Noncircadian oscillations in amino acid transport have complementary profiles in assimilatory and foraging hyphae of Phanerochaete velutina

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Summary

• Cord-forming woodland basidiomycete fungi form extensive, interconnected mycelial networks that scavenge nitrogen (N) efficiently. We have developed techniques to study N dynamics in such complex mycelial systems in vivo.
• Uptake and distribution of the nonmetabolised, 14C-labelled amino-acid analogue, \( \alpha \)-aminoisobutyrate (14C-AIB) was continuously imaged in Phanerochaete velutina growing across scintillation screens using an enhanced photon-counting camera.
• Oscillations in the 14C-AIB signal were observed for both the assimilatory hyphae in the inoculum and the foraging hyphae, but with complementary profiles. Pulses were asymmetric, with an abrupt switch between each exponential decay phase and the next rising phase. The period of the oscillations was 16 h at 21°C, but showed a strong temperature dependence with a temperature coefficient of 2.1. Oscillations occurred in the absence of obvious pulses in growth.
• Some, but not all, of the features of the oscillations were simulated using a model of amino acid accumulation and transport that included both vacuolar uptake, and release once an intravacuolar concentration threshold was exceeded. The combination of imaging and modelling provides a useful framework to understand N fluxes in vivo.

Key words: Phanerochaete velutina, amino acid transport, vacuole, \( \alpha \)-aminoisobutyrate, metabolic oscillations, metabolic modelling.

Introduction

Cord-forming basidiomycete fungi form extensive mycelial networks that scavenge inorganic and organic nitrogen (N) efficiently from the soil environment (Boddy & Watkinson, 1995; Dighton, 1997). Their main source of carbon (C) for respiration or provision of C-skeletons for metabolism is wood, which has very low levels of N and other nutrients, such as phosphate (Boddy & Watkinson, 1995; Dighton, 1997). Thus considerable fluxes of N and phosphorus (P) must occur from remote assimilation sites or following autolysis of redundant mycelium to regions of high demand that arise during colonization and exploitation of new wood resource (Levi & Cowling, 1969; Venables & Watkinson, 1989; Cairney, 1992; Jennings, 1994; Boddy, 1999; Olsson, 2001). Resources are often scarce, ephemeral and patchy. As a result, the mycelial network develops as a highly plastic, interconnected functional unit that continuously senses and responds to nutritional cues in the environment and translates them into developmental responses (Rayner et al., 1995; Boddy, 1999; Ritz & Crawford, 1999; Olsson, 2001).

Most experimental data on nutrient transport in cord-forming mycelia is based on movement of radiolabelled phosphate (Cairney, 1992; Jennings, 1994; Boddy, 1999). There are fewer reports on N-translocation as there is no readily available radioactive N-isotope. Heavy nitrogen (15N) and mass spectroscopy have been used to study transport (Arnebrant et al., 1993), while stable N isotopes have proved useful in characterizing the metabolic pathways involved in mycorrhizal N metabolism using nuclear magnetic resonance (NMR) (Pfeffer et al., 2001), but have not yet been applied in imaging mode to resolve N transport. An alternative strategy has been to use a 14C-labelled amino-acid analogue, \( \alpha \)-aminoisobutyric acid, (AIB) which is taken up but not
metabolised (Ogilvie-Villa et al., 1981; Kim & Roon, 1982; Watkinson, 1984; Lilly et al., 1990; Olsson & Gray, 1998). Because AIB is not metabolized, the $^{14}$C-label faithfully reports the distribution of the amino acid analogue. Although early studies on $^{14}$C-AIB translocation used destructive sampling, dynamics of $^{14}$C-AIB movement were recently imaged in intact colonies of Schizophyllum commune using a $\beta$-scanner with spatial resolution in the order of a few millimetres and a theoretical sampling interval of about 1 h (Olsson & Gray, 1998).

To visualize N transport in mycelial networks with higher temporal and spatial resolution, we developed a novel noninvasive technique to track movement of $^{14}$C-AIB in mycelia grown over an inert scintillation screen using a photon-counting camera (Tlalka et al., 2002). Rapid, pulsatile movement of AIB was observed with a period of 14.5 h in growing, foraging hyphae and 11–12 h along developing cords. The oscillations were maintained for a considerable period in darkness at constant temperature.

