shown), its value (as a fraction of the upper band) was subtracted from each sample in the O₂ series. The fraction of unmodified Skp1 was determined by dividing the corrected intensity of the lower Skp1 band by the sum of the intensities of the lower and upper bands.

**Results**

**Terminal differentiation at an air-water interface**

*Dictyostelium* amoebae develop to form fruiting bodies when dispersed in a low ionic strength buffer on a moist surface (Figure 2A). About 75% of the cells become aerial spores and the remainder form the structural stalk. At reduced O₂ levels (2.5-10%), the slug intermediate continues to migrate on the surface without culminating [24]. When returned to the ambient O₂ level (21%), culmination then occurs within about 5 h. To determine the minimal time required for exposure to ambient O₂, slugs were exposed to 21% O₂ for varying times before returning to low O₂. Figure 2B shows that exposure to high O₂ can be as brief as 1 h, though up to 4 h is required for maximal culmination based on spore counts. The requirement for high O₂ appeared to be selective for induction of culmination, because terminal cell differentiation occurred normally even within the fruiting bodies formed after only 1 h of exposure to normoxia (data not shown). The effect of O₂ appears to be mediated at least in part by prolyl 4-hydroxylation of Skp1, because elevated O₂ levels are required by phyA⁻ and Skp1-overexpression strains, and lower O₂ is required by PhyA overexpression and Skp1B⁻ cells [10,24].

**Terminal differentiation in submerged cultures**

When normal strain Ax3 cells were incubated at a similar density under a height of several mm of PDF buffer under room light illumination, rather than on a surface wetted with the same buffer, development proceeded only to the loose aggregate stage. However, when the atmosphere above the culture was maintained at 70 or 100% O₂, the majority of cells formed tight spherical aggregates with diameters of 100–250 μm (Figure 3A) and optically dense cores (see Figure 4D below). These cell aggregates were uniformly bounded by Calcofluor-positive stalk cells, distinguished by their polygonal shapes due to cell expansion during terminal differentiation (Figure 3A). Confocal microscopy revealed that the stalk cells comprised a cortex surrounding an interior region of spore-like cells, based on their characteristic ellipsoid profiles, with an uneven boundary at the interface (Figure 3B). Note that Figures 3 and 4 also include comparative data on phyA⁻ cells (which do not modify Skp1), which will be described below. The interior cells could be liberated under pressure and consisted of a mixture of spores and undifferentiated (Calcofluor-negative) cells (Figure 3D). In contrast, the stalk cells remained associated with the deflated cyst-like structures. Maximal spore number was achieved by 2 d
Figure 3 (See legend on next page.)
cells were agitated in high O2 [22,23]. The radially polarized
and linearly polarized aggregates formed when
ments of culmination. This contrasts with the slug-like
ion in the absence of the normal morphogenetic move-
cell differentiation occurred in radially symmetrical fash-
fruiting bodies on filters, which was 66%. Thus, terminal
spores was 70%, similar to that of spores collected from
clusters or single cells (not shown). At 40% O2, larger
under these conditions. When present they occurred as
observed in the less compacted aggregates that form
(Figures 3C,E). Though spores were not detected in this
similarly imaged, but exposure was adjusted to show fluorescence of expelled cells (absent in panel C), resulting in overexposure of the stalk cell-rich case. The point of emergence (rupture) of interior cells is indicated in panel E. (F) Spore coat formation. Spores from normal fruiting bodies
developed at an air-water interface, and from submerged cultures maintained for 3 d under 70% O2, were compared by immunofluorescence
labeling with mAb 83,5, which recognizes the fucose epitope predominantly on the spore coat proteins SP96 and SP75. Spores were labeled
before or after extraction with urea/2-mercaptoethanol to permeabilize the coat. Control samples lacking mAb 83,5 exhibited only dim internal
fluorescence (not shown).

(See figure on previous page.)

