Dynein and dynactin deficiencies affect the formation and function of the Spitzenkörper and distort hyphal morphogenesis of *Neurospora crassa*

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The impact of mutations affecting microtubule-associated motor proteins on the morphology and cytology of hyphae of *Neurospora crassa* was studied. Two ropy mutants, ro-1 and ro-3, deficient in dynein and dynactin, respectively, were examined by video-enhanced phase-contrast microscopy and image analysis. In contrast to the regular, hyphoid morphology of wild-type hyphae, the hyphae of the ropy mutants exhibited a great variety of distorted, non-hyphoid morphologies. The ropy hyphae were slow-growing and manifested frequent loss of growth directionality. Cytoplasmic appearance, including organelle distribution and movement, were ostensibly different in the ropy hyphae. The Spitzenkörper (Spk) of wild-type hyphae was readily seen by phase-contrast optics; the Spk of both ro-1 and ro-3 was less prominent and sometimes undetectable. Only the fast-growing ropy hyphae displayed a Spk, and it was smaller and less phase-dark than the wild-type Spk. Growth rate in both wild-type and ropy mutants was directly correlated with the size of the Spk. Spk efficiency, measured in terms of cell area generated per Spk travelled distance, was lower in ropy mutants. Another salient difference between ropy mutants and wild-type hyphae was in Spk trajectory. Whereas the Spk of wild-type hyphae maintained a trajectory close to the cell growth axis, the Spk of ropy hyphae moved much more erratically. Sustained departures in the trajectory of the ropy Spk produced corresponding distortions in hyphal morphology. A causal correlation between Spk trajectory and cell shape was tested with the Fungus Simulator program. The characteristic morphologies of wild-type or ropy hyphae were reproduced by the Fungus Simulator, whose vesicle supply centre (VSC) was programmed to follow the corresponding Spk trajectories. This is evidence that the Spk controls hyphal morphology by operating as a VSC. These findings on dynein or dynactin deficiency support the notion that the microtubular cytoskeleton plays a major role in the formation and positioning of the Spk, with dramatic consequences on hyphal growth and morphogenesis.

**Keywords:** ropy mutants, Spitzenkörper, *Neurospora crassa*

**INTRODUCTION**

This study is part of a project to elucidate the role of the Spitzenkörper (Spk) in hyphal growth and morphogenesis of higher fungi. In the present work, we used video-enhanced phase-contrast microscopy and image analysis to examine the behaviour of morphological mutants of *Neurospora crassa* affected in genes encoding microtubule-associated motor proteins.

The Spk is a dynamic body whose structure varies widely among fungi; it usually consists of an outer vesicle cloud and an inner core (López-Franco & Bracker, 1996). In addition to vesicles (Girbardt, 1969; Grove & Bracker, 1970), other components have

**Abbreviations:** Spk, Spitzenkörper; VSC, vesicle supply centre.
been detected in the Spk, including amorphous or granular material of undefined nature in the core region as well as components of the cytoskeleton and ribosomes (McClure et al., 1968; Turian, 1978; Bourett & Howard, 1991; Roberson & Vargas, 1994; López-Franco & Brack, 1996). The morphology of the fungal cells is determined by the way the cell wall is assembled (Barnticki-García, 1968, 1973). In hyphae, cell wall growth occurs mainly at the tip by polarized secretion of enzymes and cell wall precursors (Barnticki-García & Lippman, 1969; Trinci, 1978; Harold, 1990). There is a sizeable body of evidence that the Spk plays a central role in apical growth and morphogenesis (Girbardt, 1957; López-Franco & Brack, 1996; Reynaga-Peña & Barnticki-García, 1997). According to the vesicle supply centre (VSC) model for fungal morphogenesis (Barnticki-García et al., 1989), the Spk functions as a vesicle distribution centre. Vesicles generated in distal parts of the hypha congregate in the Spk and from there migrate to the cell surface. The linear displacement of the Spk creates a sharp gradient of exocytosis responsible for hyphal morphogenesis.

