Dear Editor,

This letter is in reference to our revised manuscript, bg-2018-375. In most specific cases in the open discussion we agreed with the reviewers and have incorporated the suggested changes into the revised manuscript. For this author response, as we have already provided a point-by-point reply to both sets of reviewer comments during public discussion, we do not repeat these responses in this document.

In the revised manuscript we have altered substantial portions of the text. The major changes, which do not include numerous small text edits, are in summary:

1. Substantial rewrites of the introduction and discussion sections of the manuscript
2. Justification for the use of a technically non-replicated design based on ecosystem-level eddy covariance footprints, and consideration of the implications of this for the overall interpretation of the experiment
3. Addition of additional data regarding pretreatment conditions
4. Addition of additional data regarding soil N:P stoichiometry during the period of the study (which also involved adding an additional author)
5. Clarification of some of the methods, including correction of ambiguous phrasing and (hopefully) clearer descriptions of the design and sampling protocol
6. Addition of a figure illustrating the experimental design and theory behind the experiment
7. Corrections to the several mistakes in the figures in the original documents as well as addition of finer scales to timeseries figures
8. Removal of some abbreviations for easier readability

We have also added a short supplement showing the N and P concentrations used to assess N:P stoichiometry. We follow with a marked up version of the revised manuscript.

Richard Nair (on behalf of co-authors)
N:P STOICHEMISTRY AND HABITAT EFFECTS ON MEDITERRANEAN SAVANNA SEASONAL ROOT DYNAMICS

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Abstract.

Mediterranean grasslands are highly seasonal and co-limited by water and nutrients. In such systems, little is known about root dynamics which may depend on plant habit-individual plant properties and environment as well as seasonal water shortages and site fertility. This latter factor is Patterns of root biomass and activity are affected by the presence of scattered trees and site management including grazing, as well as grazing, site management, and chronic nitrogen deposition, which may lead to all of which can affect nutrient ratios and potentially cause development of nitrogen:phosphorus (N:P imbalance) imbalances in ecosystem stoichiometry. In this study we combined observations from minirhizotrons collected in a tree-grass Mediterranean savanna (Spanish/ textitDehesa with root measurements from direct soil cores and ingrowth cores and along with measures of above-ground biomass to investigate seasonal root dynamics and root:shoot ratios in a Mediterranean tree-grass 'savanna'. We investigated responses to soil fertility, using a nutrient manipulation (N / NP addition) and microhabitats effects-addition and spatial microhabitat treatments between open pasture and under tree canopy locations. Root dynamics over time were also compared with indices of above-ground growth drawn from proximal remote sensing (Normalised Difference Vegetation Index and Green Chromatic Coordinate derived from near-infrared enabled digital repeat photography).

Results show distinct differences in root dynamics and biomass between treatments and microhabitats. Root biomass was higher with N additions, but did not differ from the control with NP additions in early spring, but by late season the growing season root biomass had increased with NP in open pastures but not higher than N alone. Root added alone. In contrast, root length density (RLD) in pastures responded stronger to the NP than N only treatment, but only in the pasture addition, while beneath trees RLD root biomass tended to be higher with only N. Nutrient additions reduced root. Even though root biomass increased, root:shoot ratio but less than overall increases in root biomass decreased under nutrient treatments. Timing of root and shoot growth was reasonably well paired, although in autumn root growth appeared to be substantially slower than 'regreening' of the system. We interpret this difference-these differences as a shift in community structure and/or root traits under P fertilization and reduced nutrient limitations. The timing of maximum root cover, as assessed by the minirhizotrons, did not match with above-ground phenology indicators at the site as root growth was low during autumn despite the 'greening
up' of the ecosystem. In other periods, roots responded quickly to rain events on the scale of days, matching changes in above-ground indices. Our results highlight the need for high resolution sampling to increase understanding of root dynamics in such systems, linkage with shifts in community structure and traits, and targeting of appropriate periods of the year for in-depth campaigns changing stoichiometry induced by the fertilization. We also consider seasonal (phenology) differences in the strength and direction of effects observed.

1 INTRODUCTION

Terrestrial semi-arid ecosystems are important determinants of the interannual variability in the land C sink (??) and switch from net C sources to C sinks through the year due to plant phenology (??). Large regions in this climate zone are characterized by carbon (C) sink (Ahlstrom et al., 2015) as variation in C uptake is controlled by plant phenology (Randerson et al., 1997; Richardson et al., 2010) which responds to seasonal and interannual variation in climate conditions. In Iberia, one example of these semi-arid ecosystems is a managed agro-silvopastoral 'savanna' systems, such as the Iberian ('Dehesa' and 'Montado'). These are savanna, known as 'dehesa' or 'montado' (Spanish and Portugese names, respectively). Similar systems are common in other mediterranean countries (den Herder et al., 2017) and worldwide (Campos et al., 2013; Porqueddu et al., 2016).

Dehesas consist of 20-40 % Quercus ilex. Ballota (Desf.) and Quercus suber (L.) canopy cover with seasonally variable intercanopy grassland mediated-maintained by livestock grazing (??). Regions around the Mediterranean basin are particularly vulnerable to climate change (??) due to increasing aridity and increases in temperature (??). Savanna regions are also particularly badly represented in predictive models (??) due to their complex structure, which is a result of (Moreno and Pulido, 2009). They are a man-made conversion from oak forest (Joffre et al., 1999) as a combination of low water and nutrient availability limiting seasonal water availability and typically low nutrient availability limits conversion to other cover types (??) as well as management practices including tree thinning and grazing (??). Dehesa systems are a man-made conversion from oak forest (??) and while water limits (Eagleson and Segarra, 1985). Their complex structure and multiple seasonally limiting resources (water in summer, other-nutrients in wetter periods of the year are wet enough for nutrient limitation to dominate. Hence the slow shifting of nutrient limitations in future scenarios due to ...) means these regions are also particularly badly represented in vegetation components of predictive models (Beringer et al., 2011). These areas around the Mediterranean basin are also especially vulnerable to climate change (Giorgi and Lionello, 2008; Sillmann et al., 2013) due to increasing aridity, increases in temperature (Peñuelas et al., 2018) and other environmental changes due to human activity. Particular among these are nitrogen:phosphorus (N:P) imbalances (Peñuelas et al., 2013), which result from chronic N inputs (from deposition and management) resulting in ongoing N:P imbalances (??) is of particular interest but by necessity at a higher rate than P inputs. These stoichometric imbalances may have major impacts on plant functioning, but must be considered within the context of both major structural, micrometeorological and soil fertility-associated differences between microsites (??) and the heavy impact of summer drought, tree- and grass-dominated microsites (Moreno et al., 2013) and the impact of severe summer droughts.
Most ecological attention is focused on above-ground organs, particularly in phenological studies (Radville et al., 2016), despite the fact that below-ground systems are the main source of carbon dioxide emissions to the atmosphere (Schlesinger and Andrews, 2000), contain 2/3 of the world’s carbon (C) stocks (Batjes, 1996) and are the site of plant uptake of both water and nutrients. In grassland, uptake of both water and nutrients from the atmosphere via roots. In grasslands (and by extension, grass-dominated systems such as savannas), below-ground systems are also the site of most competition between individuals (Mokany et al., 2006), the major short-term sink for recently fixed C due to high ratios of roots to shoots (Hui and Jackson, 2006; Mokany et al., 2006), the main source of litter (Casals et al., 2010), and the main contributor to long-term soil C stocks (Rasse and Smucker, 1998), and a major site of niche differentiation between plant forms (Moreno et al., 2013). In most ecosystems, root biomass changes substantially throughout the year, although understanding drivers of phenological change in this phenology is limited, especially when using quantitative metrics. This includes considerable uncertainty about both global change factors (e.g. elevated atmospheric CO₂), increasing temperatures, precipitation changes, N availability, and local fertility, other soil properties such as compaction, as well as species-specific determinants (Radville et al., 2016). In many cases, root growth is also desynchronized from production of shoots (e.g. trees). Hence phenology of biomass (Blume-Werry et al., 2017; McCormack et al., 2017; Steinaker and Wilson, 2008) and linkages between root function and root dynamics are often poorly understood. As a major function of roots is nutrient uptake, supplying resources which are often limiting, nutrient availability may play an important role in regulating the timing and magnitude of root production. In seasonally arid and Mediterranean systems, plants are thought to be co-limited by N and P (Ries and Shugart, 2008; Sardans et al., 2012; Sardans and Peñuelas, 2013) but there is considerably less information in root responses than commonly measured above-ground may not necessarily translate to seasonal patterns below-ground under both natural and altered experimental conditions. Parameters. Roots may respond in different ways to shoots, particularly under drought (Gargallo-Garriga et al., 2014), and generally, responses below-ground are less consistent than above-ground. This may relate to the balance of co-limitation of nitrogen (N) and phosphorus (P), and water, which are all acquired by roots, and vary in availability throughout the year (Ries and Shugart, 2008). Hence, while N inputs may result in generally higher biomass in Mediterranean grasslands (Dukes et al., 2005), it is unknown to what extent these above-ground patterns are reflected in below-ground development.