Figure 3 Cell differentiation in submerged conditions. Typical cyst-like structures formed by unstirred suspensions of strain Ax3 (normal) or
phyA+ cells under the atmosphere of O2 percentage and for the duration indicated. (A) An Ax3 aggregate formed in 100% O2 was imaged by
phase contrast (above), or epifluorescence microscopy in the presence of Calcofluor White 5T (below) to reveal cell walls of terminally
differentiated stalk cells at the aggregate surface. (B) Visualization of the interior of Ax3 and phyA+ aggregates using multiphoton confocal
fluorescence microscopy in the presence of Calcofluor. (C) Aggregates of Ax3 or phyA+ cells formed under 40% O2 were squashed by applying
vertical pressure to the cover slip, expelling some of the cellular contents resulting in wrinkling of the aggregate surface (evident as concentric
folds appearing as rings). Cells were imaged for Calcofluor fluorescence. (D, E) Aggregates formed under 100% O2, (D) or 40% O2, (E) were
similarly imaged, but exposure was adjusted to show fluorescence of expelled cells (absent in panel C), resulting in overexposure of the stalk cell-rich case. The point of emergence (rupture) of interior cells is indicated in panel E. (F) Spore coat formation. Spores from normal fruiting bodies
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before or after extraction with urea/2-mercaptoethanol to permeabilize the coat. Control samples lacking mAb 83,5 exhibited only dim internal
fluorescence (not shown).

(Figure 4A), and ranged from 6 to 33% of the input cell
number. These spores tended to be less elongated than
their counterparts formed in fruiting body sori (see Figure 3F below), suggesting imperfect synchronization
of spore coat assembly processes [28]. To test their au-
thenticity, spores were released by probe sonication in a
non-ionic detergent, which ruptured the cyst-like cells
and lysed non-spore cells. Spores from cysts were
on average slightly more brightly labeled than authentic
spores isolated from fruiting bodies by immunofluores-
cence probing with mAb 83,5, which binds to the fucose
epitope associated with the spore coat proteins SP96 and
SP75 (Figure 3F). Surface labeling was retained even
after boiling the spores in urea, indicating tight associ-
ation of residual coat proteins with spore coat. To test
spore function, equal numbers of spores prepared in this
way were serially diluted in a clonal assay in association
with K. aerogenes bacteria. The plating efficiency of cyst
spores was 70%, similar to that of spores collected from
fruiting bodies on filters, which was 66%. Thus, terminal
cell differentiation occurred in radially symmetrical fash-
ion in the absence of the normal morphogenetic move-
ments of culmination. This contrasts with the slug-like
elongated and linearly polarized aggregates formed when
cells were agitated in high O2 [22,23]. The radially polar-
ized organization may result from a more uniform envir-
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cell differentiation occurred in radially symmetrical fash-
ion in the absence of the normal morphogenetic move-
ments of culmination. This contrasts with the slug-like
elongated and linearly polarized aggregates formed when
cells were agitated in high O2 [22,23]. The radially polar-
ized organization may result from a more uniform envir-
oment presented by the static setting in which
polarizing gradients of O2 or NH3 fail to form.

Under 21% O2, stalk cells and spores were rarely
observed in the less compacted aggregates that form
under these conditions. When present they occurred as
clusters or single cells (not shown). At 40% O2, larger
aggregates were formed but they lacked dense cores
observed at higher O2 levels. These cyst-like aggregates
possessed a stalk cell cortex but their interior cells pro-
duced few spores, as visualized after squashing
(Figures 3C,E). Though spores were not detected in this
example, variable numbers were observed over the 5 in-
dependent trials as quantitated in Figure 4C. The vari-
ation suggests that 40% O2 is close to the threshold
required for sporulation whose exact value is likely influ-
enced by other factors, as observed for culmination [24].
To address the differentiation status of cells at the lower
O2 levels, extracts were Western blotted for the spore
coat precursor proteins SP85, SP96 and SP75 that are
markers of prespore cell differentiation [31]. Whereas all
3 glycoproteins appeared in Ax3 cells by 24 h at 70% O2,
negligible expression occurred at 20% after 3 d
(Figure 4E). Thus increasing O2 levels were required for
tight aggregate formation, terminal stalk cell differenti-
ation, and differentiation of the interior prespore cells
into spores. It is likely that metabolic O2 consumption
results in intracyst hypoxia in these unstirred cultures
which, in the submerged state, is not adequately replen-
ished by O2 diffusion. The finding that elevated O2 ten-
sion in the atmosphere above the medium can rescue
terminal differentiation indicates that O2 availability is
the limiting factor for terminal cell differentiation in this
setting. It is not evident whether the higher O2 level
required for spore compared to stalk cell differentiation
reflects a higher O2 threshold requirement for spore dif-
ferentiation or lower O2 in the aggregate centers.