Rhythmic phenomena are widespread in fungi and other simple eukaryotes, ranging from self-sustaining metabolic rhythms with periodicities of the order of minutes (Kippert & Hunt, 2000) to the archetypal circadian rhythms associated with sporulation in Neurospora (Dunlap, 1998; Ramsdale, 1999; Bell-Pedersen, 2000; Merrow et al., 2001). Circadian rhythms in amino-acid uptake have been described for yeast (Edmunds et al., 1979) and Synechococcus (Chen et al., 1991). While the period of the oscillations in $^{14}$C-AIB transport observed previously was clearly different from a canonical 24-h circadian rhythm, it is possible that the oscillations are tied into the output of an ultradian oscillator with a period of less than 24 hours (Dunlap, 1998; Kippert & Hunt, 2000).

In this paper we report on improvements in both the camera system and the scintillation screen that allow simultaneous measurement of signals from both assimilatory hyphae that grow on the agar inoculum and takeup nutrients from it, and distal foraging hyphae that extend over the surrounding nonnutrient area. With this more sensitive system we have tested: first, whether there are differences in the pulsatile behaviour within assimilatory and foraging hyphae; second whether the oscillations are coupled to pulses in growth; and third whether the rhythm observed is subservient to a central clock, and therefore reflects the output of that oscillator, or whether the oscillations are an intrinsic part of amino-acid uptake and translocation and may therefore yield information on the underlying control systems.

Materials and Methods

Organism

Cultures of Phanerochaete velutina were originally provided by Prof. L. Boddy, University of Cardiff, UK, and have been maintained in the Department of Plant Sciences, Oxford, UK, for 3 yr. Cultures were grown on 2% malt agar (2% malt extract, Oxoid Ltd, Basingstoke, UK, 2% Oxoid No. 3 agar) at 22 ± 1°C in darkness in a temperature-controlled incubator (Gallenkamp, Loughborough, UK) as previously described (Tlalka et al., 2002).

Experimental microcosms

Small artificial microcosms were prepared as in Tlalka et al. (2002). Briefly, a submarginally cut inoculum of P. velutina was placed mycelial surface down in the middle of a square Petri dish on top of either a Lite Plus intensifying screen (Sigma, Poole, UK) or a BioMax TranScreen LE intensifying screen (Sigma). Humidity was maintained by the presence of water in small (500 µl) containers and by sealing the sample dishes with Parafilm (American National Can, Neenah, WI, USA). All experiments were conducted in a temperature-controlled room, usually at 21 ± 0.5°C. During experiments investigating the temperature-dependence of the oscillations the temperature was set at different values between 19°C and 26°C, as indicated in figure legends.

Visualization of $^{14}$C-AIB transport using photon-counting scintillation imaging

Imaging of $^{14}$C-AIB was achieved using a more sensitive version of the high-resolution, photon-counting camera system (HRPCS-3; Photek Inc., St Leonards on Sea, UK) described previously (Tlalka et al., 2002). In the modified version, the three-stage microchannel-plate intensifier was equipped with a larger face-plate (40 mm diagonal) and a low-noise S20 photocathode. The larger format camera also allowed the use of higher numerical aperture photographic lenses. Data reported here used a Nikon 28 mm f/2 lens which gave a nominal (x,y) pixel dimensions of 588 µm and a field of view of $433 \times 334$ mm. This configuration, allowed for better reproducibility than before, because six to nine replicate experiments could be conducted in parallel with similar overall signal levels from each colony to those reported previously.

The increase in sensitivity also allowed a reduction in the amount of $^{14}$C-AIB to half of that used previously. Immediately after placing the inocula on the scintillation screens, 25 µl (46.3 kBq) of a 0.9-mm solution of 2-amino[1-$^{14}$C]isobutyric acid, $^{14}$C-AIB (Amersham, Chalfont St Giles, UK) in distilled water (specific activity 2.11 GBq mmol$^{-1}$) was applied to the centre of the inoculum. The chambers were then sealed and placed in continuous darkness in the camera imaging box in a temperature-controlled room. Most experiments were conducted at 21 ± 0.5°C unless otherwise stated in figure legends. The temperature within the box was continuously monitored with ‘Diligence’ data loggers (Comark Ltd, Stevenage, UK). Images were integrated over 30-min periods and experiments lasted for up to 480 h.