Typical of Mediterranean regions, grass phenology typically centres around a summer dormancy, with a dry down in late spring and a ‘green up’ in autumn following the onset of rains. In the more continental interior Mediterranean locations of Iberia (to some extent true in the experimental site used in this study (Luo et al., 2018)), cool winters lead to an additional temperature-driven winter dormancy (Milla et al., 2010; Thompson, 2005). The main growing season is in spring for oaks and pasture species (Oliveira et al., 1994; Orshan, 1989) before an arid summer with senescence of annual pasture species and all the annual components. Roots in such systems are highly spatially segregated (Moore et al., 2005) between herbaceous plants, which dominate the upper 30 cm of soil, and trees, with deep roots which can access water sources through the dry summer. Due to the relatively low canopy cover, high plant diversity, and common use as grazing pasture, microsites may differ substantially in soil properties due to the influence of recalcitrant with trees
having a strong influence due to large inputs of oak litter and nutrient transport from deeper soil layers under trees (Gallardo, 2003). Typically, this litter is more recalcitrant than grasses, tree microsites also tend to have higher nutrient availability (Gallardo et al., 2000). In general, trees also have less tightly coupled above- and below- ground phenology than grasses (Steinaker and Wilson, 2008; Steinaker et al., 2010) due different abilities to store carbohydrates and nutrients over time. A longer lifespan (Abramoff and Finzi, 2015) of root and shoot phenology found that in Mediterranean systems (a very coarse definition including both forests and deserts across only 4 studies), peak root growth tended to lag behind peak shoot growth by over a month on average. Overall, shoots were produced in a peak of during the main (spring) growing season while root production continued through the year. However, there are very few quantitative comparisons of root and shoot phenology in Mediterranean ecosystems, fewer investigating global change factors in treatment experiments, and to our knowledge none in mixed canopy oak savanna-mixed tree-grass savanna systems where both microsite factors and extremes of temperature and water availability may promote desynchronization different responses above- and below- ground.

One reason for the relative lack of information on fine root growth patterns in many systems is the difficulty of sampling an opaque, vertically distributed three-dimensional environment usually only accessible from above. Root biomass typically varies spatially due to resource environment heterogeneity (Hodge, 2004) and in biologically diverse systems, small scale patchiness is increased by individual species with different root habits. While root biomass is a direct measure of root C stocks, it is also relatively inconsistent in response to global change factors (Eitel et al., 2018) and other generally highly inconsistent in experimental responses (e.g. Arnone et al., 2000; Mueller et al., 2018). Other physical attributes (such as spatial distribution of root systems, or traits such as root length density (RLD)) may be more relevant for explaining ecosystem functions plant function and not just standing biomass as they relate to functional properties; for example as root diameter and density vary, RLD relates more directly than biomass to soil exploration. However, all field methods to measure roots have significant downsides (Mancuso, 2012); biomass methods require destructive sampling as roots must be extracted from the soil and cleaned, while visual methods such as minirhizotrons require both pre-installation of observatories, consideration of artefacts and require lengthy and somewhat subjective post-processing.

At our site, we took advantage of a nutrient (+N, +NP) experiment located in a typical Spanish oak-grass dehesa, (described in ?), where fertilization had been applied on an eddy covariance footprint scale (described in El-Madany et al., 2018) to study the effect of nutrient additions on root growth and phenology. If a basic economic analogy for plant resource assignment (Bloom et al., 1985) is applicable, resources are allocated to maximise uptake of limiting nutrients, (Fichtner and Schulze, 1992; Schmidt et al., 2005) As we expected our system to be limited by N (Dukes et al., 2005), when N is added plants should be more limited by P. When N and P were added together, the system should be similar to its original state in terms of the stoichiometry of these nutrients but with more N and P available. A summary of this 'addition-limitation' hypothesis is presented in Figure 1 (panel a).

We used a combination of minirhizotrons (providing strict repeatability and information on root phenology), and soil 'in-growth' cores (measuring root production in root-free soil analogous to recolonization around minirhizotrons) and direct
soil coring (providing direct biomass ground truth). These latter methods were direct measures but highly labor intensive and impractical in drought periods—measurements. We sampled mostly the herbaceous layer roots, which dominated root biomass in the shallow soil 0-40 cm depths we studied (Moreno et al., 2005), although these were also likely to be the pool most variable on a seasonal and interannual basis. In similar systems, most nutrients (Jackson et al., 1988) and root length density (Moreno et al., 2005) is found in surface soils.

We hypothesized that +1.1) fine root biomass followed an annual cycle, developing through a growing season which begins in autumn, and ends with summer drydown—drying the following calendar year—and 2) nutrient and that 1.2) root phenology would be closely synchronized with shoot phenology of the predominantly annual herbaceous layer. For the nutrient additions, we expected that 2.1) nutrient addition of +N and +NP would alter overall root production and 3) nutrient responses to the treatments would be less under trees due to higher intrinsic microsite fertility. If standard resource limitation theory (? was applicable, these nutrient additions. If the site was classically co-limited, we would expect to see larger increases in root mass in +NP than +N. Additionally, 2.2) nutrient addition should decrease overall root biomass relative to shoot biomass (??) as additions of N should relieve the expected site-level N limitation and shift the system towards a P-limited state which would be further relieved by the successive P addition treatment. Considering interacting root-shoot-phenology, we also investigated the null hypothesis that root phenology was closely synchronized with shoot phenology, and indexes of above-ground growth (such as Phenocam imagery—hereafter, phenocam) derived GCC (green colour coordinate) would pair with observed root development if roots were produced at a similar rate between observations as development above-ground: shoot ratios in +NP, maintaining site stoichiometric ratios. With +N, we expected the system to shift into a more P-limited state (otherwise alleviated by the P addition in the +NP treatment, so 2.3) different biomass and RLD effects were expected for +N and +NP additions. These effects were also expected to be 2.4) affected by tree and grass dominated microsites, as higher expected nutrient availabilities under trees could buffer responses to the treatment additions.