There is mounting evidence that the position of the Spk governs the growth direction of a hypha (Girbardt, 1957; Brack, 1997; Riquelme et al., 1998). Our previous work with microtubule inhibitors implicated the microtubular cytoskeleton in the positioning and movement of the Spk in hyphae of N. crassa (Riquelme et al., 1998), but its exact role in apical growth has yet to be elucidated. It has long been proposed that cytoplasmic microtubules participate in the transport of secretory vesicles to the hyphal apex (Howard & Aist, 1977, 1980; Howard, 1981; Gooday, 1983; Gow, 1989; Hasek & Barnticki-García, 1994; McKerracher & Heath, 1987; Heath, 1994). The discovery of microtubule-associated motor proteins (Hirokawa, 1982; Paschal et al., 1987) has helped us understand how organelles move inside the cell. Cytoplasmic dyneins and members of the kinesin superfamily are main motor enzymes involved in vesicle translocation along microtubules (Haimo & Thaler, 1994; Hirokawa, 1998). There is now growing evidence that both kinesins and cytoplasmic dyneins are involved in the traffic of secretory vesicles in fungal hyphae (Seiler et al., 1997; Wu et al., 1998; Inoue et al., 1998). Cytoplasmic dyneins are multisubunit enzymes involved in transport of membranous organelles towards theminus end of microtubules (Paschal et al., 1987; Schroer & Sheetz, 1991; Hirokawa, 1998). Kinesins are motor proteins involved in membrane transport towards the plus end of the microtubules (Vale et al., 1985; Hirokawa, 1982; Steinberg & Schliwa, 1996).

To examine the relationship among the Spk, the microtubular cytoskeleton and hyphal morphogenesis, we chose two roppy mutants of N. crassa, ro-1 and ro-3. Both belong to the group of true colonial morphological mutants (Garnjobst & Tatum, 1967; Vierula, 1996) and have been characterized at the molecular level. Mutant ro-1 is deficient in one of the heavy chains of cytoplasmic dynein (Plamann et al., 1994). Mutant ro-3 is deficient in p150Glued, the largest subunit of the dynactin (dynein activator) complex (Plamann et al., 1994; Tinsley et al., 1996). Dynactin is a multisubunit complex required for cytoplasmic dynein to efficiently transport vesicles along microtubules in vitro (Gill et al., 1991; Schroer & Sheetz, 1991).

Most studies on roppy mutants have primarily focused on the aberrant distribution of nuclei in ro-1 and ro-3 hyphae and on the molecular characterization of those loci (Plamann et al., 1994; Bruno et al., 1996; Tinsley et al., 1996). Since hyphal and, ultimately, colony morphology are largely established at the hyphal apex, we have focused this study on the impact of the roppy mutations on apical events, where most of the growth process is concentrated.

**METHODS**

**Strains and media.** N. crassa wild-type (FGSC 988) and roppy strains (FGSC 4351 and 43 for ro-1 and ro-3 respectively) were obtained from the Fungal Genetics Stock Center. The strains were grown and maintained in 8.5 cm plastic Petri dishes (sold as 100 x 15 mm, Fischer Scientific) containing 20 ml Vogel's Complete Medium (VCM) agar (Vogel, 1956) with 1.5% (w/v) sucrose as the carbon source. Cultures were grown at 21 °C.

**Video microscopy.** Growing hyphae were observed with an Olympus Vanox-S microscope. For low-magnification images, the fungus was grown in 8.5 cm plastic Petri dishes on a thin layer (10 ml) of VCM agar at 21 °C and observed with bright-field optics (10 x objective and 10 x WF eyepiece). For high-resolution work, the fungus was grown on a modified slide culture chamber (López-Franco, 1992; Riquelme et al., 1998) and observed with a phase-contrast 100 x oil-immersion objective (n.a. 1.25) and a 25 x WF eyepiece (American Optical).

Video images were produced with a Hamamatsu C2400-07 high-resolution camera (Hamamatsu Photonic Systems), enhanced with an Argus-10 image processor (a real-time digital contrast and low light enhancement system), and displayed on a black and white, 12-inch, high-resolution monitor (Sony; model PVM-122). Sequences were videotaped in real time with an S-VHS recorder (JVC model BR-S822U).