The signal from $^{14}$C-AIB in assimilatory hyphae growing on the inoculum disc and foraging hyphae growing out over
the scintillation screen surface were measured simultaneously in a number of selected circular regions of interest (ROIs) of identical size (one centred on the inoculum and the others on the outwardly growing foraging hyphae). The frequency of pulses from each region was determined by Fourier analysis, essentially according to Tlalka et al. (2002). Briefly, total counts over 30-min periods from each region were outputted to Excel (Microsoft Corp.) and smoothed with a rolling 3- to 9-point averaging filter. Longer-term trends were eliminated by taking the second difference of each sequence (Diggle, 1990). Visual inspection indicated that this provided a reasonably stationary time-series suitable for Fourier analysis. Each series was padded with zeros to give a total of 512–2048 values to increase the resolution of the Fourier spectrum (Smith, 1997). The period of the pulses was calculated from the Fourier frequency with the greatest amplitude and the degree of synchrony assessed from the phase difference of this frequency between each region and the inoculum.

Calibration of the scintillation imaging system

To determine the sensitivity of the new camera system in combination with the new BioMax TranScreen LE intensifying screen, images were collected from a series of 14C-AIB standards containing varying amounts of 14C-AIB from 0 to 18.5 kBq in 20 µl droplets that had been dried on to the screen. Values of 160 photons min⁻¹ nmol⁻¹ 14C-AIB were measured for the Lite plus screen and 1000 photons min⁻¹ nmol⁻¹ 14C-AIB for the BioMax TranScreen LE. This compares with around 500 photons min⁻¹ nmol⁻¹ 14C-AIB for our previous system using the Lite Plus screen (Tlalka et al., 2002), but with a useable imaging area eight- to nine-fold greater than available previously. No oscillations were observed for dried-down calibration droplets imaged over 300–400 h. The increased sensitivity of the new system produced substantial file sizes during long time courses, reaching in excess of 1 GB.

To determine the level of signal from 14C-AIB-loaded inoculum discs without mycelium, the signal from 25 µl of 14C-AIB (46.3 kBq) was recorded from labelled agar plugs placed directly on the two different intensifying screens. The signal from the plug with mycelium was at least sevenfold higher than that simply attributable to an equivalent amount of 14C-AIB allowed to equilibrate throughout an agar plug in the absence of mycelium (0.4 photons min⁻¹ µm⁻² with mycelium compared to 0.06 photons min⁻¹ µm⁻² without).

To determine the extent that the agar inoculum attenuated the signal from the hyphae growing underneath, uncolonized agar disks were randomly placed on areas of previously labelled foraging hyphae to mimic the optical path encountered by photons from 14C-AIB accumulated by the assimilatory hyphae in the inoculum. Signal losses of 94% (Lite Plus) or 87.5–90% (Biomax TranScreen LE) were calculated from the ratio of 14C-AIB signals from the prelabelled foraging hyphae in the presence and absence of the agar plug, after subtraction of the appropriate background.

Measurement of colony growth

The growth of colonies on the two types of intensifying screens was estimated following median filtering of the scintillation images using a 3 × 3 kernel and intensity-based thresholding to produce a segmented, binary image. The darker region corresponding to the central inoculum was included in the binary image using a hole-fill operation. Radial growth was estimated as half the maximum diameter across the segmented image following subtraction of the inoculum diameter.