2 METHODS

2.1 Study site, treatments and Above-Ground Phenology Nutrient Treatments

We worked at Majadas del Tiétar, a long-term experimental site in Extremadura, Spain (central location: 39°56’25.12” N, 5°46’28.70” W) from December 2016 until May 2018. The site is a typical Spanish dehesa with a low density of oak trees (Quercus ilex (L.)) at ~20 trees ha−1. The herbaceous layer which is dominated by a seasonally changing mixture of species (Perez-Priego et al., 2015) and grazed by cows (<0.3 cows ha−1) during productive seasons which are pastured elsewhere during dry periods. In this study, cows were absent from June 2017 – December 2017. The mean annual temperature is 16°C, mean annual precipitation is ~650 mm, mostly falling between October and April, with a typical Mediterranean climate of long, hot, dry summers and mild, wet winters. The soils are classified as Abruptic Luvisols Dystric Cambisols—an Abruptic Luvisol with a sandy upper layer (~5% clay, 42% silt, 80% sand-75% sand between 0-20 cm), and a clay layer between 30 and 60 cm(2). The site is a typical Spanish ‘Dehesa’ with a low density of oak trees (Quercus ilex (L.)) at ~20 trees ha−1. The herbaceous layer, dominated by a seasonally changing mixture of species (2) is grazed by cows (<0.3 cows ha−1) during

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productive seasons but during dry periods are pastured elsewhere. In this study, cows were absent from May 2017 – December 2017. The site has been an eddy covariance (EC)-instrumented FLUXNET site (‘ES-Lma’) since 2004 and since 2014 has also operated as a stoichiometry–manipulation experiment, three 15 m eddy covariance tower installations (El-Madany et al., 2018) and additions of N with two additional 15 m towers since 2013 (El-Madany et al., 2018) used for a large-scale ecosystem manipulation experiment. The objective of this site-level experiment was to understand ecosystem responses at the EC-footprint (20 hectare) scale to fertilization treatments designed to disrupt ecosystem nutrient stoichiometry. N additions (as Ca-ammonium nitrate fertilizer), and N and P additions (as ammonium nitrate and triple superphosphate fertilizer) were added to one EC footprint each in growing seasons 2014/2015 (100 kg N ha$^{-1}$, 50 kg P ha$^{-1}$) and 2015/2016 (20 kg N ha$^{-1}$, 10 kg P ha$^{-1}$). There are thus three tower treatment footprints (control; the original FLUXNET site with no nutrient additions, N; the nitrogen addition only site, and NP; the nitrogen and phosphorus addition site (?)) resulting in the three fertilization treatments (El-Madany et al., 2018) used in this study; control, +N and +NP.

The overall site-level experiment was designed to study the impact of stoichiometric N:P imbalance, with the nutrient additions providing: standard conditions (control), high N:P expected to develop P limitation (+N), and relieved P limitation (+NP). There was no direct P-only addition but previous publications at the site have demonstrated using a small scale fully factorial experiment a lack of an effect of P alone on vegetation structure and soil properties beyond increases in leaf tissue P and P turnover rates (Perez-Priego et al., 2015; Weiner et al., 2018).

The three tower footprints are instrumented with a variety of standardized instruments including phenocams (Luo et al., in review) instruments to measure large-scale ecosystem properties including phenocams (Luo et al., 2018) using standard Phenocam Network methodology (Sonntag et al., 2012) and tower-mounted SRS Decagon (NDVI) sensors (Sonntag et al., 2012; Richardson et al., 2013), as well as seasonal direct smaller-scale measurements of plant biomass and traits, and traits, including the root measurements reported in this study.

2.2 Root Observatories

We installed root observatories and made root measurements within the site-level footprints. In order to pair these measurements with the site-level design it was necessary to deploy these within the three continuous footprints, each with a different stoichiometry treatment (Figure 1). Given the large footprint of EC measurements and for logistical reasons of finding large enough areas with homogenous land cover and the high cost of instrumentation, EC measurements are typically not spatially replicated. Hence all root measurements reported are subsamples within the three footprint treatments at the site and such point measurements are replicated in space to cover site-level variation within treatments. Ecosystem properties and the results of pre-treatment measures (including soil C, N and P, leaf level nutrients (Table 1), and EC fluxes (El-Madany et al., 2018), indicated that these footprints (25 ha) were comparable before the start of the fertilization experiment.

This design means that replicates of root analyses per treatment as conducted inform about the variation within each footprint area and are therefore strictly pseudo-replicates. This violation of independence was an unavoidable constraint of
the experiment given the very different spatial scales employed at the field site and ambitions for a larger scale synthesis of data across scales. Ecosystem properties and the results of statistical tests of pre-treatment measures indicated that variation was greater within than between treatment footprints before application of fertilizers (Table 1), giving confidence on the robustness that the findings presented in this study are not an artifact of the spatial variability within the experimental area, and so we interpret differences between the three footprints as differences due to the nutrient treatments applied.

We installed 12 minirhizotron observatories per treatment (36 in total) in May 2016 (size: 1 m x 10 cm (l x d), walls of 3mm thick hard plexiglass) used in this study were installed in May 2016 (transparent plexiglass), 8 months before the first measurement. We used 12 observatories in each of ‘control’, ‘N’ and ‘NP’ nutrient treatments, was made. Observatories were arranged in sets of 4 around individual trees, randomly selected from suitable candidates. Two minirhizotrons (MRs) per tree were installed: four around randomly selected individual trees (3 per plot), from around 480 trees per 20 ha treated plot (with a few constraints based on spacing of trees, to ensure unambiguous ‘grassland’ and ‘canopy’ microhabitats were selected and to preserve spacing between sites). The median distance between trees locations within a treatment is 116 metres and 60 meters minimum. At each tree, we installed two minirhizotrons per tree in ‘tree covered’ (TC) locations (halfway between the stem and canopy edge), and two observatories in ‘open pasture’ (OP), at least three times the canopy radius from the stem and no closer to any other tree. This design resulted in six replicates in a nested design per combination of nutrient treatments (Control, N, NP) and habitats (TC, OP), shown in Figure 1, panel b). We detail the statistical implications of this design in the ‘statistical analyses’ section.

Each set of four observatories was installed on a roughly north-south axis to minimize daily variation in solar heating, and all observatories were parallel to the stem. We placed a small bag of silica gel on a piece of high-friction plastic weave within each observatory in order to reduce condensation on the inside of the tube. Despite a tight fit at installation, at first sampling (December 2016), winter soil swelling had moved some observatories in wetter microsites, which we immediately repaired with additional stabilization equipment, resulting in no movement in further campaigns. We excluded these observatories from the first sample, but included them from March 2017 onwards as processed images were not different to other sites. Similarly, two observatories (one in the +N, one in the +NP treatment) were damaged while in the field during the study. These observatories were carefully removed and replaced with a spare tube which was imaged from the subsequent campaign onwards.

2.3 Minirhizotron Method / Sampling Protocol

We used a custom built minirhizotron camera in our observatories modified from other designs (e.g., Amato et al., 2012). This was a visible light camera (FCB-MA 130-FG, JenCam GmBH, Germany) using a 45° mirror to view the outer tube surface with an adjustable handle allowing movement both on the long axis and in defined rotational positions. In the final sampling presented in this paper, this instrument was replaced by a prototype of a new design eliminating the mirror but otherwise producing comparable images. Images were collected using VREO OneView software in .bmp format at 4192 x 3104 pixel resolution and consisted of an image of the mirror surface. The camera was lit with two rows of LED lights on either side of the mirror. We trimmed images to remove image overlap which also removed areas of poor illumination, resulting in a ‘window’ of
observation 5.25 x 4.14 cm per image. We sampled all observatories on seven dates spanning December 2016 to March-May 2018, corresponding roughly to bimonthly measurements in sequential phenology phases of winter, early spring, late spring, late summer drought, winter, and early spring. Each time, we calibrated the camera against a grid of known dimensions fixed to an unburied observatory and collected a complete rotational profile of images (8 total) every 5 cm along the minirhizotron (complete coverage) with image midpoints 0 to 23 cm vertical depth and three additional depths (28, 38 and 44 cm), resulting in 2880 images per sampling date. In December 2016 we only measured two depths, while in May 2017, September 2017, and March 2018, the entire observatory was imaged. We also took a second set of images at all depth positions from the ‘control’ treatment alone in mid-May 2017, 10 days after the complete survey in early May. This was a few days after a major rain event, allowing an observation of the short-term response in fine roots to increased water availability.