**Growth rate and cell parameters measurements.** Videotaped sequences were played on a variable-tracking player (JVC model BR-S525SU) and observed on a Sony Trinitron model monitor. Individual images were captured from the videotaped sequences in 8-bit greyscale with an Imagase/Chroma frame grabber (Imagraph). With Image Pro Plus Software for Windows (Media Cybernetics), we traced the cell profiles of the images captured by video microscopy. xy coordinate values were automatically collected into a text file with a Windows application program interfaced with the Argus-10 analyser (Bartnicki et al., 1994). The text files were then imported into Microsoft Excel spreadsheets and analysed.

Growth rate was measured in terms of cell area increase. Typically, measurements were made during growth periods of 1–4 min depending on growth rate and availability of well-defined profiles. For convenience and accuracy, only images within the same field of view were compared. The area delimited by the cell profiles was computed by using Green's formula (Marsden et al., 1993). Spk diameter was measured
Morphogenesis in \textit{N. crassa} ropy mutants

\textbf{Fig. 1.} Colonies of \textit{N. crassa} wild-type (a), ro-1 (b) and ro-3 (c) grown on VCM (20 ml) for 24 h at 28 °C.

\textbf{Fig. 2.} Bright-field micrographs of hyphae of \textit{N. crassa} wild-type (a, d), ro-1 (b, e) and ro-3 (c, f). Top row after 14 h growth; bottom row after 24 h growth.

Directly on the monitor screen with the line command of the measure option in the Argus-10 menu.

**Spk trajectory analysis.** The centre of the Spk was mapped at 2 s intervals from the videotaped sequences. To quantitate the erratic behaviour of the advancing Spk, we calculated a steadiness index ($S$), namely the ratio of the minimal distance between the initial and final position of the Spk in a given videotaped sequence over the total distance travelled by the Spk. A steadiness index of 1 would correspond to a perfectly straight path. The smaller the steadiness index the less steady the trajectory of the Spk.

**Computer simulation.** From videotaped sequences of hyphae of wild-type and ropy mutants, we mapped the Spk position (every 2 s) and traced the cell profiles at various intervals. These data were fed to the Fungus Simulator (a Windows program for fungal morphogenesis (Bartnicki et al., 1994))
available through the Internet (http://boyce3427.ucr.edu). The Fungus Simulator generates hyphal shapes by a process that mimics a vesicle-based mechanism for cell wall growth.

RESULTS

Colony and hyphal morphology

On agar plates, the two ropy mutants formed colonies that were drastically different from the wild-type. Radial growth was restricted and the aerial mycelium showed the characteristic ropy phenotype (Fig. 1).

Under the microscope, the ropy mutants showed remarkable differences in hyphal morphology compared to the wild-type (Fig. 2). In a young colony (8–14 h), the morphology of ropy hyphae was highly irregular, with extensive curling—an indication of a major loss in growth directionality (Fig. 2b, c). In contrast, the hyphae of the wild-type were more regular, and despite some minor meandering they exhibited long stretches of mostly straight growth (Fig. 2a). In older colonies (> 14 h), the morphology of hyphae growing on the agar surface remained the same but the ropy mutants produced large cottony tufts of aerial hyphae at the periphery of the colony. This aerial mycelium consisted of some very long and straight primary hyphae (Fig. 2e, f) with numerous short curly branches. In the wild-type, mycelium spread faster on the agar surface, producing a flatter and more uniform mat of aerial hyphae. Both surface and aerial hyphae grew rather straight (Fig. 2d).

At high magnification, most wild-type hyphae (Fig. 3a, b) showed a regular apical profile that approximated the ideal shape defined by the hyphoid equation \( y = x \cot(xV/N) \). There were also some slow-growing wild-type hyphae that deviated appreciably from this hyphoid morphology (Fig. 3c). In contrast, the hyphae of the ropy mutants exhibited a great variety of distorted, non-hyphoid morphologies: some apices were blunt, others more pointed; often, the hyphal profile was highly irregular with conspicuous swellings (Fig. 3d–i).

