Temporal tracking of quantum-dot apatite across in vitro mycorrhizal networks shows how host demand can influence fungal nutrient transfer strategies

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Abstract
Arbuscular mycorrhizal fungi function as conduits for underground nutrient transport. While the fungal partner is dependent on the plant host for its carbon (C) needs, the amount of nutrients that the fungus allocates to hosts can vary with context. Because fungal allocation patterns to hosts can change over time, they have historically been difficult to quantify accurately. We developed a technique to tag rock phosphorus (P) apatite with fluorescent quantum-dot (QD) nanoparticles of three different colors, allowing us to study nutrient transfer in an in vitro fungal network formed between two host roots of different ages and different P demands over a 3-week period. Using confocal microscopy and raster image correlation spectroscopy, we could distinguish between P transfer from the hyphae to the roots and P retention in the hyphae. By tracking QD-apatite from its point of origin, we found that the P demands of the younger root influenced both: (1) how the fungus distributed nutrients among different root hosts and (2) the storage patterns in the fungus itself. Our work highlights that fungal trade strategies are highly dynamic over time to local conditions, and stresses the need for precise measurements of symbiotic nutrient transfer across both space and time.

Introduction
Underground, arbuscular mycorrhizal fungi form massive physical networks of hyphae connecting roots of diverse host plants [1]. A single gram of soil can contain ten to hundreds of meters of hyphae that function as a conduit for nutrient transport [2, 3]. These networks, also called common mycorrhizal networks (CMNs), have tremendous effects on nutrient cycling, transferring up to five billion tons of carbon (C) per year worldwide [4–6]. Partnerships with mycorrhizal-type fungi have facilitated major evolutionary events across the globe [7, 8], driving the evolution of complex root traits [9, 10], and creating habitats for various other organisms [11, 12].

Past work has shown that mycorrhizal fungi mediate the success of their hosts, influencing which plants survive and reproduce [13–15]. The fungus does this by forming an intricate hyphal network, which explores the soil and gains access to soil-bound mineral nutrients, such as phosphorus (P) and nitrogen (N). The fungus then exchanges these nutrients for C compounds from the host plant [16–18]. Transfer of resources from a fungus to the host has been shown to be highly variable, and depend on factors including available soil nutrients [19, 20], host species [21], host age [22], light availability [23–27], and even host sex [28]. Because mycorrhizal networks can connect several plants simultaneously, the transfer of resource across a shared fungal network can shift...
depending on these factors. Pioneering work has shown how changing the environment [29] or host composition [30, 31] across a shared network will change resource allocation by the fungus. However, the factors determining exactly which plants get what resources remains an open question.

An emerging body of research is now asking whether we can predict fungal trading strategies across networks containing multiple host plants with diverse demands [32–34]. We define trading strategy as a “conditional strategy that prescribes trading behavior under all circumstances regularly encountered by members of the trading agent’s species” [35]. Trading strategies of non-cognitive agents, such as plants and fungi, are assumed to be shaped exclusively by natural selection. Understanding these trading strategies is important if we hope to manipulate rates of nutrient transfer from the fungus to the host in agricultural settings [36]. For example, it is unknown how the host’s nutritional needs influence the amount and timing of resource transfer by the fungus. When connected in a multiple plant network, do arbuscular mycorrhizal fungi provide more nutrients to plants with higher nutrient demands? And if yes, is there a clear fitness benefit to fungi providing more nutrients to more nutrient demanding hosts?

Until recently, we have been limited in our ability to study the relationship between host demand and fungal trading strategies across shared fungal networks because we have been unable to effectively track the temporal dynamics of the nutrient transfer in the arbuscular mycorrhizal symbiosis. Understanding temporal dynamics is important because fungal strategies are likely to be transient, such that trade with one host, in time, does not imply a consistent supply of nutrients to the same host over an extended period of time [37]. We recently developed a technique to resolve this constraint that allows us to follow P resources tagged with differently colored nanoparticles from a fungal network to host roots [38]. We tagged apatite, a form of mineral P, with quantum dots (QDs). QDs are nanoparticles that fluoresce bright and pure colors when excited with UV light. We used a class of QDs that were highly fluorescent, stable and can fluoresce in different colors depending on chemical composition of the core of the QD [39, 40]. Each color has the same size and weight, making them physically indistinguishable [38]. The outer layer of carboxyl polymers protects the organisms from the heavy metal core, and allowed us to conjugate the QDs to P. To be able to study transfer strategies over time, we synthesized three colors of fluorescent P that were added in a time series. Using confocal microscopy and epifluorescence analyses on plant tissue, we could study where and when P was transferred to different hosts over time, and/or stored by the fungal network.