2.4 Image Analysis

We rescaled each trimmed image to a standard 1123 x 1434 pixel resolution. After manually filtering images for quality (removing those with image artefacts or ambiguity in cover), a 10 x 13 grid (i.e., 112 by 110 pixel) was superimposed over each image, producing 130 squares. Each square was marked as either ‘roots present’ or ‘roots absent’ following the criteria that a visually unambiguous, apparently living (i.e. with clearly defined form and no obvious structural breaks) root, crossing at least half the square, was present. The proportion of squares with roots per image was calculated (Root Cover Index, RCI). This required 0-2 minutes per image and had significant time advantages over comparative methods which focus on markup of all roots in the image. We validated this method against a standard methodology using the open source minirhizotron interpretation software Rootfly 2.02 (Birchfield and Wells, 2011) to mark up all roots in the images, using 200 randomly selected images from each of the March 2017, May 2017, September 2017 datasets for calibration.

2.5 Direct Soil Measurements

We collected ancillary measurements of root biomass from two non-concurrent methods, sampling within 1 m of the minirhizotron tubes; soil cores (Dec 2016, March 2017) and two rounds of ingrowth cores installed in December 2016 (removed May 2017, December 2017) and December 2017 (removed March 2018, May 2018). These were direct measures of root properties but highly labor intensive and impractical in drought periods. Ingrowth cores were installed by removing a 13 cm, 4.5 cm wide soil column with an auger, homogenizing the soil and removing the live roots and replacing the soil inside a metal 13 cm core with three large root windows. Two ingrowth cores were recovered from each location at each date using a soil auger (i.e., total n = 72). Soil cores were 4.5 cm wide and 30 cm deep in December 2016 (n = 36, one per minirhizotron) but reduced to 20 cm depth (n = 108, 3 per minirhizotron) on subsequent samplings. In this manuscript we only consider the top 13 cm, to pair with root ingrowth cores. The December 2016 Experiment set of ingrowth cores was amended with a separate amount of replacement amount of litter designed to equalize the total root litter in each core to a previously observed seasonal site mean. This was part of a separate isotope labelling experiment not detailed in this paper (Nair et al in prep.).
Root decomposition rates in dehesas are typically very high (Casals et al., 2010) and when combined with a relatively coarse threshold for sieving we judged that these data were acceptable to use along with unamended cores as most root litter would be fragmented by the time of sampling. In all cases, roots were extracted from soil samples by passing through a 2 mm mesh, and picking through the remaining material for intact roots. The extracted root material was cleaned in distilled water and dried at 40°C until weight loss ceased. The final weight was recorded as root biomass.

In addition we made measurements of extractable N and P content in surface soil during the period of the experiment. These were not from the same cores as root measurements but closely paired (< 1 m) at the same sampling location. Two thirds of the minirhizotron and root sampling locations were used with an even distribution of habitats and treatments. Soil from 3, 0-5 cm cores was bulked and sieved (2 mm) and stored overnight at 4 °C. Sub-samples of 20 g were extracted by shaking for 1 hour in 100 mL of 2M KCl (for inorganic N), or 0.5 M NaHCO₃ for (phosphate-P), then filtering the supernatant through Whatman no. 1 (N) and no. 42 (P) filter papers that were preleached with 30 mL distilled water. Extracts were analyzed for ammonium, nitrate, and phosphate using standard colorimetric methods on a Lachat QuickChem 8500 (Lachat Instruments, Hach Company, Loveland CO, USA). Additional 7 g sub-samples were oven dried for 48 hours (until mass loss ceased) at 45°C to determine gravimetric water content.

2.6 Comparisons with site-level and above-ground measurements

Additional to the minirhizotron measurements collected in this study, we also compared root profiles both a) over time and b) with depth to site-level instrumentation. These were i) harvests of above-ground understory biomass made in each treatment area, dried to determine dry weight on 23 March 2017 and 25 May 2017, used to calculate root:shoot ratios (henceforth, RSR) by comparison with the direct biomass measurements made at comparable times and ii) phenocam derived GCC products (Luo et al., in prep.) and grass NDVImeasured from the towers, green colour coordinate (GCC) (Luo et al., 2018) and normalized difference vegetation index (NDVI) of the herbaceous layer measured with an infrared enabled digital camera (StarDot technologies, USA). These two properties are common methods of assessing above-ground phenology from proximal remote sensing. In both cases these above-ground measurements were not directly paired with root measurements, so RSR—root:shoot ratio and phenological synchronicity is examined on the treatment-level. Statistical Analyses

2.7 Statistical Analyses

All statistical analyses were conducted in R (R Core Team, 2018), version 3.50. We—

As previously mentioned, in this study, we assumed that all measurements at individual locations containing paired minirhizotron, root cores, and soil measurements were independent, given the large distance between locations and pretreatment similarity.

We modelled the distribution within treatments as a nested structure, so that the sets of four ‘locations’ (individually either OP or UC) was associated within a particular central tree within a particular nutrient (control, +N or +NP) treatment. Due to the lower number of replicates in soil measurements, we removed this nested term to avoid overfitting the statistical models used for these data. As individual minirhizotrons were always at least 5 m apart and arranged linearly, we expected this spatial
co-variation associated around any particular tree to be minimal. This was especially true between the OP locations, on opposite sides of the tree, where the biggest treatment-level differences tended to be found.

We used the R package lme4 (Bates et al., 2015) to fit mixed effect models and performed a series of linear and generalized-linear regressions, aiming to explain i) relationship between image RCI with image RLD, volume per area or root number in order to validate this fast markup and ii) use observed differences between treatments (control, \( N + N + NP \) and locations TC (tree canopy) and OP (open pasture) to explain root dynamics (both from minirhizotron and direct soil cores) in this system. For the validation of the mark-up methodology we tested different empirical models (linear to polynomial) and we chose the best model for further analyses using BIC model selection, bearing in mind that the granular RCI measurement would always fall within a range of 0-100 so polynomial models with inflection points outside this range could be valid for conversion between RCI and RLD. Data was transformed where appropriate to meet assumptions of normality (Shapiro-Wilks test) and reduce the clumping of low observations—the assumptions of models used, primarily by transformations using Tukey’s ladder of powers using the rcompanion package (Mangiafico, 2018).

We used mixed effects models to understand the effect of nutrient and habitat on both minirhizotron observations (predicted RLD) and soil measurements (root biomass, soil extractable nutrients). In these, we assumed that individual minirhizotrons locations and sampling dates were crossed random effects but did not implement a time series correlation structure due to the large interval between observations and the rapid turnover of species (Fernández-Moya et al., 2011) throughout the year. We performed simple linear model comparisons within individual sampling dates, reducing the models to their most parsimonious form via Akaike Information Criterion (AIC). We tested for appropriateness of including interaction effects between treatment and location by comparing models with and without interaction terms, checked residuals for normality, and report P values for these models using Satterthwaite approximation of degrees of freedom (Satterthwaite, 1946).