Cytoplasmic appearance

As visualized by phase-contrast microscopy (Fig. 3), there were major differences in cytoplasmic appearance between hyphae of the ropy mutants and wild-type. In the apex of wild-type hyphae, the dominant structure was a large phase-dark Spk (Fig. 3a, b) that corresponds to the organizational pattern 8 described by López-

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**Fig. 3.** Variation in hyphal tip morphology in wild-type and ropy mutants of *N. crassa*. Phase-contrast micrographs of wild-type (a–c), ro-1 (d–f) and ro-3 (g–i) hyphae. Numbers indicate growth rate in \( \mu \text{m}^2 \text{s}^{-1} \). Arrows, Spk; m, mitochondria; v, vacuoles; l, lipid bodies. Arrowheads (in d and g) point at a phase-dark globular body.
Table 1. Cell parameters of N. crassa wild-type, ro-1 and ro-3

Data are mean values ± sd.

<table>
<thead>
<tr>
<th>Strain</th>
<th>No. of hyphae analysed</th>
<th>Growth rate (µm² s⁻¹)</th>
<th>Hyphal diameter (µm)†</th>
<th>Spk diameter (µm)‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type</td>
<td>4</td>
<td>3.47 ± 0.63</td>
<td>10.67 ± 1.03</td>
<td>1.85 ± 0.14</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>1.42 ± 0.20</td>
<td>9.82 ± 0.60</td>
<td>1.35 ± 0.15</td>
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<tr>
<td></td>
<td>4</td>
<td>0.84 ± 0.07</td>
<td>8.28 ± 0.88</td>
<td>1.32 ± 0.30</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>0.61 ± 0.16</td>
<td>7.36 ± 1.97</td>
<td>1.05 ± 0.32</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>0.28 ± 0.04</td>
<td>8.92 ± 1.98</td>
<td>$§$</td>
</tr>
<tr>
<td>ro-1</td>
<td>5</td>
<td>1.38 ± 0.48</td>
<td>8.96 ± 1.54</td>
<td>1.02 ± 0.31</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>0.77 ± 0.06</td>
<td>7.19 ± 0.48</td>
<td>0.82 ± 0.23</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>0.50 ± 0.07</td>
<td>8.29 ± 2.36</td>
<td>0.93 ± 0.28</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>0.24 ± 0.10</td>
<td>6.60 ± 0.82</td>
<td>0.67 ± 0.36</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>0.09 ± 0.02</td>
<td>6.97 ± 2.70</td>
<td>$§$</td>
</tr>
<tr>
<td>ro-3</td>
<td>5</td>
<td>0.83 ± 0.22</td>
<td>8.02 ± 0.89</td>
<td>0.81 ± 0.12</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>0.57 ± 0.03</td>
<td>7.85 ± 0.46</td>
<td>0.96 ± 0.12</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>0.41 ± 0.08</td>
<td>7.13 ± 1.24</td>
<td>0.84 ± 0.05</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>0.29 ± 0.04</td>
<td>7.05 ± 1.44</td>
<td>0.72 ± 0.04</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>0.15 ± 0.03</td>
<td>5.69 ± 0.12</td>
<td>$§$</td>
</tr>
</tbody>
</table>

a For each strain, the values for individual hyphae (Fig. 4) were organized in subgroups in descending order of growth rate.
† Hyphal diameter was calculated by dividing the hyphal area by the length of the hypha.
‡ During each growth sequence, Spk diameter was measured when it was most clearly defined.
§ No Spk was visible.

Franco & Bracker (1996). In the ropy mutants, the Spk was smaller and less phase-dark (Fig. 3d, e, g, h). In the subapex, there were also marked differences between wild-type and ropy. In general, organelle distribution was more uniform in the wild-type. A remarkable difference was in the appearance of mitochondria. In the wild-type strain, these long, phase-dark organelles were the dominant structure of the cytoplasm. In the ropy mutants, the mitochondria were round and less conspicuous, giving the cytoplasm a more finely granular appearance, particularly in slower-growing hyphae.