Requirement of PhyA for sporulation in
submerged conditions
A previously described mutant strain disrupted at its
phyA locus [24] was analyzed to determine the involve-
ment of Skp1 prolyl 4-hydroxylation in submerged de-
development. phyA+ cells formed cyst-like structures at 40-
100% O2 with outer layers of differentiated stalk cells,
similar to the normal Ax3 strain (Figure 3C, D). How-
ever, interior cells failed to differentiate as spores, even
after extended periods, as shown in the side-by-side
comparisons in Figures 3B, D, 4A, and D. Instead, they
remained as prespore cells, based on Western blot anal-
ysis showing abundant expression of the spore coat pre-
cursors (Figure 4E). Failure to sporulate was due to the
PhyA deficiency, because phyA− cells complemented
with ecmA::phyA or cotB::phyA, which overexpress PhyA
activity in prestalk or prespore cells respectively [24],
Figure 4 (See legend on next page.)
were rescued at high O₂ (Figure 4B). ecmA::phyA/phyA⁻ cells formed normal numbers of spores compared to Ax3, while cotB::phyA/phyA⁻ only partially rescued spore formation to about 30% of Ax3 levels. The difference suggests that prestalk cells may be important in mediating the role of PhyA in sporulation, consistent with evidence for a role of prestalk cells in processing or mediating sporulation signals during normal culmination [32-34]. While overexpression in prespore cells (cotB promoter) was also partially effective, the possibility that PhyA signals autonomously in prespore cells is not proved because on filters, cotB::PhyAoe cells tend to migrate to the tip in chimeras with normal cells [24]. Successful complementation from these developmental promoters confirmed that cells had differentiated into prestalk and prespore cells in the absence of PhyA, and showed that PhyA is required only after their appearance. Since spore formation selectively depended on high O₂ and the threshold for spore (but not stalk cell) differentiation was specifically affected by the absence of PhyA, PhyA activity appears to have a novel function in mediating O₂ regulation of sporulation differentiation.

Since overexpression of PhyA in a phyA⁺ (wild-type) background reduces the O₂ level required for culmination on filters [24], the effect of PhyA overexpression on sporulation was investigated. As shown in Figure 4C, modestly increased sporulation was observed at 70% O₂ when PhyA was overexpressed in prespore cells. However, overexpression in prestalk cells inhibited sporulation, without affecting cyst formation per se. As noted above, PhyA overexpression under the ecmA promoter in a phyA⁺ background rescued sporulation better than under the cotB promoter, so the inhibitory effect of overexpression in phyA⁺ cells appears to be dependent on a complex interplay between relative levels of expression in the different cell types rather than a cell autonomous effect on prestalk cells.