Our aim was to understand what determines the temporal dynamics of when, and how much, P is transferred to individual root hosts connected by a single mycorrhizal network. Specifically, we asked: (1) does the mycorrhizal network transfer more P to host roots with a higher nutrient demand? (2) Do patterns of P allocation change over time? (3) Is there evidence that the fungus benefits from transferring P to host roots with the highest need? To answer these questions, we designed an in vitro root organ experiment in which a single fungal network colonized two physically separated transformed carrot roots of different ages, of which one was more P limited than the other (Fig. 1). We used a double plate system to physically isolate the roots from each other in separate compartments, while maintaining a living fungal connection between them. The older roots were grown for 6 weeks before the younger roots were transferred into the network. We added QD-tagged apatite at three time points in the compartment of the established root over 3 weeks: first injection, 21 days before harvest (green), second injection, 14 days before harvest (yellow), and the third injection, 7 days before harvest (red).
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25 mL 10 mM sodium citrate solution. After incubating for 2 h at 65 °C to allow any substrate on the outside of the hyphae to be washed away, we separated the extraradical hyphae from the roots and placed the roots in paper bags to dry for a minimum of 48 h at 50 °C. From the dissolved MSR, we took a 1 mL subsample for confocal microscopy analysis of the extraradical hyphal network. We vacuum filtered each sample over a 0.45 µm, 47 mm cellulose nitrate Whatman membrane filter (GE Healthcare, Chicago, USA) to isolate the extraradical hyphae, and then freeze dried the network for 24 h [42]. We recorded dry weight of extra-radical hyphae and roots. We manually homogenized the roots and divided them into two subsamples. From both the established and young roots, we subsampled ~7 mg for epifluorescence analyses, confocal microscopy, light microscopy, and to determine the overall phosphorus concentration of the roots. We subsampled ~20 mg of the larger established roots, and ~1 mg of the young roots for DNA extraction because of their smaller size. We pulverized the root samples for epifluorescence and DNA extraction using glass beads and a bead-beater for 40 s on speed 4 (Thermo Savant FastPrep Fp120 Cell homogenizer, Thermo Fisher Scientific, Waltham, MA, USA).

DNA isolation and real-time quantitative PCR (qPCR) analysis

We isolated fungal DNA from the roots for intraradical hyphal abundance using the DNeasy Plant Mini kit (Qiagen, Hombrechtikon, Switzerland), with the exception that after the lysis step we added 10 µL internal standard, a vector of the cassava mosaic virus, to control for DNA extraction efficiency [42]. To measure the intraradical hyphal abundance, we used a TaqMan probe-based qPCR (LightCycler CFX96, Bio-Rad, Hercules, CA, USA), with probes and primers to target the lesser subunit of mitochondrial DNA of *R. irregularis* [47]. We analyzed each sample on internal standard and *R. irregularis* abundances and exported Cq values at a baseline threshold of 500 relative fluorescent units. We used the internal standard abundance to control for DNA extraction efficiency [42, 47] and converted Cq values to *R. irregularis* mtDNA copy numbers per mg of host root in R version V.3.3.1 [48] with the calibration curves described as described in [47].

Visual root colonization measures

We visually determined the percentage of hyphal colonization, arbuscules, and vesicles by staining a subsample of ~9 mg in a subset of the roots (five root samples per treatment) with trypan blue [49]. We rehydrated the roots in dH₂O and added 2.5% KOH solution to the dried root samples (w/v) and mixed well. We incubated the roots for 5 min in a water bath at 90 °C. After 5 min of cooling, we removed the KOH and rinsed the roots on a sieve with dH₂O. We added 2 mL of staining solution (0.05% trypan blue in destaining solution: lactic acid/glycerol/dH₂O 2:1:1) and mixed well. We incubated the roots for another 10 min at 90 °C in a water bath and after 5 min of cooling outside the water bath, we removed the staining solution by rinsing the roots with dH₂O on a sieve. We stored the roots in Eppendorf tubes in glycerol/dH₂O (1:1) at 4 °C. We analyzed the roots by placing roots on a microscopic slide under an Olympus CX41 microscope with a ×10 ocular and ×40 objective. We scored intraradical hyphae, arbuscules, and vesicles at gridline intersections.

Fluorescence analysis

We analyzed root samples for fluorescence intensity to measure QD-apatite concentrations of three spectra in the roots. We prepared the samples by adding borate buffer (10 mM at pH 7.4) per 1 mg dry root per 150 µL buffer. We then placed five times 1 mg root of each subsample into a 96-well glass bottomed plate (Eppendorf AG, Hamburg, Germany). We obtained spectra from each sample well in the wavelength range of 450–800 nm with 2 nm intervals using a BioTek Synergy MX plate reader at 325 nm excitation (BioTek Instruments, Bad Friedrichshall, Germany). To prevent edge effects, we left the outermost wells empty. To convert photon count to QD concentrations of different colors, we made calibration curves from each color based on seven QD concentrations (13.1, 9.83, 7.37, 5.53, 4.15, 3.11, and 2.33 mM [38]).

We used Matlab R2016a (The MathWorks, Natick, MA, USA) and a custom designed script in Matlab Code based on emission finger printing to deconvolute overlapping emission spectra, quantifying the three colors of QD-apatite simultaneously as described in detail in [38]. This approach allowed us to analyze low levels of QD-apatite (>1 femto mol QD per mg plant tissue) that would otherwise be undetectable using traditional filter and channel-based techniques. We subtracted background emission, and unmixed the overlapping emission spectra with linear models using reference spectra for the autofluorescence and the three QD colors. After unmixing, we optimized and smoothed the spectrum curves to reduce noise and normalized the data. We summed the photon counts of each color separately and converted the total photon counts to nmol of QD-apatite using the above described calibration gradients. To avoid pseudoreplication, we averaged over the five wells.