Another common difference was in vacuolation. The cytoplasm of both ropy mutants was more vacuolated than in wild-type. In ropy hyphae, large, phase-light and phase-dark structures of irregular shape (probably vacuoles and lipid bodies) accumulated at a shorter distance (as close as 10–15 µm) from the tip. Occasionally, one or more small phase-dark globular bodies were seen immediately behind the Spk (Fig. 3d, g; arrowheads). These bodies moved continuously within the area surrounding the Spk but appeared to be physically attached to the Spk by thin filaments and remained behind the Spk for the entire observation period. These granules were usually more conspicuous in the ropy mutants but were sometimes seen in tips of wild-type hyphae. Similar structures have been seen in other fungi (López-Franco & Bracker, 1996) but their function remains unknown.

There was a pronounced difference in overall movement of cytoplasmic structures between wild-type and ropy mutants. To judge differences in organelle movement, the videotaped growth sequences were examined in real time and at three times the original speed. The cytoplasm in the hyphae of ropy mutants was ostensibly less dynamic than that in wild-type hyphae. In the latter, cytoplasmic organelles were in constant motion over the entire length of the subapical area observed (at least 20–25 µm from the apex). In ropy hyphae, motion was less active and was usually restricted to the proximal subapex (5–15 µm from the apex); beyond that, the cytoplasm became static and extensively vacuolated (Fig. 3; see also video in http://boyce.3427.ucr.edu/cytoskel.htm). In both ropy mutants, the movement of cytoplasmic structures appeared less organized than that in wild-type hyphae. In growing wild-type hyphae, the mitochondria moved continuously in a back and forth fashion roughly parallel to the longitudinal axis of growth. In the ropy mutants, mitochondria did not move preferentially along the longitudinal axis but in a more erratic manner.

Table 2. Cell parameters of N. crassa wild-type, ro-1 and ro-3

Values for each strain are the mean of all cells analysed in Table 1. Means within each column followed by the same letter are not significantly different (P = 0.05) according to Student’s t-test analysis.

<table>
<thead>
<tr>
<th>Strain</th>
<th>No. of hyphae analysed</th>
<th>Growth rate (µm² s⁻¹)</th>
<th>Hyphal diameter (µm)</th>
<th>Spk diameter (µm)</th>
<th>Ratios</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Growth rate/hyphal diameter</td>
<td>Growth rate/Spk diameter</td>
<td>Hyphal diameter/Spk diameter</td>
<td></td>
</tr>
<tr>
<td>Wild-type</td>
<td>19</td>
<td>1.40*</td>
<td>9.94*</td>
<td>1.37*</td>
<td>0.13</td>
</tr>
<tr>
<td>ro-1</td>
<td>24</td>
<td>0.63*</td>
<td>7.21*</td>
<td>0.87*</td>
<td>0.09</td>
</tr>
<tr>
<td>ro-3</td>
<td>20</td>
<td>0.50*</td>
<td>6.54*</td>
<td>0.83*</td>
<td>0.07</td>
</tr>
</tbody>
</table>
Growth rate and Spk size

Hyphal elongation is traditionally used as a measurement of the growth rate. This method is accurate provided the hypha maintains a relatively constant shape during elongation. For the *ropy* mutants, with their distorted hyphal morphology, this method was clearly not valid. Instead, we measured growth rate as increase in cell area per unit of time (Table 2) and calculated that the mean growth rate of hyphae of *ro-1* and *ro-3* was 45 and 34%, respectively, of that in wild-type hyphae. There was no significant difference in growth rate between the two *ropy* mutants (Table 2).

Growth rate was correlated with the presence and size of the Spk in hyphae of both wild-type and the *ropy* mutants (Table 1; Fig. 4a). In general, faster-growing hyphae exhibited a larger Spk; this was true for both the wild-type and the mutants. A large well-defined Spk was the rule in wild-type hyphae (Figs 3a, b); only a few hyphae with growth rates lower than 0·35 µm² s⁻¹ (i.e. <10% the rate of the fastest-growing hyphae) failed to exhibit a Spk (Fig. 3c). In the two *ropy* mutants, even the fastest-growing hyphae tended to show a Spk that was less phase-dark and smaller than the wild-type (Fig. 3d, e, g, h). Slow-growing hyphae showed only a very small Spk and none was observed in the slowest hyphae. On average, the Spk diameter of *ro-1* and *ro-3* was 62% and 60%, respectively, of that in wild-type (Table 2). Expressed as cross-sectional area, the relative size of the Spk of *ro-1* and *ro-3* becomes 46% and 41%, respectively, of the wild-type Spk, values that approximate the comparative growth rates calculated above.