Skp1 modification is O₂ dependent
To determine if Skp1 hydroxylation is affected by O₂ availability, its modification status was assessed by Western blotting with pan- and isoform-specific Abs. Extensive analysis of soluble Skp1 from growing and developing cells shows that ≥90% of the steady state pool is homogenously modified by the pentasaccharide, and ~5% exists in unmodified form. Fully modified and unmodified Skp1 migrate as a doublet in SDS-PAGE and, though the resolution of the doublet is compromised when whole cell extracts are analyzed, isoform-specific Abs indicate that total cell Skp1 is modified to a similar extent [5,10]. After 1 d of submerged development, total Skp1 from 40, 70 or 100% O₂ cells migrated mainly as the upper band using mAb 4E1 that recognizes all Skp1 isoforms (Figure 5B). In comparison, 5% O₂ cells accumulated substantial Skp1 in the position of the lower band. This band corresponds to unmodified Skp1 based on reactivity with pAb UOK87 (Figure 5A). UOK87 preferentially binds unmodified Skp1 but exhibits weak reactivity with all Skp1 isoforms, so the upper band is also labeled. The lower band was not recognized by pAb UOK85 or mAb 1C9, which are specific for HO-Skp1 and GlcNAc-O-Skp1, respectively (data not shown). Quantitation of 5 independent samples indicated that the fraction of unmodified Skp1 decreased from 41% at 5% O₂ to 24% at 21% O₂ and 5% at 40% and higher levels (Figure 5D). Similar results were observed after 2 d of development except that the fraction of unmodified Skp1 at the lower O₂ levels was slightly increased (data not shown). Since Skp1 turns over slowly with a half-life of 12–18 h during filter development [5,35], it is likely that the appearance of non-glycosylated Skp1 was the result of new synthesis and that at 5 and 21%, O₂ is rate limiting for Skp1 hydroxylation. As shown in panel E, sporulation depended on higher levels of O₂ than required to hydroxylate Skp1. Although 40% O₂ was sufficient to ensure that the steady-state pool of Skp1 was maximally hydroxylated within the sensitivity of our assay, a delay in hydroxylation of nascent Skp1 of several hrs would have escaped our detection, and may be biologically relevant for sporulation (see Discussion).

Role of glycosylation in submerged development
Disruption of phyA also blocks hydroxylation-dependent glycosylation of Skp1, which occurs according to the scheme in Figure 6A. To investigate the role of
glycosylation per se, gnt1.3, pgtA⁻, gmd⁻, pgtA-N/pgtA⁻, and agtA⁻ cells, which accumulate Skp1 with zero, one, two, two, or three sugars respectively [5,8,26] on account of enzyme gene disruptions, were analyzed. The strains expressing up to two sugars formed cyst-like structures which, however, failed to acquire dense-cores or induce spore formation, like phyA⁻ cells (Figure 6B, C). In contrast, agtA⁻ cells, which accumulate the trisaccharide form of Skp1 [25], were inconsistent in spore formation with numbers ranging from essentially zero to more than Ax3. Thus although the final two sugars were not always required for sporulation, their absence appears to make sporulation vulnerable to an unknown variable. Potential sources of variation include NH₃ and light, which were previously shown to influence the O₂ threshold for culmination on filters [24], and conditioned medium factors previously detected during submerged development [30]. Taken together, the results suggest that the role of hydroxylation may be simply to support glycosylation. This contrasts with culmination, in which hydroxylation alone partially rescues the normal O₂ requirement of phyA⁻ cells [5], an effect that is reversed by the action of PgtA in the absence of AgtA [9].

**Role of Skp1 and its modifications in submerged development**

The role of Skp1 itself was investigated by overexpression in different genetic backgrounds. Native Skp1 sequences were employed because a previous study showed that N- or C-terminal peptide tags interfere with its hydroxylation and activity in cells [10]. Overexpression of Skp1B under the ecmA (prestalk) promoter inhibited tight aggregate formation even at 100% O₂ (Figure 7A-2). No spores (Figure 7B) and few stalk cells (not shown) were observed, confirming inability to progress past this early stage. Similar results were observed with a strain overexpressing the closely related isoform Skp1A (which differs by a single amino acid), or when either Skp1 was expressed under control of the cotB promoter (Figure 7B). However, overexpressing mutant Skp1A3(P143A), which cannot be modified, did not interfere with aggregation (Figure 7), and wild-type Skp1 overexpression failed to inhibit cyst formation in the absence of PhyA (Figure 7A-4). These strains did not form cyst-like structures or spores at lower O₂ levels (data not shown), implying that high O₂ also provides an additional, possibly metabolic, function important for development. The opposing effects of Skp1 overexpression and blocking its modification suggests that modification stimulates Skp1 activity, which can be modeled as breakdown (by a specific E3SCF⁻ubiquitin ligase) of a hypothetical activator of cyst formation.