3 RESULTS

3.1 Treatment Effects on Soil N:P stoichiometry

During the period of this study there was a strong effect on the ratios of extractable N and P measured in surface soil. Extractable inorganic N content \((2.37 \pm 3.8 \text{ (sd.) } \text{mg g}^{-1} \text{ soil (UC), } 1.79 \pm 3.1 \text{ mg g}^{-1} \text{ (OP), Figure S1})\) was significantly different between habitat types \((P < 0.05)\) but only had a borderline significant effect of \( +N \) \((P = 0.07, \text{ driven by very high N contents in some samples in March 2017})\). Olsen-extractable P \((\text{Figure S2})\) was very different between both treatments and habitat type, with more P available in the \( +NP \) treatment \((P < 0.001)\) and in OP locations \((P < 0.001)\). In the control and \( +N \) treatment, mean extractable P over the period of the experiment was \(3.00 \pm 2.78 \text{ g}^{-1} \text{ in TC locations and } 1.44 \pm 0.9 \text{ g}^{-1} \text{ in OP locations. When P was added in } +NP, \text{ these phosphate-P concentrations were } 7.03 \pm 5.6 \text{ g}^{-1} \text{ in TC locations and } 3.5 \pm 1.54 \text{ g}^{-1} \text{ in OP locations. Overall these differences led to a strong difference in the ratio of bioavailable N and P for the treatment types (P}}\)
<0.001, Figure 2), with a higher (P = 0.06) ratio in +N and a lower (P < 0.001) ratio in +NP than the control. This difference was bigger than the habitat effect (P = 0.08) on available N:P.

3.2 Validation of Minirhizotron Markup

We found a good correlation of our cover-based markup method at our site against compared to all Rootfly-derived indices RLD (Figure ??), RLD, volume per area, and root number were well predicted using the fast cover markup with the best predictive models (3rd order polynomials (RLD, vol area), 2nd order polynomials (root number) using BIC model selection) having $R^2 = 0.77$ (Figure 3) and 0.78, and 0.67 respectively. Particularly high root density caused saturation in RCI but not in Rootfly these measurements, but as this affected only a small number of images in the validation dataset and residuals were otherwise normally distributed, we converted treatment mean root covers RCI to RLD using the observed relationship in all further analyses.

3.3 Depth and seasonal profile from minirhizotrons

Minirhizotron-Derived Root Length Density

RLD from the minirhizotrons-

In general, the minirhizotron images contained less roots in TC than in OP locations (across the whole dataset, P < 0.001, Figure 4). RLD decreased with depth at all periods of the year (Figure 225), with the deepest soil (<40 cm) having a mean RLD of 0.07 ± 0.01 (S.E.) mm mm$^{-2}$ in the most abundant period (May, in contrast to maximum above-ground biomass in March), and 0.02 ± 0.01 in the least abundant period (December). The seasonal cycle of biomass, peaking in March, This peak in deep soil was in May while the overall seasonal cycle (including above-ground biomass, shown later) peaked in March. The seasonal cycle was most evident in the shallow soil surface soil, where maximum RLD was 0.50 ± 0.03 mm mm$^{-2}$ in OP, and 0.40 ± 0.03 mm mm$^{-2}$ in TC. Following March, root biomass declined through May to September (mean 0.05 ± 0.01 mm mm$^{-2}$ in OP, 0.04 ± 0.01 mm mm$^{-2}$ in TC), and stayed relatively constant until December (OP: 0.07 ± 0.01 mm mm$^{-2}$, TC: 0.06 ± 0.01 mm mm$^{-2}$). In general, the minirhizotron images contained less roots in TC than in OP locations (Figure ??).

3.4 (Seasonal) treatment differences from minirhizotrons

Differences between the nutrient treatments were smaller than variation between locations or within time but evident in some periods of the growing season. Minirhizotron calculated RLD tended to be RLD was higher in all measured depths in the +NP treatment plot during the growing season, peaking in the March 2017 sample (Figure 225). Taking the cumulative RLD in the top 13 cm of soil (corresponding to the depth of our ingrowth cores and containing the majority of roots, we compared the treatment x location effect on RLD. Both location (P < 0.005) and +NP treatment (P < 0.005) had significant effects on the RLD calculated from the minirhizotrons over the experiments, but the +N treatment did not differ from control (P = 0.33). Differences between the nutrient treatments were smaller than variation between locations or within time but evident in some periods of the growing season. This difference tended to be greater during the spring as low average RLDs
outside the main growing season limited the potential for variation meant absolute differences between nutrient treatments, if they existed at this time, were impossible to detect using our methodology.

3.4 (Seasonal) differences from direct soil measurements Root Biomass and Root Ingrowth Measurements

The two methods of direct soil measurement (soil cores and ingrowth cores) produced similar results. While these were less frequent than the minirhizotron measurements, they also indicated seasonal changes in roots (Figure 22). Root- indicating a seasonal cycle of root biomass similar to that measured by the minirhizotrons (Figure 6). The top 13 cm root biomass in December 2016 in the top 13 cm (canopy (TC median 2020 kg ha$^{-2}$, grassland OP median 1140 kg ha$^{-2}$) was substantially lower than in the following March (TC median 6390 kg ha$^{-2}$, OP median 5670 kg ha$^{-2}$). In December, while root biomass was low, there were no significant treatment effects but a difference between locations (P < 0.05). As the ecosystem developed into the spring growing period there was a difference in these absolute stocks in March as following transformation to fit the assumptions of linear models, the 2017 (TC median 6390 kg ha$^{-2}$, OP median 5670 kg ha$^{-2}$). Here +N treatment had significantly more (P < 0.05) roots than the +NP and control (although +NP tended to be higher than control cores). This treatment difference was strong enough that the most parsimonious (AIC) model at this date did not include a location effect unlike all other comparisons.

From the ingrowth cores, we also found significant effects. The most parsimonious model for recovery in May 2017 found an effect of both nutrient treatments, where both nutrient amended treatments increased over control (P < 0.05) and also significantly less production in OP compared to TC (P < 0.001) but no interaction. For the year-round ingrowth cores the treatment effect was lost in December 2017 but the highly significant (P < 0.001) location effect remained. Likewise, in 2018, the most parsimonious models showed that in March 2018 (where an interaction term remained in the model), the +N treatment had significantly more roots root biomass in the cores (P < 0.001) and differed between locations (+N-location interaction, P < 0.01). In May 2018, both location (P < 0.01) and both nutrients nutrient treatments (+N, P < 0.05; +NP, P < 0.05) had significant effects. Post-hoc Tukey HSD groupings for linear models for all individual dates for direct soil measurements are shown in Figure 226.

3.5 Short-Term Rain Pulse Responses Pairing of Biomass Dynamics Above- and Comparison with High-Resolution Site Measurements Below-Ground

The short-term minirhizotron measurements in May 2017, separated by 7-10 days around a rain pulse (1 day pre-pulse, 6 days post-pulse) showed a clear proliferation of roots following the pulse (Figure 227). This increase was significant (P < 0.001) in both OP and TC locations and evident in all soil depths measured with the minirhizotron. Similar short-term responses were evident in NDVI and GCC during this period (Figure 228). The relatively sparse distribution of minirhizotron campaigns means we were unable to diagnose similar responses to other rain events although from these site-level above-ground indices the May event was the largest shift against overall trend in the overall trend for the year. From comparison with site indices it is also notable that while the minirhizotron root-cover time series correlates well with both NDVI and GCC in respect to the March
peak and decline into the summer dry period, root cover was not in sync with either of these indices in the autumn growing season where root cover was low in autumn. In both 2016 and 2017 sampling winters, RLD was low but had recovered in both March 2017 and 2018 (Figure 8) by the March of the following year (Figure 8). This indicates that the majority of root growth was in the period of December to March unsampled in either year by the minirhizotron campaigns and after the apparent ‘green up’ of the ecosystem from near-surface remote sensing.