In addition, the growth of P. velutina on the two different screens was measured under low level white-light illumination (12 W energy-saver; Osram, München, Germany) using time-lapse video-imaging at a photon flux of 150 µmol m⁻² s⁻¹ measured with a light power meter (model 1815-C; Newport Corp., Irvine, CA, USA). Colonies were grown under the same conditions as for the scintillation imaging except that the inoculum was loaded with either 25 µl of 0.9 mCi cold AIB to match the concentration of 14C-AIB, or 25 µl distilled water. Images of the growing hyphae from four replicate experiments were captured simultaneously using monochrome cameras (Cohu 4910; Brian Reece Scientific Ltd, Newburg, UK) equipped with macro zoom lenses (18–108 mm, f/2.5; Edmund Scientific, York, UK) adjusted to give field sizes of between 19 and 110 mm diagonal. Signals were digitized using a Matrox Meteor 2 framestore controlled by KCS300 software (Karl Zeiss Vision GmbH, Jena, Germany). Brightness and contrast levels were manually adjusted to achieve contrast from the fine, translucent hyphae growing across the white screens. Typically, 50 frames were averaged over 5 s to reduce noise and images collected at 30-min intervals over 250 h. Images were imported into Lucida v4.0 (Kinetic Imaging Ltd, Liverpool) and median filtered using a 3 × 3 kernel to reduce noise. The contrast for growing hyphal tips was accentuated using a high-pass filter. The linearity of growth was visualized by sampling line-transects drawn parallel to the growth direction and writing the resultant intensity plot into successive lines of a second image representing intensity along the x-axis and time on the y-axis. The rate of growth was measured as the slope of the colony profile and the uniformity of growth estimated by the linearity of the profile.

Simulation modelling of amino acid uptake and transport

Several simulation models were set up to explore the conditions under which a typical complement of amino acid transport systems and intracellular compartments might generate the observed kinetics for 14C-AIB using MODELMAKER (Cherwell Scientific Publishing, Oxford, UK). Uptake of
Materials and Methods).

absorption and/or scattering by the agar plug (see the was about tenfold greater, but was attenuated through (Fig. 1b). We estimate that the true signal from these hyphae particularly for colonies grown on the BioMax T ranScreen LE

The 14C-AIB signal was clearly visible in emissions excited by β-particles colliding with the underlying scintillation screen. The 14C-AIB from the agar by assimilatory hyphae in the inoculum was assumed to operate via a saturable transport system obeying Michaelis–Menten kinetics with a Km of 100 µM (Ogilvie-Villa et al., 1981; Kim & Roon, 1982). In simulations that included a vacuolar compartment, vacuolar uptake was assumed to operate via a saturable carrier with a Km of 400 µM (Zerez et al., 1986). In addition, a vacuolar efflux system with first-order kinetics was introduced in parallel on the tonoplast that was activated once a threshold level (10 mM) of amino acid had accumulated in the vacuole. The efflux system was inactivated once the vacuolar content had fallen below a second, lower threshold set at 60% of the maximum pool size. The 14C-AIB was transferred to the foraging hyphae at a rate proportional to the growth-rate, to reflect the increasing volume of the mycelium. With these constraints, the only additional variables were the apparent Vmax for the transporters and the rate constant for the tonoplast efflux carrier. The sensitivity of the system across a range of values for each of these variables was explored. The series of interlinked differential equations were solved numerically using the fourth order Runge–Kutta method within MODELMAKER.

Movie and image presentation
Movie files were output directly from the IFS32 Imaging Software (Photek) or assembled from individual images using QuickTime Pro 4.0 (Apple Computer Inc.). Images for publication were assembled in PhotoShop 4.0 (Adobe Systems, San Jose, CA, USA).

Results
Oscillations in 14C-AIB transport show complementary profiles in assimilatory and foraging hyphae

The 14C-AIB was taken up by hyphae of P. velutina when applied to the inoculum plug and was imaged from photon profiles in assimilatory and foraging hyphae

Fourier analysis confirmed that the dominant frequency was the same for both sets of oscillations and was similar between colonies growing on the two types of screen (Fig. 1g,h). The corresponding period of the oscillations at 21°C was around 16 h (Table 1).

Oscillations are asymmetric
Closer inspection of the oscillations revealed that they were not sinusoidal, but showed a pronounced asymmetry that was particularly apparent once the plateau phase was reached in colonies grown over the Biomax TranScreen LE (Fig. 2). In the foraging hyphae, there was a slow rise lasting around 11 h during the first phase (P1) of the cycle, followed by an abrupt switch to rapid decay lasting about 5 h in the second phase (P2) of the cycle (Fig. 2 and Table 1). The opposite pattern was observed for the assimilatory hyphae, starting with a slow, exponential-like drop and followed by a rapid recovery. This asymmetry was reflected in the Fourier analysis that showed additional peaks in the spectrum, notably at the first harmonic (Fig. 1g,h).