In comparison, the requirement of Skp1 glycosylation for sporulation suggests that for this later developmental step, Skp1 contributes to the breakdown of a hypothetical inhibitor of sporulation. Without modification, Skp1 is not activated and the inhibitor accumulates. However, overexpression of Skp1 in the phyA⁻ background (thereby bypassing the block to cyst formation) allows sporulation, which can be interpreted as providing additional activity to compensate for lack of activation by modification (Figure 7B, blue bars and inset; data not shown).
shown). Similar effects were observed irrespective of the promoter used, or whether wild-type Skp1A or B, or mutant Skp1, was overexpressed (data not shown). However, overexpression of Skp1 at very high levels did not rescue sporulation in phyA– cells as well, which might reflect a dominant negative effect toward SCF-complex formation. Separate effects on activators and inhibitors may depend on involvement of distinct F-box proteins.

Discussion

Three novel observations regarding development under submerged conditions are presented here: i) In the presence of high O2 and absence of stirring, cell differentiation occurs in a radially symmetrical rather than the typical linearly polarized pattern. With their outer husk-like cortex and interior germinative cells, these structures have the organization of multicellular cysts as occur in animal tissues. The cyst-like structures are distinct from other terminal states formed by Dictyostelium, including the dormant unicellular microcyst and the multinucleated macrocyst [36]. Although conditions leading to the formation of cyst-like structures are not known to occur naturally, its O2 dependence is likely to be relevant to interpreting O2 signaling in normoxia as outlined below. ii) Skp1 hydroxylation is limited by O2 availability. iii) Certain developmental transitions that occur during submerged development, including tight aggregate formation and terminal spore differentiation, critically rely on hydroxylation and glycosylation of Skp1. Together, these findings reinforce a role for environmental O2 for influencing polarity and key developmental transitions, and strongly implicate the Skp1 modification pathway in decoding the O2 signal.

Significance of O2 for control of polarity and terminal differentiation

Formation of the novel cyst-like structures is compared to normal development at an air-water interface as a backdrop to interpreting the role of Skp1 modification in O2 signaling. During normal development at an air-water interface, the tip emerges at the apex of the hemispherical aggregate and exerts a dominant role in

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**Figure 6** Dependence of spore differentiation on Skp1 glycosylation. (A) Schematic of the Skp1 modification pathway. Enzymes are indicated by gene names; G = Gal; F = Fuc; G = GlcNAc. (B) Normal cells (Ax3) and modification pathway mutants were developed at 100% O2 in submerged conditions for 72 h. All strains formed similar tight aggregates, except that Ax3 aggregates exhibited dense cores; agtA– aggregates formed few spores in this trial. (C) Spore numbers were determined as in Figure 4. Average values ± SEM from 5 independent trials are shown. The wide error bar for agtA– cells results from a range of outcomes from near zero to more than Ax3.
controlling elongation into a slug, slug migration, internal cell dynamics, and the induction and orchestration of the morphogenetic movements of culmination [13,14,16,20,37]. The tip, composed of prestalk-type cells [38], senses environmental signals, including O2 potentially, and relays the information to the other slug cells to follow suit (Figure 8, upper track). In previous submerged development studies, cells were shaken under an atmosphere of high O2 and the aggregates elongated into slug-like structures in which prestalk and prespore cells segregated toward opposite ends and terminally differentiated in situ [22,30,31,39]. In the absence of stirring as described here, cell aggregates instead become spherical cysts in which internal prespore and spore cells are surrounded by stalk cells. These findings suggest that O2 contributes to patterning and terminal differentiation, as follows (Figure 8, lower track). In previous submerged development studies, cells were shaken under an atmosphere of high O2 and the aggregates elongated into slug-like structures in which prestalk and prespore cells segregated toward opposite ends and terminally differentiated in situ [22,30,31,39]. In the absence of stirring as described here, cell aggregates instead become spherical cysts in which internal prespore and spore cells are surrounded by stalk cells. These findings suggest that O2 contributes to patterning and terminal differentiation, as follows (Figure 8, lower track). Given that O2 is metabolically depleted in the aggregate center, a gradient of O2 occurs with the highest levels at the aggregate surface [39] where the O2 level is expected to be uniform all they way around. Based on studies in capillaries [40] and in agar immobilized aggregates [39], it is likely that the higher O2 level at the aggregate surface attracts spontaneously differentiated prestalk cells and triggers their terminal differentiation. This is consistent with the transient existence of a monolayer of prestalk-like cells that has been observed at the slug surface [41]. Higher than ambient O2 might be required as a consequence of the submerged condition in which replacement diffusion of O2 lags behind metabolic consumption. In the absence of orienting signals in this isotropic setting, the aggregate remains radially-polarized. However, at the air-water interface, tip formation initiates at the apex of the aggregate owing to highest O2 accessibility, which becomes stabilized as its smaller radius of surface curvature ensures greatest gas exchange with the underlying cells. The interior prespore cells, experiencing relative hypoxia owing to metabolic consumption of O2, might not normally differentiate until culmination permits aerial exposure to atmospheric O2 levels or modulates metabolites that regulate PhyA and the glycosyltransferases. The idea that hypoxic niches regulate cell differentiation has precedent in studies on animal stem cells and maize germ cells [42,43].