Hyphal growth rate and Spk size were further correlated with hyphal diameter in both wild-type and *ropy* mutants (Tables 1 and 2, Fig. 4b, c). In *ro-1* and *ro-3* hyphae, hyphal diameter was difficult to assess because of the irregularities of the hyphal shape. Therefore, we estimated mean diameters by dividing hyphal area by hyphal length. In the wild-type hyphae, there was a positive correlation between growth rate and hyphal diameter. The faster-growing hyphae, which had a large Spk, had the largest diameter (Table 1, Fig. 4b). In the mutants, the same tendency could be observed (Table 1, Fig. 4b).

The appearance and size of the Spk varied greatly during growth, particularly in the *ropy* mutants. For instance, during the observation of a hyphal tip of *N. crassa* *ro-1* growing at 0·6–0·7 µm² s⁻¹, a smaller and less phase-dark Spk was visible for about 220 s (Fig. 5), then the Spk disappeared, coinciding with a sharp decrease of the growth rate to 0·1 µm² s⁻¹. When the growth rate returned to almost the original values (0·5 µm² s⁻¹), the Spk became visible again.

Spk trajectory

The Spk of wild-type hyphae of *N. crassa* follows an intricate trajectory produced by a dominant forward motion accompanied by frequent, short, transverse
Morphogenesis in N. crassa ropy mutants

Fig. 5. Growth rate and Spk changes in a hypha of N. crassa ro-1. The small Spk observed from (a) to (c) is no longer visible in (d), coinciding with a drastic decrease of the growth rate. As growth rate recovers, a small Spk starts being visible again (e).

Fig. 6. Computer simulation of hyphal morphogenesis in wild-type and ropy mutants. The grey areas are the shapes generated by the Fungus Simulator programmed to follow the Spk paths of (a) wild-type hypha growing for 384 s; (b) ro-1 hypha growing for 514 s; (c) ro-3 hypha growing for 491 s. For each hypha, the cell profile (dark solid line) was reconstructed by assembling profiles from three screen displacements. A correction factor was included to compensate for the slight tendency of the microscope stage to drift during the course of the observations (Bracker, 1995). The straight dashed line corresponds to the axis of growth of the cell. Arrows in (c) indicate a data gap in Spk trajectory that occurred during manipulation of the microscope stage. The gap (about 30 s) was filled with a straight line for the simulation. The percentage of mismatch between simulated shape and real shape was 45% for wild-type, 42% for ro-1, and 6.05% for ro-3.

oscillations. The trajectory tends to follow the longitudinal cell axis (Fig. 6a). In the distorted hyphae of the ropy mutants, the Spk advances with an erratic trajectory that also tends to follow the longitudinal cell axis but with larger and longer-lasting departures (Fig. 6b, c). Minor oscillations in Spk trajectory cancelled each other and had no apparent impact on the cell profile. However, larger departures in the trajectory of the Spk resulted in corresponding distortions in the morphology of the hypha.

We compared the behaviour of the Spk in ropy vs wild-type hyphae by analysing trajectory and growth efficiency. To compare trajectories, we calculated the steadiness index (S) (see Methods). In general, the S values for wild-type hyphae were higher than those of the ropy mutants hyphae (Table 3). In the wild-type, the S values were close to 1, indicating a rather steady Spk path, whereas in the ropy mutants, the S values were considerably lower, indicating a much more erratic movement of the Spk.