3.6 **Comparison with Above-Ground Biomass Root:Shoot Ratios**

For the two campaigns where above-ground biomass data was available for the vegetative herbaceous layer, even only using the top 13 cm of soil indicated that RSR leaned heavily towards roots root:shoot ratio was very large (in control treatments in March 2017, these were 20:1 (OP) and 15:1 in OP and TC respectively (UC), while in May 2017 these were 21:1 and 22:1). Nutrient treatments showed a typically lower higher ratio in the +N compared to +NP treatment in March but not in grassland in May and generally although by May this difference had been lost from OP locations. Generally, root:shoot were higher in ratio was higher in +N than control but equal or lower in +NP than control (Table 2). Regardless of this potential change in root:shoot ratios, the absolute between treatments, the magnitude of the difference between amount of roots and shoots across all treatments was substantially larger than any changes induced by nutrients in root biomass induced by our nutrient treatments.

4 DISCUSSION

4.1 **Methodological Considerations**

We found effects on RLD and root biomass that differed between treatments in this experiment. Interestingly, the effect of habitat type reversed between RLD (measured from minirhizotrons) and root biomass (measured directly) (Figure 4, 6). However, measurements of roots must always be carefully interpreted (2) to assess ecological meaning (Mancuso, 2012) as all procedures are both affected by methodological biases and time consuming to perform. Notably, we found a different directional effect of habitat type between open pasture and under canopy locations which could possibly be due to methodological limitations. Using our direct methods, we treated all root biomass remaining within the sieve as roots, while the visual MR method allowed roots to be ignored if broken or clearly dead. On the other hand, artefacts due to MR subject to logistical constraints. Hence before discussing nutrient treatments in depth, we will briefly address the interpretative trade-offs between methods.

Minirhizotrons are non-destructive measurements but require the presence of an observatory. Artefacts due to observatory presence are particularly acute close to installation (2). MR equilibrium time can range from a few months (see 2) to years (2) the time of installation (Joslin et al., 2001) and there is little consensus towards an appropriate time to leave observatories before good data is collected. This is especially true in manipulation experiments but measurements during an ‘stabilization’
phase after installation can be assumed to have similar biases between treatments and thus relative differences in this period used to understand effects on root dynamics, if not absolute equilibrium root presence (see Johnson et al., 2001; Mueller et al., 2018; Strand et al., 2018). Slow-growing species may also take considerably longer than this time to equilibrate (Strand et al., 2018) but most (perennial) Q. ilex roots probably reached deeper soil layers than our observations (Moreno et al., 2005), causing both of our methods to mostly sample herbaceous layer roots. In any case, the eight months before first measurements included summer drought and almost total annual mortality of the senescence of this herbaceous layer, followed by autumn rewetting which we expected (as shown in Figure 4). We expected this to have stronger effects than installation on root presence around minirhizotrons. Hence we expected observatory presence on root growth around minirhizotrons and so time since installation to be was unlikely to have impacted the observed trends.

Rapid processing of roots from soil cores does not allow architectural properties to be easily examined in dry systems, as roots are often fragmented during sieving and breaking up of soil clusters. On the other hand, minirhizotron measurements do not alter the position or distribution of roots once they have colonized the area around the tube. Our processing method for minirhizotrons (which was calibrated for our site only) validated well against most of the range of data (Figure 3, $R^2 = 0.77$), with a polynomial fit due to the highest root density being smaller than the resolution of our markup. Using our direct methods, we treated all root biomass remaining within the sieve as roots, while the visual minirhizotron method allowed roots to be ignored if broken or clearly dead. Excluding this possibility of misdiagnosed root biomass in soil cores at different periods of the year (i.e. more root litter later in the season), we found that RLD decreased from March 2017 to May 2017 and root biomass increased slightly. A similar, but smaller difference was observed in 2018. We assume that this difference was due to difference in responses of root traits such as RLD when compared to trends in traits such as root biomass / diameter between treatments and habitats through seasonal changes in weather and water availability.

Similarly, another potential source of errors in our study is the effect of low replication, particularly acute in systems such as Dehesa sites, which are highly diverse and heterogeneous (Moreno, 2008). Below-ground systems additionally cannot be seen before sampling and representative locations may often be assigned based on above-ground properties. We used 6 replicates per treatment vegetation combination for MRs nutrient-vegetation combination for minirhizotrons (36 in total), informed by installation and sampling effort (2 days to fully sample and also lengthy post processing). This level of replication was similar to other multifactorial field experiments using MRs and sites were relatively consistent between measurements. Additionally, with the notable exception of the reversed location effect between MR and soil cores, our treatment trends tended to be consistent between methods (Figure 3, Figure 4) minirhizotrons (Ziter and MacDougall, 2013; Arndal et al., 2017) and, we accounted for consistency in resampling microsites in the statistical models for root data. We hence treat the differences shown by different methods as being both real and ecologically relevant for the rest of this discussion.

4.2 Treatment and Location Effects on Root Measurements Between Methods Effectiveness of Nutrient Treatments

The nutrient treatments used at each site were designed to induce a N:P stoichiometric imbalance with N addition and reduce it by restoring N:P ratios when adding P alongside N (El-Madany et al., 2018). The soil sampling during this experiment (Figure 2, S2,S3) indicated that the intended stoichiometric differences between treatments
were maintained, particularly in the case of +NP, where the increased bioavailable P from the fertilizer (Weiner et al., 2018) was still evident across both treatments and habitat types during the period of this study. Therefore differences in +N and +NP treatments together can be interpreted as N effects and differences in N treatments +N alone can be interpreted as P deficiency, while NP effects +NP effects above control without corresponding N effects indicate co-limitation of both the nutrients together. We expected these. As this study was conducted several years after fertilization, and N is generally more prone to ecosystem losses than P, the overall trend in decreasing ratios in +NP, and less strong increases in +N with our expectations.

4.3 Treatment and Microhabitat Effects

We expected root production (which we measured in terms of RLD and biomass) to differ between habitat types: both habitat types and nutrient treatments at our site. Tree-grass systems, combining short-term and long-term optimality in vegetation habit (Eagleson and Segarra, 1985), usually occur in areas with major seasonal variation in water availability, leading to major variation in soil and understory properties between microhabitats (Moreno et al., 2013). These typically include altered soil water storage (Joffre and Rambal, 1993), and water stress as trees (Joffre et al., 1987). Trees increase shading, allowing reduced transpiration beneath the tree, while extending their root systems to obtain water from both LC-TC and OP locations (Cubera and Moreno, 2007). Additionally, litter input and waste from animals congregating beneath trees result in higher SOC, N soil organic carbon (Howlett et al., 2011), N (Gallardo, 2003) and sometimes P concentrations (Gallardo, 2003; Rolo et al., 2013) beneath trees and in many cases lead to higher herbaceous layer above-ground biomass. In March, this was also observed in our study, alongside an increase in root biomass in soil cores beneath trees. Nevertheless, in May (Moreno, 2008; Rivest et al., 2011). Our site had higher C, N and P contents between tree- and grass-dominated areas (Table 1) which corresponded with higher total above-ground production is frequently higher in open pasture than under canopies, as we found in May (Table 3) biomass in March 2017 (Table 8) and root biomass (Figure 6) throughout the experiment beneath canopies than in pastureland. A positive effect of tree on above-ground yield has been reported in many tree grass systems (e.g. Puerto, 1992; Frost and McDougald, 1989). However in May, we found greater above-ground biomass in OP locations (Table 2) compared to TC locations. This inversion of directional effect, which is probably the due the combination of sustained grazing pressure through the season and increasing water demands into the summer dry-down, microhabitat effect in late Spring above-ground but not belowground in pasture locations was possibly due to differing grazing pressure or plant responses to water availability; the tree-covered microhabitats probably depleted the water in the summer drying faster than in open areas (Moreno and Cubera, 2008; Moreno, 2008), which resulted in more root production for water uptake compared to above-ground growth (and increasing root:shoot ratios as observed in Table 2). This seasonal shift in observed effect highlights the importance of seasonal observations in this system to understand biomass dynamics in such systems.