Oscillations are not tightly linked to an internal clock
Colonies were exposed to a short (1 h) light break before transfer to continuous darkness for the imaging experiments. There was no evidence that this pretreatment was sufficient to entrain the subsequent oscillations as pulses recorded from mycelia in replicate experiments conducted simultaneously in the same chamber were not synchronized (Fig. 3a). The lack of synchrony also rules out the possibility that oscillations were associated with cyclical changes in environmental conditions or instrument artefacts.
To investigate the temperature dependence of oscillations further in this study, the period was measured using Fourier analysis from foraging hyphae grown at a range of different temperatures between 19°C and 25.5°C. The period decreased markedly with increasing temperature (Fig. 3b). The temperature coefficient ($Q_{10}$) was calculated as 2.1 by extrapolation from the ratio of the frequency at a given temperature to the frequency at a temperature 10°C higher (Fig. 3b, inset). In several of these experiments, colonies were also shifted between temperatures. The period of the oscillations adjusted to the temperature within a few cycles but there was no evidence that the phase of the oscillation was reset by the temperature jump (data not shown). The timing of the temperature jump differed with respect to the phase of the cycle in parallel colonies, but, in the absence of clear support for an underlying circadian or ultradian rhythm, we have not systematically...
tested whether temperature jumps or light breaks could entrain the rhythm at defined phases of the cycle.

Radial growth of the colony does not have a clear pulsatile component

To test whether the pulses of $^{14}$C-AIB detectable in the foraging hyphae were associated with periodic changes in growth rate or growth habit, the colony diameter was estimated following intensity-based thresholding of the scintillation images. On both screens, growth was relatively slow for the first 30–40 h then increased to a maximum around 70–100 h before tailing off again by 200 h (e.g. Fig. 4a). The apparent growth, determined as the maximum colony diameter, was greater for colonies grown on the Biomax TranScreen compared with the Lite Plus (Table 2). However, these figures are slightly misleading because part of the apparent difference in growth arises from stochastic variations in colony morphology. Radially symmetrical colonies (Fig. 4b) had a lower diameter than those where symmetry was broken (e.g. Fig. 4c). These measurements of growth can provide only a crude descriptor of the true colony behaviour.

No obvious oscillations or pulsatile surges in growth were conducted in darkness, we first ascertained that growth of $P$. velutina (on agar) was not markedly affected by light intensity up to 150 µmol m$^{-2}$ s$^{-1}$ – the highest intensity tested (Table 2) – and showed little or no dependence on wavelength (data not shown). Over the first 70–100 h growth was mainly in the form of relatively infrequently branched thin hyphae growing on or near the surface of the screen. There was no detectable pulsatile component for growth during this period whether colonies were grown on agar or on scintillation screens and in the presence or absence of cold AIB (Fig. 4d–f and Table 2). From this point forward, colony growth became progressively more complex. Radial expansion continued at roughly the same rate but branching became more frequent, some hyphae appeared to regress and the initial radial symmetry altered in favour of a more limited number of discrete point growth foci (compare Fig. 4b,c) typical of the early stages in development of a cored system. There was no obvious synchronization of these events throughout the colony with a
periodicity that might be associated with oscillations in $^{14}$C-AIB transport. However, we cannot rule out subtle changes in growth pattern due to the technical difficulties of imaging morphological changes within the mass of thin, white hyphae superimposed on the white scintillation screen. For example, we might not detect cycles where a limited number of unsuccessful foraging hyphae regress and their contents are remobilized to support growth at the margin.

Discussion

$^{14}$C-AIB is taken up rapidly by the assimilatory hyphae in the inoculum

The improved sensitivity of the camera system used in this study compared with our previous system (Tlalka et al., 2002) enabled simultaneous measurement of $^{14}$C-AIB dynamics for

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Table 2 Growth of *Phanerochaete velutina* on different substrates