To compare Spk efficiency, we assessed the increase in hyphal area per total distance travelled by the Spk and also the rate of Spk advance (total distance per unit time). By both criteria, the ropy Spk was less efficient than the wild-type. In area generated per travelled distance, the ro-1 and ro-3 Spk were only 71% and 60%, respectively, as efficient as the wild-type Spk (Table 3). In rate of advance, the values were 83% and 72%. These two criteria combined gave overall efficiencies of 59% and 43% compared to wild-type. The latter values correspond to the overall growth rate differences between wild-type and ropy mutants (Table 3).

Computer simulation

To analyse the effect of Spk movement on hyphal morphogenesis, the traced Spk trajectories and cell profiles of wild-type and ropy mutants were fed to the Fungus Simulator program. The VSC of the simulator was programmed to follow actual Spk trajectories. Cell profiles were used to calculate the amount of vesicles to be released by the VSC at each point in the trajectory. With these two sets of input data, the simulator
The hyphae are those in Fig. 6.

Table 3. Spk behaviour and efficiency in three representative hyphae of N. crassa wild-type, ro-1 and ro-3

<table>
<thead>
<tr>
<th>Strain</th>
<th>Hyphal growth rate</th>
<th>Spk steadiness</th>
<th>Spk efficiency</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total increase of area (µm²)</td>
<td>Time (s)</td>
<td>Growth rate (µm s⁻¹)</td>
</tr>
<tr>
<td>Wild-type</td>
<td>556</td>
<td>384</td>
<td>1.40</td>
</tr>
<tr>
<td>ro-1</td>
<td>472</td>
<td>514</td>
<td>0.92</td>
</tr>
<tr>
<td>ro-3</td>
<td>315</td>
<td>491</td>
<td>0.64</td>
</tr>
</tbody>
</table>

* Overall efficiency is the product of (Area/total Spk travelled distance) times (Total Spk travelled distance/time).

Duplicated the morphologies of wild-type hyphae and the highly distorted shapes of ropy mutants hyphae (Fig. 6).

DISCUSSION

High-resolution video microscopy has allowed us to analyse quantitatively key cellular details of the pheno-
type of two ropy mutants of N. crassa (ro-1 and ro-3). We focused this study on the effects of these mutations (dynein and dynactin, respectively), on the behaviour of the Spk and its consequences on hyphal growth and morphogenesis.

Impairment of dynein or dynactin caused major effects at different levels: reduced growth rate, distorted morphology, aberrant cytoplasmic organization, disrupted organelle motility, and a smaller Spk with erratic trajectory. The effects of these mutations on nuclear distribution and mitosis have been previously document-
ed for this and other fungi (Xiang et al., 1994, 1995; Plamann et al., 1994; Tinsley et al., 1996; Bruno et al., 1996; Inoue et al., 1998). Apparently, the reduction in the overall movement of intracellular components along the microtubular cytoskeleton, caused by dynein/dynactin deficiencies, had a general effect on all cyto-
plasmic activities and hence led to an overall reduction in growth rate. But diminished growth rate need not result in distorted morphology; it should result in slower-growing hyphae with similar morphology. We believe the morphogenetic effects caused by these mutations can be directly ascribed to their impact on Spk formation and behaviour. Invariably, the Spk of ropy hyphae was smaller and lacked the stability of the wild-type Spk. The fact that the ropy mutants grew poorly and produced deformed hyphae on Petri dish cultures in the dark eliminates the possibility that the Spk deficiencies we observed were caused by stresses imposed on the mutants during microscopy.

Spk size

Presumably, the smaller size of the Spk of ropy hyphae is a consequence of a diminished supply of vesicles to the apical region caused by the dynein/dynactin deficiency in the mutants. This conclusion differs from that made by Seiler et al. (1999), who described the ro-1 mutant as having a ‘prominent’ Spk. However, they apparently did not take into account that the Spk of ro-1 is considerably smaller than that of the wild-type, and at times may not even be visible, as shown here in Fig. 4 (see also http://boyce3427.ucr.edu/cytoskel.htm for video).

Consequently, their conclusion that ‘apical transport was intact’ in this ropy mutant runs contrary to our evidence, which clearly shows that the processes responsible for Spk formation, including apical vesicle traffic, are affected by dynein deficiency. Inoue et al. (1998) also found that dynein is required for normal secretory vesicle transport to the hyphal apex of Nectria haematococca.