The same difference in roots between habitat type was not found from mini-rhizotrons, where we consistently observed greater RLD in open pasture throughout the year. The major alteration by tree canopies on environmental properties also affects-Between tree-covered and pasture areas of dehesas there are major differences in herbaceous layer diversity (López-Carrasco et al., 2015) and community vegetation composition (Marañon, 1986), which affects plant trait distribu-
tions. RLD and specific root length are important belowground plant traits linked to plants ability to explore soil and acquire resources (7) and may be (Fort et al., 2014) thus potentially important to competitive success in herbaceous layer communities. Our MR observations indicated a significant increase in RLD in the NP treatment, present in the growing season of both years but strongest during the spring of 2017. However, the direction of the habitat effect difference differed between RLD calculated from minirhizotrons and root biomass measured from cores. This was not a methodological effect as the RCI method fit well with RLD across most of the range of data (figure 1), albeit with a polynomial fit due to high root density within some individual squares. It is important to note that this correlation was a calibration for our site only, and that MR observations may notably not correlate with biomass differences between treatments as our pre-treatment data was available for this study indicated little differences in basic soil properties between habitats (slight differences in texture), surface soil measurements (Table 1, Figures 2, S1, S2) suggest that soil conditions were different in these two habitat types, potentially promoting more RLD-producing species in the open pasture.

Alongside these induced changes, and primarily in the most productive parts of the year, we observed increased RLD in +NP (P < 0.05) compared to the N and control treatments (Figure 4), particularly in open pasture locations. The direct biomass (pooled from direct measurements and ingrowth cores, Figure 6) showed a stronger effect of +N (P < 0.05 over the whole experiment) than +NP on total root mass recovered, particularly under canopies. Biomass and other root traits and properties such as root length may be affected in dissimilar ways by the nutrient and habitat treatments. This is especially important for the uppermost soil layer, where fine roots appeared disappear progressively with soil dryness but remaining roots must leaving remaining roots to grow in diameter to support developing water-harvesting architecture in deeper layer-s. Rapid processing of roots from soil cores does not allow architectural properties to be easily examined, as roots are often fragmented during sieving and breaking up soil clusters whereas MR measurements, while invasive, do not alter the position or distribution of roots once they have colonized the area around the tube. Excluding this possibility of misclassified root biomass in soil cores at different periods of the year (i.e. more root litter later in the season), we expect the difference between minirhizotron RLD and absolute root mass and the different time structure of observations (MR RLD decreased from March 2017 to May 2017 and root biomass increased slightly) to be due to difference in responses of root traits such as RLD when compared to trends in traits such as root diameter between treatments. Stoichiometric changes may provoke layers (and hence, a decrease in RLD in May but an increased or similar biomass, Figure 4, 6). Stoichiometric variation provoke modulation of these responses in terms of root system architecture and traits (2) (Drew, 1975), potentially due to either a shift in species community or in traits themselves (e.g. root diameter). Thus, plants exploring shallow soil in the N-limited N treatment but producing more primary roots in the NP treatment grasslands alongside general increased production (Figure 3) would explain the methodological difference between treatments. Unlike N ions, P is relatively immobile in soil, so in general, P availability has a major effect on root system architecture, with availability promoting primary root growth at the expense of lateral development (2) (Williamson, 2001). Less surface soil exploration is necessary under high availability of relatively-immobile P, reducing the need to access P-rich plant residues (2) and promoting (Lynch, 2011). P addition hence promotes development to reach deeper soil layers and water-soluble nutrients (such as NO3-). Hence the distinct e.g. nitrate. Thus, a proliferation of roots in the topsoil in the P-limited +NP treatment effect from minirhizotrons. N treatment but more
primary roots (and hence, root length from the minirhizotrons) in +NP grasslands alongside a general increased production as a result of the nutrient additions can explain the difference between treatments. The observed difference between +NP and control in RLD, despite the intended difference in stoichiometric conditions may be due to more primary roots under the NP treatment, following the greater loss of added N than P since the start of the experiment. The available N and P (Figure 2) suggests that in +NP, ratios were actually lower than the control (i.e. relatively more P available than N) which may have prompted this increased production of RLD in search of additional N from deeper in the soil.

An increase in primary roots and increased RLD from grasses may be easily detected by the minirhizotrons, as primary roots follow areas of disturbance, previous root channels or soil objects (2) (Rasse and Smucker, 1998) to allow penetration of deeper soil areas. Indeed, we observed a relative shift towards deeper located roots in the MR measurements. Proliferation in surface soil layers already heavily colonized by roots is more difficult to measure using our root presence method. Likewise, our method did not allow an assessment of thickness of individual roots, which may indicate adaption for water uptake. This explanation is supported by the observation from minirhizotrons that roots shifted towards deeper locations late in the growing season (Mar-May 2017, Figure 2) with most roots lost in shallow soils as they dried out 5). Root biomass declined in drying surface soils at the onset of summer. This but production continued in deeper areas, presumably where water was still accessible. Both changes in root diameters and dense proliferation would potentially increase root biomass, as observed in the ingrowth cores, without necessarily changing the RLD observed from the minirhizotrons. This seasonal shift could be in terms of traits or overall species composition and associated traits due to individuals or shifts in species composition; little information on specific species root phenology in Dehesas is available although dehesas is available, but it is clear that both traits relating to root growth seasonal timings and exploratory properties differ substantially between species. In the soil cores, the elevation of both N and NP biomass may be indicative of overall increases in root biomass (including lateral roots) in the N treatment, which are not detected by our minirhizotrons. The methodology allowing us to process large numbers of replicated observations did not allow an assessment of root thickness (potentially leading to exponential increases in mass compared to diameter) and thus potentially, a denser root system under N, supporting foraging for water mobile N ions in deeper soil. Exploratory and water properties can differ substantially both between species (e.g., Luke McCormack et al., 2014; Fitter, 1986) and in time.

A better pairing between difficult to measure root traits and relatively accessible above-ground plant traits may allow a more diagnostic understanding of root behavior under a highly diverse (5) (Moreno et al., 2016) above-ground system and further differentiation between these treatments.

A trait effect of nutrient treatments would also explain the difference observed between OP and TC locations throughout the experiment where directional effects reversed between reversal of directional effects between habitats when measured with minirhizotrons and cores. Similar experiments at our site have previously reported changes in canopy functional traits in response to nutrient addition (5) but are unable to (Migliavacca et al., 2017) but cannot diagnose changes belowground. A greater abundance in ‘high RLD producing’ species in OP, where RLD and soil exploration is a critical competitive trait may have led to a greater response for the minirhizotron observatories. On the other hand, an increase in lateral roots under P deficiency—High RLD is potentially a strong nutrient uptake strategy if roots are cheap (i.e. our +N treatment)
may have had minimal effects on RLD observed compared to the shift in traits when productivity was increased by extra P (in the +NP treatment) restoring stoichiometric ratios. Tree cover has major functional effects on herbaceous layer species (with high specific root length (Hodge, 2004)) and capable of high nutrient uptake. Interestingly, tree cover tends to promote more (relatively high RLD) graminoids compared to over legumes and forbs in dehesas (López-Carrasco et al., 2015), and N abundance (comparatively higher under canopies in dehesa systems (Gallardo, 2003)) tends to shift communities towards grass dominance (?). As nutrient limitation is lower (due to thick community composition towards grasses (Bobbink et al., 2010). However, in heterogeneous environments, high RLD may provide competitive advantages, even if high RLD individuals are less focused on particular nutrient hotspots (Mommer et al., 2011). Hence the relatively more homogeneous environments under canopies with thicker organic layers and relatively high nutrient abundance) in these microhabitats, we may expect an overall weaker response in nutrient-acquiring strategies compared to grasslands when nutrient additions are applied as generally observed in RLD in this study, even if extra nutrient and more abundant background levels of nutrients may mean that nutrient-searching ('high RLD') strategies are less important compared to grassland areas even if nutrient re-cycling through recalcitrant litter means root biomass increases is higher, as observed, in shallow soil."