Total P content of roots

As a second metric to confirm that QD-apatite was being transferred as a P source to host roots, we determined total
root P concentration via acid digestion and spectrophotometry. This total P content includes P present in the phytagel substrate in which the roots grew. We digested a subsample of root material by adding 2 mL digestion mixture (HNO₃/HCl 4:1) per root sample to a Teflon cylinder. After 15 min we closed the cylinders, and placed them for 7 h at 140 °C. We then added 8 mL dH₂O to each cylinder and transferred the contents of the cylinder to a test tube. We left the test tubes for 1 day at room temperature after which we closed the tubes with plastic foil and placed them at 7 °C for 7 days. We then determined the total P content with spectrophotometry. We added 4 mL reagent (1000 mL H₂O, 13.33 mL H₂SO₄, 1.14 g (NH₄)₂Mo₇O₂₄·4H₂O, 1.00 g C₆H₈O₆, 2.6 mg K) to 150 µL of each root sample. After 30 min, we measured the absorbance at 880 nm UV–visible spectrophotometer UV-1601PC (Shimadzu, Kyoto, Japan) and calculated the concentration of P per mg sample. Total nmol of P per mg of root was then used as the denominator to calculate the % of P in the host root derived from QD-apatite, based on the ratio (QD-apatite:P = 1:708) as determined previously [38].

Raster image correlation spectroscopy (RICS) analysis

Epifluorescence analysis on root tissue cannot distinguish between QD-apatite in the root cells and in the intraradical hyphae. Therefore, we used RICS, a technique that can measure particle concentrations from confocal images, to quantify the amount of QD-apatite in root cells versus the amount retained in fungal hyphae [50, 51]. We randomly selected five replicates per treatment and placed these on a microscope slide with 250 µL bicarbonate buffer. We analyzed all samples using an Olympus FluoView TM FV1000 confocal microscope (Olympus, Tokyo, Japan) with a water immersed ×60 UPLSAPO, NA 1.2 objective (Olympus, Tokyo, Japan). Excitation was conducted using a 20 MHz pulsed 405 nm laser (Picoquant, Berlin, Germany). Per sample, we collected five images of intraradical hyphae and five images of uncolonized root cells. The images we collected were 4.096×4.096 µm at a ×50 zoom, with an individual pixel sizes of 16 nm (256×256 pixel frames). We used an optimal pixel dwell time of 20 µs per pixel⁻¹ and a raster line time of 1.25 ms. With the SIM-FCS software [51], we calculated the RICS autocorrelation function, fitted with a 3D diffusion model, and corrected with the moving average intensity correction. We then could calculate the number of particles in nmol per bio-volume root [50, 51].

Growth rate of roots

In order to determine growth rates of roots in our confined system, and confirm that roots were not limited by carbon over the time frame of our experiment, we imaged the growth of Ri T-DNA transformed carrot (D. carota) root organ cultures in a separate experiment. We imaged noncolonized and roots colonized with R. irregularis for 5 months. We filled standard 9-cm Petri dishes with MSR medium, and placed a young piece of ~2 cm root on top. We inoculated the root by positioning a circular agar plug, containing R. irregularis spores and hyphae on top of the roots. To provide similar conditions between the inoculated and non-inoculated plates, we added a sterile and geometrically identical agar plug to the non-colonized roots. We closed the Petri dishes with a double layer of Parafilm M and placed the Petri dishes in incubator at 25 °C in complete darkness.

The imaging system was setup in the incubator. It consisted of a rotary mount, allowing us to image up to six plates automatically. The plates were illuminated exclusively during each capture using dark-field illumination. A single picture per plate was taken with a DSLR camera (EOS 1100D Canon, Tokyo, Japan) placed on top. We imaged the Petri dishes every 2 h for 5 months. The growth rate of the roots was calculated by binarizing the images, in which pixels containing root are one, and the others are zero. The fraction of the gel surface area covered by root was calculated by summing the pixels in the binary image, and dividing by the total number of pixels in the imaged gel area at every time point (Fig. S1).

Videos of nutrient flows in fungal hyphae with and without QD-apatite

To visually confirm that QD-apatite had been uptake by the fungal hyphae, we recorded changes in the flow inside the hyphae with and without the addition of QD-apatite, monitoring for changes due to large particles. We prepared plates of Ri T-DNA transformed carrot (D. carota) root organ cultures colonized with R. irregularis, as above. When the extraradical network was formed (~45 days), 100 µL of QD-apatite solution was added to half the plates. Seven days later, we captured videos of the cytoplasmic flow in the hyphae in the treatments with and without QD-apatite. Videos were made using an inverted microscope (Nikon eclipse ti-E, Nikon, Tokyo, Japan) at ×100 magnification (CFI Plan Apo Lambda 100X Oil, Nikon, Tokyo, Japan). Bright-field illumination was used for plates without QD-apatite solution, while synchronous bright-field and fluorescence illumination was used when QD-apatite was also present. Plates were illuminated with an ultraviolet LED light source at 365 nm (CooLED pE-4000). Grayscale videos of 1 min were recorded using a Hamamatsu Orca-Flash 4 camera at 100 fps with a resolution of 15 pixels per micrometer. For all videos, plates were opened and placed facedown to visualize the fungal hyphae directly.
Statistical analysis

We performed all statistical analysis in R version 3.6.1 [48]. All data, scripts, and analysis are available at: https://github.com/anoukvantpadje/Two_roots. A summary of the data can be found in the Supplementary material (Table S1). We studied the effect of the treatment (control and low-P), compartment (established and young), and time (days since QD injection) on the amount of QD-apatite in the roots. We calculated the ratio of QD-apatite in young/established roots by dividing the amount of QD-apatite in young over the amount in established roots as pairs in a shared plate, and analyzed the effect of treatment and time. To analyze the location of QD-apatite in hyphae/roots, we used our RICS analysis. We calculated the ratio of QD-apatite present in the intraradical hyphae divided by the QD-apatite in the roots per biovolume, for both established and young roots and studied the effect of treatment, compartment and time. We calculated the intraradical fungal abundance per root by taking the logarithm of the total root weight multiplied by the R. irregularis copy numbers per gram of root, and studied the effect of treatment and compartment on the extra- and intraradical fungal abundance and the root weight.