<table>
<thead>
<tr>
<th>Substrate</th>
<th>AIB</th>
<th>Light</th>
<th>Measurement technique</th>
<th>Maximum growth rate$^2$ ($\mu$m h$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agar</td>
<td>-</td>
<td>-</td>
<td>Direct measurement</td>
<td>238 ± 14 ($n = 12$)</td>
</tr>
<tr>
<td>Agar</td>
<td>-</td>
<td>+</td>
<td>Direct measurement</td>
<td>211 ± 17 ($n = 12$)</td>
</tr>
<tr>
<td>BioMax TranScreen</td>
<td>+</td>
<td>-</td>
<td>Scintillation imaging</td>
<td>317 ± 66 ($n = 6$)</td>
</tr>
<tr>
<td>Lite Plus</td>
<td>+</td>
<td>-</td>
<td>Scintillation imaging</td>
<td>186 ± 72 ($n = 12$)</td>
</tr>
<tr>
<td>Lite Plus</td>
<td>+</td>
<td>+</td>
<td>Bright-field image analysis</td>
<td>157 ± 39 ($n = 4$)</td>
</tr>
<tr>
<td>Lite Plus</td>
<td>-</td>
<td>+</td>
<td>Bright-field image analysis</td>
<td>168 ± 12 ($n = 4$)</td>
</tr>
</tbody>
</table>

$^1$AIB, $\alpha$-aminoisobutyrate. $^2$Values are given as the mean ± SD, with the number of separate experiments given in brackets.
Asymmetric oscillations highlight physiological differences between assimilatory and foraging hyphae

Following uptake, $^{14}$C-AIB was distributed to the foraging hyphae growing out from the inoculum. Asymmetric oscillations were observed for both the assimilatory and foraging hyphae with complementary profiles. It is difficult to infer the underlying mechanisms that would generate such asymmetrical pulses, particularly the sharp transitions between the two apparently distinct phases of the cycle. We can exclude a tight coupling to an ultradian/circadian clock as, although the oscillations persisted in constant conditions, they showed a marked temperature dependence. Similarly, we have not been able to detect synchronized changes in marginal growth that might drive cyclical changes in flux, although we cannot rule out more subtle cycles of regression and redistribution within the foraging hyphae. Even if such morphological changes did occur, it is not immediately apparent how the complementary profile of the rhythm in the assimilatory and foraging hyphae and the distinct switch in behaviour during the cycle would be generated. Previously (Tlalka et al., 2002) we suggested that cyclical changes in the fungal structure or local distribution of $^{14}$C-AIB might alter the efficiency with which photons were either generated or detected and thus generate artefactual pulses. This explanation now seems less likely, unless the underlying process also operates asynchronously in the assimilatory and foraging hyphae. We have therefore focused on physiological and biochemical explanations of the pulsing behaviour.

Oscillatory behaviour can be simulated with a pool-refilling model

To link the in vivo observations in this study with existing biochemical information, we developed a simulation model that included saturable AIB uptake across the plasma membrane, followed by distribution throughout the mycelium by a mass-flow mechanism (Fig. 5a). Because there is also extensive evidence that fungi accumulate amino acids in their vacuoles under conditions of high N-availability (Kitamoto et al., 1988; Watkinson, 1999), vacuolar transport was also included in the simulation. Although vacuolar uptake may be mediated by both high-affinity (Zerez et al., 1986) and low-affinity (Keenan & Weiss, 1997) tonoplast amino-acid transporters (Klionsky et al., 1990; Horák, 1997; Wipf et al., 2002), only a single saturable tonoplast transport system was included in the model. The kinetic parameters for the transporters and the constants used in the simulation are given in Fig. 5b.

Under conditions of N demand, stored vacuolar amino acids are mobilized and there is growing biochemical and molecular evidence for the involvement of a specific set of tonoplast efflux transporters (Keenan & Weiss, 1997; Russnak et al., 2001). Molecular insights into control of the switch between accumulation and release are also beginning to emerge, particularly from studies on selective permeabilization of the plasma membrane to allow direct access to the tonoplast (Keenan & Weiss, 1997; Roos et al., 1997). While vacuolar amino acid uptake appears to be independent of the intravacuolar concentration, efflux is initiated immediately vacuolar amino acid levels exceed a threshold value (Keenan & Weiss, 1997; Roos et al., 1997). The consequences of such a system operating were explored in the simulation. In permeabilized cell models, efflux is also stimulated by changes in the cytoplasmic nucleotide pools (Roos et al., 1997; Steighardt et al., 2000), but this level of control has not yet been incorporated into the model.