Since mutations in kinesin are known to affect Spk formation (Seiler et al., 1997; Wu et al., 1998), it was proposed that cytoplasmic microtubules are oriented with their plus ends towards the apex (Lehmler et al., 1997; Seiler et al., 1997, 1999). By the same reasoning, our studies showing that deficiency in cytoplasmic dynein or dynactin affects Spk formation could lead us to the opposite conclusion, namely that the microtubules are oriented with their minus ends towards the apex. Clearly, none of these observations can reveal conclusively the polarity of cytoplasmic microtubules in a hypha. The similarity of phenotypic effects caused by the impairment of opposite motor proteins suggests that both motors are necessary for the maintenance of the Spk and apical growth, but whether or not both are directly involved in the apical transport of secretory vesicles remains to be seen. Possibly, a deficiency in cytoplasmic dynein or kinesin may also impair the endocytotic processes that contribute to the recycling of membranous components from apex to subapex (Hoffmann & Mendgen, 1998), and thus affect Spk formation.

Overall, our analyses showed positive, though not necessarily linear, correlations between Spk size, hyphal growth rate and hyphal diameter in both the wild-type and the ropy mutants of N. crassa. In general, fast-growing hyphae share a tendency to have a larger hyphal diameter and a larger Spk than slow-growing hyphae.
Similar tendencies were observed in other fungi (López-Franco & Bracker, 1996). Wu et al. (1998) found a similar correlation between Spk size, growth rate and hyphal diameter in a kinesin-deletion mutant of Nectria haematococca. In other tip-growing cells such as pollen tubes and root hairs, the size of what would be their Spk equivalent (so-called tip body or clear cap) has also been correlated with high growth rate (Reiss & Hert, 1979; Sievers, 1963).

**Spk behaviour and morphogenesis**

The highly erratic behaviour of the Spk in *ropy* mutants is difficult to interpret since we do not know for sure which cellular components determine the positioning and advance of such a complex and dynamic structure as the Spk. Previously, based on comparative results with benomyl and cytochalasin, we proposed that the microtubule cytoskeleton was directly involved in maintaining the trajectory of the Spk (Riquelme et al., 1998). Our present observations with the dynein-deficient *ropy* mutants lend support to the notion that the microtubule cytoskeleton plays a major role in the formation and behaviour of the Spk. Similarly, Wu et al. (1998) showed that kinesin was essential for normal positioning of the Spk in hyphae of *N. haematococca*. All this leads us to conclude that microtubule-associated motor proteins are necessary for maintenance of a high growth rate, a rather steady Spk and a near-perfect hyphoid shape.

Regardless of the exact mechanism controlling the position of the Spk, we have reason to conclude that the erratic trajectory of the Spk is directly responsible for the distorted morphology of *ropy* hyphae. As we did in studying other morphogenetic processes (Bartnicki-García et al., 1995; Reynaga-Peña & Bartnicki-García, 1997; Riquelme et al., 1998), we used the Fungus Simulator to test the correlation between Spk trajectory and cell shape. The simulator generated forms that reproduced the distorted morphology of the *ropy* hyphae. This indicates that the fungal Spk, operating as a VSC, is the structure that ultimately controls the shape of fungal hyphae. In wild-type hyphae, the Spk advances in a fixed direction with only minor transverse oscillations. The spatially uniform vesicle traffic emanating from such a Spk produces a smooth regular shape that approximates the ideal ‘hyphoid’ shape stipulated by the hyphoid equation (Bartnicki-García et al., 1989). In the *ropy* hyphae, sustained departures in the trajectory of the Spk result in corresponding distortions in the morphology of the hypha.

Barring pleiotropic effects caused by the *ropy* mutations, the dynein/dynactin deficiency probably causes morphogenetic effects by (1) diminishing the traffic of secretory vesicles from their synthesis site (endoplasmic reticulum) either to intermediate secretory compartments or to the hyphal apex; (2) affecting the organization and movement of the growing microtubules; and/or (3) impeding the proper recycling of material needed for normal apical growth to proceed.

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