We tested all data for normality with a Shapiro–Wilk test and for homogeneity of variances with a Levene’s test. We checked the distributions of the residuals per eye using QQ plots. Due to non-normality and heteroscedasticity, we transformed the QD-apatite in roots, the ratio of QD-apatite in young/established roots, the extraradical and intraradical fungal abundance by taking the square root. We transformed the location of QD-apatite in hyphae/root by taking the logarithm.

We analyzed QD-apatite in roots, the ratio of quantum-apatite in young/established roots, the extra- and intraradical fungal abundance and the root weight with linear mixed effect models, using the R package lme4 [52]. We assigned Petri dish as a random effect to correct for the non-independent measurement of the two root systems growing on the same plate. We produced type II ANOVA tables with the Kenward–Roger method to the denominate degrees of freedom and F-statistics using the R Package lmerTest [53]. We also assessed the difference in the ratio of QD-apatite in young/established roots at specific time points with three separate Student’s t tests. We analyzed the location of QD-apatite in hyphae/roots with a linear model and produced type II ANOVA tables using the R Package car [54].

Results

We first visually confirmed that QD-apatite was taken up by the fungal hyphae by comparing videos in which hyphae were exposed or not exposed to QD-apatite. We noted the large (~5 µm) vacuoles inside the hyphae when QD-apatite was present (Video S1), but never in the plates (ten replicates) when QD-apatite was absent (Video S2). We then quantified the distribution of QD-apatite across the established and young roots using epifluorescence measurements to determine how much was transferred by the mycorrhizal network from the established root compartment (where the QD-apatite was pipetted) to the younger roots. We found that QD-apatite per mg root (weight including intraradical fungal biomass) was significantly influenced by time, compartment (established versus young), and the interaction between time and treatment (Table 1). Specifically, in the first injection (i.e., green QDs, 21 days), we found more QD-apatite in the established roots compared to young roots, across both treatments (Fig. 2). However, this was a small difference given that the established roots had direct access to nutrients, while the young roots had access only indirectly via the fungal network. For example, in the control treatment, the young root had only 13% less QD-apatite per mg of root than the established, and in the low-P, the young root had only 15% less established root (Fig. 2).

Second, we found that by the termination of the experiment, the nutrient condition for the host roots affected the nutrient allocation patterns of the fungus (Fig. 3). We plotted the ratio of QD-apatite in two host roots (young/established), allowing us to study the transfer strategies of the fungus. We found a significant effect of time in both treatments (Table 1 and Fig. 3). We then compared the allocation patterns in each treatment for each time point. We found no statistically significant difference in allocation patterns of QD-apatite based on the first (Student’s t test: t = −0.067, p = 0.946) nor the second injection (Student’s t test: t = −0.506, p = 0.617). However, by the third injection, the ratio was significantly higher, such that there was more P allocated to young roots in the low-P treatment compared to the control (Student’s t test: t = −2.223, p = 0.032, Fig. 3). Over time, differences in nutrient transfer to roots could depend on P condition of the young root, with more nutrients being allocated to those roots in low-P conditions.

To confirm this trend using a different metric, we determined total P concentration of host roots via acid digestion technique (Fig. S2) and compared this to the % of P from fluorescing QD-apatite. We found that the % of P from the QD-apatite over total root P was significantly higher in roots growing under low-P conditions (~6.4%) compared to control (~4.7%) (Fig. 4a and Table 1). When plotting this as a time series, we found a significant effect of compartment (established versus young) and the interaction between treatment and compartment in by the third QD-apatite injection (7 days), with more P from QD-apatite in young roots in the low-P treatment (Fig. 4b, c and Table 1).
Table 1 | Analysis of variances.