In the simulation, the plasma membrane uptake step alone, in the absence of vacuolar transport, was not capable of generating oscillations, even if feedback-inhibition by increasing cytoplasmic concentrations was included in the model (Fig. 5c). Similarly, inclusion of a vacuolar compartment yielded a greater net accumulation in the assimilatory hyphae, but did

hyphae performing different physiological roles, namely nutrient uptake by assimilatory hyphae in the inoculum and exploratory growth by the foraging hyphae. The assimilatory hyphae growing under the inoculum plug accumulated $^{14}$C-AIB avidly from a relatively low concentration (22 µM), presumably through the action of a plasma membrane amino acid transporter(s). In fungi, including saprotrophic basidiomycetes (e.g. Kersten et al., 1999), a range of amino acid uptake systems exist in parallel on the plasma membrane, including high-affinity transporters for specific amino acids with $K_m$ values in the 1–1000 µM range, and more general amino acid permeases with slightly lower affinities (Horák, 1997; Chalot & Brun, 1998). These transporters share homology with members of amino acid transporter families described for animals and plants (Wipf et al., 2002). Although AIB most closely resembles the neutral amino acid alanine in structure, uptake experiments suggest that it may be transported by a general amino acid permease in both Neurospora (Olgilvie-Villa et al. 1981) and Sacharomyces (Kim & Roon, 1982) with $K_m$ values of 98.7 µM and 270 µM, respectively. The situation in basidiomycetes is not yet resolved with the limited data available from competition studies. Thus, AIB uptake is inhibited by high levels of glutamate in Serpula lacrymans (Watkinson, 1984) or t-asparagine in Schizophyllum commune (Lilly et al., 1990). These amino acids are likely to be taken up both by a general amino acid permease and by more specific transporters. Apparently nonsaturable uptake systems may also contribute to assimilation of some amino acids (Chalot & Brun, 1998; Anderson et al., 2001), although these typically operate at higher substrate concentrations than those used here. The absence of a detectable lag period in uptake (Fig. 1c,d) suggests that the transporter(s) was already present here. The absence of a detectable lag period in uptake (Fig. 1c,d) suggests that the transporter(s) was already present in the membrane rather than requiring induction and de novo expression.

not give rise to oscillations (data not shown). However, the type of hysteresis required to generate a continuous train of pulses was observed if a threshold-dependent transition between uptake and release from the vacuolar storage pool was included (Fig. 5d). In this simulation, uptake into the vacuolar storage pool was allowed to proceed through a saturable carrier with a $K_m$ of 400 µM (Zerez et al., 1986) and efflux was via a system with first-order kinetics, which was activated once a threshold level of amino acid accumulated in the vacuole. The efflux system was inactivated once the vacuolar content fell below a second, lower threshold, set at 60% of the maximum pool size. A range of values for each of the variables were explored (the results shown in Fig. 5d are illustrative). This combination of cyclical uptake and release from the vacuole plus continuous removal to the foraging hyphae was sufficient to simulate several of the features observed in vivo, including the rapid increase in signal in the assimilatory hyphae, the delayed increase in signal from the foraging hyphae and the superimposed complementary asymmetric oscillations.

It is encouraging that a relatively simplistic model using generic components provides a reasonable description of the pulsatile behaviour in an intact system. If validated by further work, it suggests that the combination of in vivo analysis and simulation modelling might prove a useful route to understand the control of fluxes between the cytoplasmic and vacuolar amino acid pools operating in vivo. Equally, we cannot rule out, at this stage, that pulses are generated by a more complex series of control systems involving a greater number of compartments and/or differentiated hyphae.