**Figure 7 Dependence on Skp1 expression level.** Strains overexpressing Skp1A or Skp1B under control of either the *ecmA* or *cotB* promoter, in either a normal or *phyA*– background, were developed in submerged conditions beneath 100% O2 for 72 h. Typical results from selected strains are shown. (A) All tested strains formed tight aggregates except those overexpressing wild-type Skp1 in a *phyA*+ background, which remained as small, loose aggregates. (B) Spore numbers were counted and normalized to Ax3 spore counts in the same trial. Average values ± SEM from 2–3 independent trials are reported. Inset shows a Western blot for Skp1, showing its level of overexpression in the *phyA*– strains.
NH₃, a volatile metabolite released during the massive breakdown of protein during development [44], has also been implicated as a polarity factor and inhibits the slug-to-fruit switch [16]. Since NH₃ is expected to diffuse away most at the same surfaces that O₂ is expected to diffuse in, the two compounds may play complementary inhibitory and activating roles that tune developmental decisions. Thus, while hypoxic or phyA–precursors may still form tips at the air-water interface [24] due to the NH₃ effect, the spherical shapes assumed by phyA–slugs after long periods of migration [5] might reflect eventual depletion of the NH₃ signal as protein is finally consumed. The isotropic environment during static submerged development may thwart formation of orienting NH₃ as well thereby resulting in radial polarization, and high NH₃ in the interior is expected to promote sporulation [45]. Since NH₃-signaling is mediated in part by NH₃-transporter/sensors [16,17], investigation of genetic interactions with phyA may allow understanding of the interplay with Skp1 modification.

Role of Skp1 prolyl hydroxylation in tight aggregate formation

Tight aggregate formation depended on an elevated O₂ level of ≥40%, but this was inhibited when Skp1 (either isoform) was overexpressed under either developmental promoter (Figure 7A). This correlates with the 7-hr delay of the loose-to-tight aggregate transition of these overexpression strains at the air-water interface [10]. Interestingly, inhibition of tight aggregate formation was partially relieved when Skp1 was overexpressed in a phyA-mutant background, which also relieved the delay on filters. Consistent with a requirement for modification, overexpression of Skp1A3(P143A), which cannot be hydroxylated, is not inhibitory (Figure 7A, B). The opposing effects of Skp1 overexpression and inhibiting its modification are consistent with a model in which modification activates Skp1 and its role in polyubiquitination and breakdown of a hypothetical activator of cyst formation.