|                          | df | res     | F value  | p value |
|--------------------------|----|---------|----------|---------|
| **QD-apatite per mg root** |    |         |          |         |
| Time                     | 2  | 222.439 | 1070.87  | <0.0001*|
| Treatment                | 1  | 46.701  | 2.321    | 0.134   |
| Compartment              | 1  | 229.886 | 18.751   | <0.0001*|
| Time × treatment         | 2  | 222.462 | 3.424    | 0.034*  |
| Time × compartment       | 2  | 222.468 | 2.891    | 0.0576  |
| Treatment × compartment  | 1  | 230.33  | 2.611    | 0.108   |
| Time × treatment × compartment | 2 | 222.479 | 0.636    | 0.53    |
| **Ratio of QD-apatite in young/established roots** |    |         |          |         |
| Time                     | 2  | 83.852  | 13.968   | <0.0001*|
| Treatment                | 1  | 42.634  | 2.995    | 0.091   |
| Time × treatment         | 2  | 84.123  | 1.838    | 0.165   |
| **P per mg root**        |    |         |          |         |
| Treatment                | 1  | 58      | 11.478   | 0.001*  |
| Compartment              | 1  | 58      | 1.382    | 0.245   |
| Treatment × compartment  | 1  | 58      | 0.003    | 0.957   |
| **% P from QD-apatite**  |    |         |          |         |
| Treatment                | 1  | 48      | 5.174    | 0.027*  |
| Compartment              | 1  | 48      | 0.09     | 0.766   |
| Treatment × compartment  | 1  | 48      | 0.771    | 0.381   |
| **% P from 1st QD-apatite addition (21 days)** |    |         |          |         |
| Treatment                | 1  | 48      | 7.066    | 0.011*  |
| Compartment              | 1  | 48      | 0.127    | 0.723   |
| Treatment × compartment  | 1  | 48      | 0.386    | 0.538   |
| **% P from 2nd QD-apatite addition (14 days)** |    |         |          |         |
| Treatment                | 1  | 48      | 0.08     | 0.778   |
| Compartment              | 1  | 48      | 2.764    | 0.103   |
| Treatment × compartment  | 1  | 48      | 1.301    | 0.26    |
| **% P from 3rd QD-apatite addition (7 days)** |    |         |          |         |
| Treatment                | 1  | 48      | 2.802    | 0.1     |
| Compartment              | 1  | 48      | 6.812    | 0.012*  |
| Treatment × compartment  | 1  | 48      | 6.245    | 0.016*  |
| **QD-apatite retention** |    |         |          |         |
| Treatment                | 1  | 42      | 5.419    | 0.025*  |
| Compartment              | 1  | 42      | 4.676    | 0.036*  |
| Time                     | 1  | 42      | 3.129    | 0.054   |
| Treatment × compartment  | 1  | 42      | 0.959    | 0.333   |
| Treatment × time         | 2  | 42      | 5.421    | 0.008*  |
| Compartment × time       | 2  | 42      | 1.352    | 0.27    |
| Treatment × compartment × time | 2 | 42 | 0.2 | 0.82 |
| **Root biomass**         |    |         |          |         |
| Treatment                | 1  | 56.59   | 0.008    | 0.928   |
| Compartment              | 1  | 56.204  | 846.59   | <0.0001*|
| Treatment × compartment  | 1  | 56.59   | 0.783    | 0.38    |

Table 1 (continued)

|                          | df | res     | F value  | p value |
|--------------------------|----|---------|----------|---------|
| Extraradical hyphae biomass |    |         |          |         |
| Treatment                | 1  | 56.114  | 11.659   | 0.001*  |
| Compartment              | 1  | 55.681  | 0.419    | 0.52    |
| Treatment × compartment  | 1  | 56.114  | 0.003    | 0.954   |
| Intrapradical colonization (copy numbers) |    |         |          |         |
| Treatment                | 1  | 52.496  | 5.318    | 0.025*  |
| Compartment              | 1  | 37.803  | 305.721  | <0.0001*|
| Treatment × compartment  | 1  | 52.496  | 0.653    | 0.423   |
| Percentage colonization  |    |         |          |         |
| Treatment                | 1  | 103.918 | 1.894    | 0.202   |
| Compartment              | 1  | 216.767 | 3.951    | 0.078   |
| Treatment × compartment  | 1  | 34.191  | 0.623    | 0.45    |
| Percentage arbuscles     |    |         |          |         |
| Treatment                | 1  | 8.0     | 0.0426   | 0.8411  |
| Compartment              | 1  | 169.056 | 8.9945   | 0.0150* |
| Treatment × compartment  | 1  | 0.8     | 0.995    | 0.995   |
| Percentage vesicles      |    |         |          |         |
| Treatment                | 1  | 1.6815  | 0.34     | 0.574   |
| Compartment              | 1  | 31.142  | 6.302    | 0.033*  |
| Treatment × compartment  | 1  | 2.9314  | 0.593    | 0.461   |

Degrees of freedom (df), residuals (res), F and p values are given for the effect of the variables. p values with a * have a significant effect of the variable (p < 0.05).

Fig. 2 | nmol QD-apatite per mg of host root. a Established roots in the control treatment contained significantly more QD-apatite per mg of root than young roots across all three time points: first addition, 21 days before harvest (green), second addition, 14 days before harvest (yellow), and the third addition, 7 days before harvest (red). b Established roots contained more QD-apatite from the first injection in the low-P treatment, but this significant difference disappeared by the last injection, 7 days before harvest (red) (* ncontrol = 24, nlow-P = 25). Means ± SEM.
Next, we were able to identify the exact spatial location of the QD-apatite (i.e., retained in the fungal hyphae versus transferred to the root) using RICS analysis. We found that spatial location of the QD-apatite was significantly influenced by treatment, compartment, and by the interaction between treatment and time (Table 1). From the first and second injection, we found no significant difference in the ratio of QD-apatite located in hyphae/roots between low-P versus control treatment. However, by the final injection, the ratio of QD-apatite in hyphae versus root was significantly lower in the young roots of the low-P treatment versus the control (Fig. 5 and Table 1). This result suggests that, rather than retaining the QD-apatite in the hyphae, the QD-apatite was more likely to be transferred to the roots, when there was a higher P limitation for roots. Over the time frame of our experiment, we found no significant effect of our treatment on root growth. Root biomass was not significantly different between the control and low-P treatments in both the established and young roots. We found that established roots were larger than young roots, as expected because they were planted 6 weeks earlier (Table 1). We then analyzed data from the root growth imaging experiment to confirm that roots were not C limited in the time frame of our experiment. We found root growth increased rapidly for the first 2 months, with root growth rate decreased to almost zero after ~3 months (Fig. S1a). Even after 3 months, we documented that the hyphal network continued to grow and sporulate (Fig. S1b), suggesting the allocation of C to the fungi was still not limited.