Fig. 5 Simulation for generating oscillations based on a vacuolar pool refilling model. The $^{14}$C-labelled $\alpha$-aminoisobutyrate ($^{14}$C-AIB) dynamics were modelled as the interaction between fluxes across the plasma membrane and tonoplast in the assimilatory hyphae and transport to the foraging hyphae (a). Uptake across the plasma membrane and tonoplast were modelled using Michaelis–Menten kinetics, using apparent $K_m$ values drawn from the literature (b). Efflux from the vacuole was modelled with first-order kinetics that were triggered once the intravacuolar concentration reached a threshold level. Efflux was shut down once the intravacuolar concentration had fallen to 60% of its maximum value. Translocation to the foraging hyphae was modelled at a rate proportional to the growth of the mycelium measured experimentally and the cytoplasmic concentration. Superscripts refer to the compartment as follows: c, cytoplasm; a, agar; v, vacuole; the direction of the arrow indicates the direction of the flux. The output of the model for similar sized areas of assimilatory and foraging mycelium is shown in (c) for the case with no vacuolar transport, and in (d) for the full model. The combination of cyclical uptake and release from the vacuole and continuous removal to the foraging hyphae was sufficient to simulate several of the features observed in the in vivo imaging experiments.
Are the observed oscillations unique to rapid 
¹⁴C-AIB exposure?

While such simulations may highlight the importance of coordination between transport and vacuolar storage in amino acid homeostasis, a number of important caveats remain. First, it might be expected that if the sudden uptake of readily-available amino acid exceeds throughput to the rest of the mycelium, the system will be prone to oscillate (Rayner et al., 1995). The oscillations would then map the trajectory of the displaced system as it returns to its unperturbed state rather than highlight an intrinsic regulatory feature of N uptake and translocation in this fungus.

It should be noted that AIB is also not without biological effect. The extent that AIB interferes with normal N-sensing and metabolism is not well established. At high (mM) concentrations, AIB suppresses exploratory growth, tricking the fungus into an assimilatory growth pattern by inhibiting cord development and hyphal extension and increasing branch frequency (Elliot & Watkinson, 1989), even though AIB cannot be metabolized (Watkinson, 1984). It is possible that AIB may directly interfere with external N-sensing systems at the plasma membrane, such as a homologue of amino acid sensing permease, Ssy1p (reviewed by Gagiano et al., 2002), or disturb intracellular N sensing by competing for transporters or energy gradients involved in maintaining normal amino acid pool sizes. If this were to impact on the level of glutamine, for example, it might perturb intracellular N sensing through interactions of uncharged glutamine tRNA_{CUG} molecules with a kinase cascade (reviewed by Gagiano et al., 2002). Alternatively, because AIB is not metabolized, it may not be recognized by the normal regulatory control mechanisms, and consistently overshoot the normal homeostatic set points.

Evidence from yeast indicates that AIB may act as an effector of N catabolite repression in the presence of poor nitrogen sources, but not with good N sources (Kim & Roon, 1982). Whether this is a specific effect is less clear, as the concentrations used (1–10 mM) were also sufficient to inhibit protein synthesis (Kim & Roon, 1982). The external concentration of AIB used here (22 µM) was significantly lower than this and did not affect the growth of 

\textit{Phanerochaete} in this study. In our previous report (Tlalka et al., 2002), we attributed the reduction in growth observed after several days to the presence of AIB. It now seems more likely that the major effect on growth was the higher temperatures (up to 26°C) that arose during the imaging experiments with the previous camera system.

Any inhibitory effects of AIB may also be masked by the high levels of other amino-acids in the relatively rich malt agar inoculum used. It will be interesting in future work to bring the overall AIB : N ratio under experimental control to see if it is possible to directly correlate changes in growth pattern with the intracellular N status throughout the entire, intact foraging mycelial network.

In summary, oscillations in 
¹⁴C-AIB uptake and transport were not associated with a circadian or ultradian rhythm in 

\textit{P. velutina}, but showed a marked temperature-dependence consistent with a self-sustaining metabolic rhythm. We infer from the asymmetric shape of the pulses, particularly the abrupt transition between each exponential decay phase and the next rising phase, that a switch-like phenomenon takes place in the underlying uptake and transport systems. A model that included known properties of transporters in other fungal systems and sequestration/release from the vacuole simulated several aspects of the observed behaviour, including the complementary profiles of the pulses in the assimilatory and foraging hyphae. This model focuses attention on the possible role of vacuolar uptake and release of amino acids as a regulatory step in the pathway of N from outside the cell into intracellular metabolism, and may, in future, help us in the investigation of the morphogenetic effect of nutrients that is such a striking feature of fungal development. It is well established that facultative development of multihyphal aggregated structures responds to external nutrient triggers and available evidence (reviewed by Watkinson, 1999) implicates intracellular amino acid levels in the signalling pathway.

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