Role of Skp1 prolyl hydroxylation and glycosylation in sporulation

A second function of the pathway was revealed by the essentially complete failure of the interior prespore cells to differentiate in the phyA– strain, whereas stalk cell differentiation was qualitatively unaffected (Figures 3, 4). The blockade was overcome when PhyA was overexpressed in prestalk and to a lesser extent prespore cells

![Figure 8 Model for O₂ regulation of development and dependence on the Skp1 modification pathway](image-url)

Cells form loose aggregates that condense into tight aggregates by a Skp1-associated, O₂ dependent mechanism based on submerged development studies (Figures 4D, 7A-2). At a conventional air-water interface (upper track, depicted by dashed line), the exposed apical surface supports maximal O₂ transport which is proposed to induce tip formation whose smaller radius of curvature encourages even more O₂ transport, stabilizing the tip as an organizer and the zone where cells differentiate as prestalk cells. The Skp1 modification pathway, under regulation of O₂ and other factors, regulates culmination and sporulation at the air-water interface. Under submerged conditions (lower track), metabolism consumes O₂, which becomes depleted owing to slow diffusion in the unstirred cultures. In this isotropic and hypoxic environment, all (not just apical) surface cells become prestalk cells and, in the absence of a polarizing O₂ gradient, cells differentiate in situ and those that sense the lowest O₂ level (which occurs in the center) become spores. O₂ action may be complementary to NH₃, a volatile inhibitor that is generated during development and is preferentially lost from the same surfaces by diffusion (see Discussion).
(Figure 4B), so control by O2 may be mediated via pre-stalk cells. This is consistent with evidence that pre-stalk cells can regulate sporulation via processing of spore differentiation factor-1 and –2 [33,34]. However, the role of PhyA appears complex because overexpression in pre-stalk cells in the phyA+ (wild-type) background inhibited sporulation, as if relative levels of O2 signaling between cell types could be important. The blockade was also partially overcome when PKA activity was promoted by overexpression of its catalytic domain under its own promoter (Figure 4B). Since PKA expression in prespore cells was previously shown to be sufficient for activating sporulation [46], PhyA may signal upstream of PKA as suggested for its role in culmination on filters [24].

Hydroxylated Skp1 is a substrate for Gnt1 that in turn generates a substrate for PgtA, and then AgtA, resulting in formation of the pentasaccharide on Hyp143 (Figure 6A). Mutants lacking enzymes to extend to the trisaccharide state were also unable to sporulate at high O2 (Figures 6B,C), suggesting that hydroxylation supports extension of the glycan chain to three or more sugars to trigger sporulation. Though the preceding culmination step (on filters) exhibited more modest dependence on addition of the first two sugars (at lower O2 levels) [5], the more dramatic difference in the static submerged model may simply result from failure to achieve a critical threshold of O2 in the cyst interior. The greater difference was in the role of AgtA, whose contribution was almost as important for culmination as PhyA [9] but was unnecessary for submerged sporulation. Thus the role of AgtA appears to be specialized for culmination compared to sporulation.

The requirement of PhyA for sporulation was partially overcome by overexpression of Skp1 (Figure 7). This suggests that PhyA action normally promotes Skp1 activity, and its absence can be bypassed by excess Skp1. A related effect was observed on filter development, where Skp1 overexpression inhibited sporulation at high O2 levels that allowed culmination, but removal of PhyA blocked inhibition [10], indicating that PhyA tunes Skp1 activity. This is consistent with activation of Skp1 polyubiquitination activity toward an inhibitor. In comparison, the effect of Skp1 modification on culmination implied inhibition of Skp1 breakdown activity toward a hypothetical activator [10,11], and the effects on cyst formation (assessed morphologically) above suggested activation of breakdown activity toward an activator. These disparate effects are consistent with what is known about the SCF family of E3 ubiquitin-ligases, which polyubiquitinate different substrates depending on which F-box protein is present. Furthermore, these Ub-ligases can have opposite effects via auto-polyubiquitination of the F-box protein itself, which results in protection of the substrate receptor [6,7]. Conceivably, Skp1 modification may selectively affect these different activities.