We then studied the biomass of the fungi to determine if the way the fungus distributed nutrients among the two roots affected their own success. We asked whether providing more nutrients to the young root growing under the low-P conditions provided a benefit to the fungi, as measured by their extra and intraradical growth. We found overall higher biomass of extraradical hyphae in the control treatment compared to low-P, but no other statistically significant differences between young and established roots (Table 1 and Fig. 6a).

Finally, we analyzed intraradical fungal colonization per mg of root, as measured by *R. irregularis* copy number and P from QD-apatite (%)

| Treatment | Control | Low-P |
|-----------|---------|-------|
| Days since QD-apatite addition | 21 | 14 | 7 |
| Ratio of QD-apatite in young/established roots | 0.5 | 1.0 | 1.5 |

Fig. 3 Ratio of QD-apatite in young/established roots. In the first two QD additions, there were no significant differences in allocation patterns of QD-apatite between young (light blue) and established roots (dark blue). However, by the third addition, the ratio was significantly higher, meaning more QD-apatite allocated to young roots, in the low-P treatment compared to the control (n<sub>control</sub> = 24, n<sub>low-P</sub> = 18). The asterisk indicates a p value < 0.05. Means ± SEM.

Fig. 4 Percentage of P from QD-apatite in host roots. a Across all time points and treatments, an average of 5.6% of the P in the host roots originated from a QD-apatite source. The percentage of P from the QD-apatite was significantly higher in roots growing under low-P conditions. b, c In the first QD-apatite injection (21 days), %P from QD-apatite was significantly affected by treatment, with more P from QD-apatite in the low-P treatment. In the second QD-apatite injection (14 days), we found no significant effect of treatment, compartment, or the interaction. In the third QD-apatite injection (7 days), we found a significant effect of compartment and the interaction between treatment and compartment, with more P from QD-apatite in young roots in the low-P treatment (n<sub>control,established</sub> = 32, n<sub>control,young</sub> = 2, n<sub>low-P,established</sub> = 24, n<sub>low-P,young</sub> = 3). The asterisk indicates a p value < 0.05. Means ± SEM.
We found a statistically significant effect of treatment and compartment on intraradical fungal colonization as measured by mtDNA copy number, but no effect in the interaction between treatment and compartment (Table 1). The colonization of control roots was on average 125 times higher than that of low-P roots. The colonization of the younger roots was on average 32 times higher than that of established roots (Fig. 6b). We visually quantified the % of hyphae, arbuscules, and vesicles using microscopy and found levels of hyphal colonization ranging from 18 to 38%, and arbuscules and vesicles around 0.5–10% of root length (Table 1 and Fig. S3).

**Discussion**

Our aim was to understand the allocation dynamics of P transfer from fungal networks connected to two host roots...
using a recently developed QD-nutrient tagging approach. We asked (1) whether the distribution of P from a mycorrhizal network to a host root depends on the P needs of the host, specifically asking if fungal networks transfer more P to hosts growing under lower P conditions, (2) if fungal P allocation patterns changed over time, and (3) if the fungus benefited from providing more P to roots with a higher P demand.

**Nutrient need of the host plant influences the nutrient transport of fungi**

First, our data suggest that QD-apatite can be taken up in one part of the fungal network and transferred through the network to a different host root. In our case, this movement was from an established root to a younger root (Fig. 2). We found that the P needs of the host can influence this distribution; however, these distribution differences take time to emerge. Specifically, we found that after 3 weeks, the fungus transferred more QD-apatite to young roots growing under low-P conditions (Fig. 2). By the third injection, the ratio of QD-apatite in young/established roots was significantly higher under lower-P conditions than the control treatment (Fig. 3), and more of the total P was coming from QD-apatite (Fig. 4). Before this time point, we saw no effect of the nutrient status of the younger root. This small delay is not surprising given the nutrients were added directly to the compartment of the established roots, whereas the younger roots had access only through the shared network.

While we found that P allocation by the fungus is influenced by the P status of the host, the physiological mechanisms and regulating gene networks underlying this effect are still unknown. Key genes, such as MtPT4 the mycorrhiza-inducible phosphate transporter, play an important role in regulating the P transfer at the symbiotic interface between the root cells and the fungus [55]. However, many additional (and complimentary) candidate genes in nutrient sensing and regulation are now emerging [56], including those involved in fatty acid transfer [57]. One major constraint in identifying underlying patterns in these regulation mechanisms is that similar physiological reactions can be driven by different molecular mechanisms in distantly related plant species [58].

Although we observed a higher amount of QD-apatite per mg of root in the established roots from the first injection, this was only 13–15% higher compared to younger roots (Fig. 2). This is a surprisingly small difference given that the established roots had direct access to the nutrients (including P readily available in the MSR medium), and thus these roots could potentially utilize a “direct pathway” via root transporters [59, 60], rather than relying on the fungus for P uptake and transfer. However, direct uptake of nutrients can be limited when the plants are colonized by arbuscular mycorrhizal fungi [60, 61, 62]. While direct uptake of QD-apatite by whole plants in the absence of mycorrhizal fungi has been documented, it was shown to be significantly less per mg of root compared to when mycorrhizal fungi are present [38]. This is because the rock phosphate apatite as a nutrient source is generally difficult for roots to solubilize.