O2 is limiting for Skp1 hydroxylation in submerged culture and mechanistic implications

In submerged development, substantial levels of unmodified Skp1 (Figure 5D) accumulated at 5% and 21% O2. Since i) there is no evidence for enzymatic reversal of hydroxylation or glycosylation, ii) the level of Skp1 was similar at different O2 levels, and iii) Skp1 turns over with a half-life of 12–18 h [5], it is likely that appearance of unmodified Skp1 was due to failure to hydroxylate nascent Skp1. Since the total Skp1 pool becomes 95% hydroxylated at ≥40% O2 (Figure 5D), O2 is likely rate-limiting for Skp1 prolyl hydroxylation. This is consistent with co-expression evidence that PhyA is rate limiting for Skp1 hydroxylation [10]. Since sporulation is minimal at 40% O2 even though the steady-state pool of Skp1 appears fully modified, it may be that O2 and PhyA have additional or alternative mechanisms for controlling sporulation. However, it should also be considered that a several hour delay in the hydroxylation of nascent Skp1, which might be most important for partnering with nascent F-box proteins, would have escaped detection against the background of total Skp1 using our methods.

Since the Skp1/F-box protein complex is characterized by a high affinity [29] that is increased by hydroxylation as suggested in Figure 1B (M.O. Sheikh and C.M. West, unpublished data), we propose that even transient accumulation of unmodified Skp1 will influence the spectrum of complexes with one or more of the ~38 predicted F-box proteins that are strongly up and/or down-regulated at various times during development based on RNAseq data [47] (unpublished studies). This in turn may affect the timing of developmental transitions via effects on the stability of F-box proteins and hypothetical F-box protein substrates (activators and inhibitors) that normally control aggregation, slug formation, culmination and sporulation [e.g., 48]. Figure 2B shows that O2 exposure of 1–3 h can rescue culmination of hypoxic slugs, consistent with a transient role that might correlate with expression of a specific F-box protein. Current studies are focused on how Skp1 modification influences E3SCF ubiquitin-ligase assembly and activity.

These findings in social amoebae may be pertinent to numerous protist groups, including other amoebae (e.g., Acanthamoeba), plant pathogens (Phytophthora), diatoms (brown algae), green algae (Chlamydomonas), ciliates (Tetrahymena), and apicomplexans including Toxoplasma, whose O2 dependence have been little studied but whose genomes harbor Skp1 modification pathway-like genes [11]. For example, recent studies [4]
showed that the related Skp1 modification pathway supports growth of *Toxoplasma* in cultured fibroblasts especially at low O₂.

**Conclusions**

In an isotropic submerged environment under high O₂, starved *Dictyostelium* cells form cyst-like structures in which terminal differentiation occurs in a radially symmetrical pattern consisting of external stalk cells and internal spores. Low O₂ is rate-limiting for the hydroxylation and subsequent glycosylation of Skp1, which correlates qualitatively with inhibition of spore differentiation. Genetic perturbations indicate the importance of Skp1 hydroxylation and glycosylation for activating Skp1 activity in regulating cyst formation and sporulation, in addition to previous evidence for its inhibition in regulating culmination at an air-water interface. The findings support a model in which environmental control of Skp1 modification differentially influences sequential developmental transitions via poly-ubiquitination and degradation of F-box proteins and their respective regulatory factor substrates.

**Abbreviations**

Hyp: (4R,2S)-pro-hydroxyproline (aka 4-trans-hydroxy-L-proline); mAb: Monoclonal antibody; pAb: Polyclonal antibody; PhyA: Prolyl 4-hydroxylase-1 from *D. discoideum*; PKA: Protein kinase A; SCF: E3 ubiquitin ligase sub-complex consisting of Skp1, a cullin-1, an F-box protein, and Rbx1; SEM: Standard error of the mean; Ub: Ubiquitin.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

RSG performed initial experimentation including optimizations. ZAW broadened the scope of the study to include the complete mutant panel, and wrote the first draft. YX confirmed all the findings and conducted most of the molecular characterizations. CMV trained the students, coordinated the study, conducted some experiments, and wrote the manuscript which was approved by all authors.

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