Past work has shown that arbuscular mycorrhizal fungi can increase the dissolution and uptake of apatite [63, 64]. While the apatite uptake mechanism for arbuscular mycorrhizal fungi is still unknown, fungi generally use endocytic pathways to take up large particles [65–67]. For example, clathrin-mediated endocytosis has been shown in the fungus *Candida albicans*, with invagination cells reaching diameters of 100 nm [68]. Using bright-field and fluorescence imaging of flows, we found large vacuoles were formed inside the hyphae when the fungus was given access to QD-tagged apatite (Video S1). In hyphae not exposed to QD-tagged apatite, we noted the absence of these large vacuoles (Video S2). While endocytic pathways are likely to be responsible, we assume there is an initial dissolution of the apatite by mycorrhizal fungi prior to uptake. This process likely reduces the size of the tagged apatite crystals and facilitates translocation. There are many apatite solubilizing bacteria described [69, 70], but the ability of arbuscular mycorrhizal fungi to dissolve apatite in absence of any bacteria is still an active area of research. Previous studies have shown that mycorrhizal species *R. irregularis* and *Glomus custos* are able to dissolve apatite when in symbiosis with *Plantago lanceolata* in sterilized soil [63], and *Glomus clarum* increased the dry weight of maize plants fertilized with rock phosphate [71]. Likewise (although in non-sterile soil), the distantly related *Glomus tenuis* significantly increased P uptake from rock phosphate in four different crops [72]. In agreement with these past studies, we found that, across treatments, ~5.6% of the root P content was from derived from QD-apatite (Fig. 4).

Although our results confirm that larger, established mycorrhizal roots directly adjacent to nutrient sources may initially dominate nutrient uptake [73], we found that this can change over time as the fungal network continues to relocate nutrients to other host roots. Past work has shown how seedlings receive nutrients from mycorrhizal networks, despite not being the main sinks [74–76]. However, it has also been shown that smaller plants can suffer when connected to larger plants via a mycorrhizal fungus [29]. A review of the literature found no clear pattern in how seedlings respond to being incorporated into mycorrhizal networks, with a meta-analysis finding that roughly half (42%) of the seedling species investigated responded positively to the presence of a CMN, while negative or neutral effects were found in the remaining cases [77]. We argue that this lack of consistency may be driven by a time
component. Our work suggests that the transfer of P from the fungal network is highly dynamic and that nutrient allocation dynamics need to be measured across multiple time points, and potentially over even longer time scales than the 3 weeks measured here.

A second issue is the sink dynamics of the host roots. If the established roots had depleted their C source by week 3, then there is the question of whether they could still “pay” for their P. This could result in more P being allocated to the younger root simply because of C limitation in the established root. However, by adding excess sucrose to the MSR medium (10 gL⁻¹), we ensured there was no C shortage. This was confirmed by our imaging data (Fig. S1), which demonstrated that root size increased at high rates, (between ~1 and 3 mm²/h, for the first ~3 months) with no evidence of C shortage to fungal allocation even after 5 months.

**Fungal nutrient allocation strategies: retention or transport**

Using confocal microscopy, we were able confirm that the QD-apatite was taken up by the fungal hyphae and transferred from intraradical hyphae to the host roots (Fig. 5). Specifically, our confocal RICS technique allowed us to visually determine the proportion of QD-apatite physically retained in the hyphae versus transferred to the host roots.

Based on data from the first addition, we observed that the ratio of QD-apatite retained in the hyphae did not differ in established or young roots, or under control and low-P treatments (Fig. 5). This ratio of QD-apatite retained in hyphae divided by amount transferred to roots was slightly below one, suggesting that equal amounts of QD-apatite per biovolume were present in both tissue types. However, by the third injection, we saw a different pattern: a lower ratio indicated that more QD-apatite was transferred to the roots than retained in the hyphae. This ratio was significantly lower in the young roots under low-P conditions versus control conditions, suggesting low resource conditions stimulated more nutrient transfer to roots and less hyphal storage (Fig. 5). Because historically there has been an inability to separately quantify nutrients into those transferred to the roots and those retained in the hyphae, it has been difficult to study nutrient allocation strategies from a fungal perspective. This new resolution offers the potential to quantify these strategies over space and time using differently colored nutrients in combination with precision RICS techniques [38, 50, 51, 78, 79].

**Fungal benefit**

We have shown that transfer of nutrients from fungal networks to hosts can be influenced by the host’s nutrient status: more QD-apatite was transferred to young roots under low-P conditions (Figs. 2 and 3). We suggest that the fungi employ this strategy not to “help hosts” as is often invoked in the literature, but rather because this strategy has the potential to increase direct benefits to the fungus. Theoretical [34] and empirical [24, 25] work suggests that, rather than a fixed exchanged of nutrients between partners, allocation of resources depends strongly on context, such as host growth conditions [23–26], externally available nutrients [19, 20, 80, 81] and number of competing fungi [82]. Specifically, biological market theory predicts that as a resource becomes more limiting, the value and hence price, of that resource potentially increases [34, 83, 84].

One of our aims was to test whether the fungi benefited from providing hosts growing under low-P conditions with an increasingly valuable resource. Specifically, we were interested in cumulative measures of C allocation by the host root to the symbiont. However, C allocation from host to arbuscular mycorrhizal fungi is notoriously difficult to measure. Physiological limitations will likely prevent the development of QD-tagged C because organisms consume C faster than P, potentially degrading QDs over time, exposing the heavy metal core of the QD, and leading to toxicity. Also, it is questionable as to whether QD-tagged C, for example, QD labeled hexose injected into root organ cultures, could ever serve as a relevant proxy for host C allocation dynamics. Traditional C labeling techniques likewise make it difficult to obtain accurate “cumulative” patterns as C is rapidly respired. However, new approaches using NanoSIMS (Nanoscale Secondary Ion Mass Spectrometry), and even NanoSIP (Nanoscale Stable Isotope Probing), may prove useful in tracking C pathways at the nanoscale in the future [85].

Given the limitations of cumulative C tracking, we measured the extraradical biomass of the fungi as a proxy for fungal benefit. Biomass can be a useful proxy because arbuscular mycorrhizal fungi are obligate biotrophs, meaning all their C is supplied directly from the host. Using the biomass measured from destructive harvesting of the samples, we did not find higher total biomass of extraradical hyphae in the low-P conditions (Fig. 6a). However, biomass measurements are imperfect because it is unknown how much C was allocated per each root system (i.e., established versus young) because C can be transferred across the fungal network. Therefore, we cannot exclude that C gained in the younger root compartment was transferred to the established root compartment, and vice versa. Ideally, we would use a method to quantify continuous C allocation that did not rely on destructive harvesting. New high-resolution imaging techniques, that convert the visual structures of mycorrhizal hyphal networks to biomass proxies, are a potential way forward.
Open questions regarding QD-tagged nutrients

While we found support for our hypothesis that mycorrhizal networks transfer more P to the hosts with a higher nutrient demand, we still have many open questions regarding the QD-nutrient tagging technology. For example, it has not yet been possible to quantify the rate at which P is dissociated from the QD core across different biological tissue. One approach would be to use anisotropy techniques, which involve measuring the rotational speed of partly dissolved versus fully intact QD-apatite in different biological tissue. However, this requires that the QD-tagged nutrients have a fixed dipole moment, which is unlikely. A second approach is to create “radioactive QDs” to determine if the P from the nanoparticles becomes incorporated into DNA or phospholipids. However, the dual radioactive labeling + QD approach still fails to give a direct measurement of how much of the P remains attached to the QD. Rather, it can only tell us that the radioactive P initially attached to the QD core was eventually used in building biological tissue. Lastly, Raman microscopy is theoretically possible, although typical Raman signals are weak, and thus require long measurement times. Since Raman microscopy relies on monitoring vibrations, it will be necessary to find a specific vibration energy different enough from the vibration energies of other covalent bonds of other molecules in the cell. Therefore, it is an open question whether QD vibrations are both specific and strong enough to overcome the signals from the rest of the cell.

While exact decay measures of P dissolution from the QD tags are difficult to obtain, there is both direct and indirect evidence that QD-tagged nutrients are being used by the root cells to build biological material. For example, there is data showing that *Medicago* plants colonized by arbuscular mycorrhizal fungi grow equally well on apatite versus QD-apatite tagged when these nutrients are their sole P source. If the P from the QD-apatite was not available or even toxic, we would expect a growth depression in replicates grown on QD-apatite compared to apatite [38]. Furthermore, by measuring total P concentrations of roots versus P contribution from QD-apatite (Figs. 4 and S2), we show that host roots will use significantly more QD-apatite when grown in low-P conditions. This effect would not emerge if the tagged P was biologically unavailable. Lastly, we have direct evidence of QD-apatite being translocated in root cell compartment using RICS. Translocation from the fungus to roots cells is expected only when P remains conjugated to the QDs. Past work has shown that when fungal hyphae are exposed to unconjugated carboxyl terminated QDs, there is no uptake or translocation of QDs to the root: after 60 days of exposure values remained lower than the detection limit; <0.000001 nmol QD mg\(^{-1}\) plant tissue [38].

Conclusion

Studying the nutrient allocation strategies of symbiotic microbes to their hosts remains a major challenge [84, 86]. In the case of arbuscular mycorrhizal fungi, slight alterations to the composition or growth conditions for the fungal network can result in shifts in nutrient trading strategies [87]. We have shown that time and host nutrient demand both play an important role in fungal allocation patterns. Under natural conditions, allocation patterns are likely to very dynamic and difficult to predict. This is especially true as host species composition changes, and abiotic factors vary seasonally [26, 29, 37, 88].

Utilizing QD techniques to study symbiotic trade is still in its infancy, but the technique can begin to be a useful tool to help us understand how fungal allocation patterns change under these various environmental contexts, across both space and time. Until we obtain exact QD-apatite decay measurements across different tissues, and we successfully characterize the QD-apatite uptake mechanisms by fungi and transfer from fungi to roots, caution is still needed. As these approaches become more refined, we can test predictions of resource exchange in host-symbiont relationships [34, 89], and determine the conditions under which partnerships are likely to bring direct and measurable benefits.

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Author contributions AP, LOG, and MK performed the experiments and measurements. AP and LOG performed all statistical analyses, created figures, and drafted the initial paper. MAH and MP were involved in RICS and fluorescence analysis. ETK and TS were involved in the experimental design and paper writing.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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