### 4.4 Changes in Root:Shoot Ratio

Conclusions about the NP treatment a +NP effect on root architecture rather than absolute amount relative to the N treatment an absolute increase in biomass compared to +N are supported by general increases in RSR in the N treatment (Table 2). These shifts in ratio were particularly extreme in the Ntreatment but not the NPtreatment. In NP treatments, RSR mostly decreased, following standard understanding of resource limitation below-ground and decreased whole-plant investment into nutrient acquisition, but ratios did not change so dramatically so this relative decrease in RSR was still a large absolute increase in (biomass-based) root:shoot ratio (Table 2) in +N, but not +NP. Compared to the other treatments, root-shoot ratio decreased in +NP, as expected with a decreasing plant biomass investment below-ground with greater supplies of both nutrients, despite the increasing root biomass. With increased biomass, but 'ambient' stoichiometric ratio of these elements, the plant community appeared to invest less biomass below-ground. The only large increases in RSR were in the Ntreatment root:shoot ratio were in +N, potentially indicating that plants were investing more below-ground producing relatively more roots to alleviate the induced P limitation. Nutrient stress may increase RSR root:shoot ratio if these deficits are the major growth constraint (both (Erikson, 1993; Ågren and Franklin, 2003) and can occur on the ecosystem and individual plant level (7), suggesting (Fichtner and Schulze, 1992). This suggested that while the overall system may have shifted in terms of traits or community in the towards higher RLD species in the +NP treatment, as observed in RLD, the system was more ‘nutrient stressed’ under the Nonly treatment, resulting in greater investment of C below-ground. The difference between control and N treatment RSR was notably +N, as soil N:P diverged from ambient conditions (Figure 2). Notably, from the two dates where we could calculate root:shoot ratio, the difference induced by +N was not found in May in OP open pasture locations. As with the overall biomass responses, this was potentially due to the faster late-season effect of the dry down in more exposed pasture locations areas away from tree canopies, where the herbaceous layer was already in decline.
Generally, the root:shoot ratios (15-30:1) found in this experiment were very large and this was driven by a high root biomass at our site (peaking at around 8000 kg ha$^{-1}$, despite being calculated based on only the top 13 cm of soil), peaking at around 8000 kg ha$^{-1}$. Other studies in similar landscapes in southwestern Europe found maximum root masses of around 2000 kg ha$^{-1}$ (dependent on cover type; Rolo and Moreno, 2012), 2500 kg ha$^{-1}$ (Jongen et al., 2013), or under in nearby walnut forestry, 300-400 kg ha$^{-1}$, despite pasture production being kg ha$^{-1}$ (López-Díaz et al., 2017), where pasture production was around 2 x higher than our site (G. Moreno, pers. comm). Root:shoot ratios at these sites were considerably lower than ours (e.g., 4:1 Jongen et al., 2013) and more consistent with global means (‘temperate grassland’ mean 4.2:1, and ‘savanna’ mean 0.6 (Mokany et al., 2006)) and more consistent with global means (‘temperate grassland’ mean 4.2:1, and ‘savanna’ mean 0.6 (Mokany et al., 2006)). We are confident in the magnitude of root masses reported in this study due to their consistency both over time and between treatments (Figure 4) and RSR in systems such as Dehesas can be shifted in favour of roots by multiple factors such as the (seasonally) dry environment, (8) where root 6. In seasonally dry systems, biomass allocation may shift towards roots due to multiple factors (Chapin et al., 1993). Root distribution in these systems tend to be shallow and wide (9) (Schenk and Jackson, 2002), and grazing of above-ground vegetation (9) which reduces LAI and transpiration, hence favouring nutrient rather than water acquisition, but also requiring as (McNaughton et al., 1998) both reduces transpiration water losses (through reduced leaf area) and requires root foraging for nutrient for nutrients to replacing this removed biomass. As the year progressed towards the summer drought and grazing pressures increased, RSR root:shoot ratio shifted towards roots, and root profiles became more evenly vertically distributed (Figure 25), agreeing with this explanation of the ratios at the site, while a combination of weather and grazing pressure during the year studied may have shifted the balance in towards root production. Ecological effects of seasonal and fertility-related changes are also likely heavily modified by annual variation (Vaughn and Young, 2010) in weather and the particular conditions of the 2017 growing season (a dry year following a particularly productive previous year at the site (Luo et al., 2018)) may have contributed to these high ratios. However, similar root: shoot ratios to ours have been reported earlier in the season (9) (Puerto, 1992) at dehesa sites. More frequent sampling is necessary over multiple years to disentangle whether these ratios are representative of the rest of the growing season.

**Seasonality of Root Biomass and Linkage to Above-Ground Phenology**

### 4.5 Seasonality of Root Biomass and Linkage to Above-Ground Phenology

Our site has a highly seasonal climate with severe deficits of water in summer and an excess in winter (9). As well as a long-term seasonal cycle of root biomass observed from all treatments, in (Perez-Priego et al., 2017) and hence short-term root dynamics are particularly interesting. In the control treatment we observed that root growth responded quickly to a short-term rain event in the late growing season (Figure 57) as RLD increased in all soil depths following a rain pulse in May. This event was paired by a clear response in both NDVI and GCC interrupting the general decline in the late growing season dry down as both shoots and roots responded simultaneously near-simultaneously. However, measurements in autumn implied a considerable desynchronization between above- and below-ground during this time in both 2016 and 2017—the early growing season in both years studied. Most of the root production appeared to occur after the measurement in December and before
the measurement in March, indicating that the key periods of root production were overwinter rather than early or late in the growing season. The initiation of major periods of root growth had not begun by December in either year, while both GCC and NDVI of grassland areas had reached growing season levels by this point in all treatments (Figure 5). This difference was presumably due to high water availability but decreasing light availability in autumn leading to prioritizing of above-ground LAI development, compared to decreasing water availability but abundant light in spring. As both GCC and NDVI are commonly used to track plant phenology (e.g. in a semi-arid grassland, Browning et al., 2017), this was particularly interesting as grassland systems are expected to be relatively highly synchronized above- and below-ground (Steinaker and Wilson, 2008). This difference was substantially larger than the 2-4 weeks observed in other grasslands (Steinaker et al., 2010), and, coupled with the implicit high root:shoot ratio during the later growing season, we observed during the spring, indicates that overall productivity and C status (i.e. the coupling between GPP and RECO) of this system may depend on the coupling between ecosystem-level productivity and respiration. Additionally, unlike the four Mediterranean studies in Abramoff and Finzi (2015)’s meta-analysis, this desynchronisation also indicated that leaf growth was before root growth in our system. This may be linked to the very severe summer drought that our dehesa site experiences, with extensive root systems being more important for water uptake in the late growing season than nutrient uptake for biomass production in the early growing season. On the other hand, both roots and above-ground vegetation responded to a rain pulse the tight coupling of above- and below-ground vegetation in May (during the dry down late in the growing season). This indicated that at some points in the year, these dynamics are highly coupled in the short term. This is likely when water stress is high, short-term events are highly coupled. While our MR, while our minirhizotron data was not high resolution enough to study this at other points, as high resolution at other times, seasonal effects did not appear to be so closely tied to rain pulses above-ground at other points earlier in the year (Figure 6), and ecological effects of seasonal and fertility-related changes are also likely heavily modified by annual variation (Steinaker et al., 2018). At our site, the expectation of root production in direct synchronicity with shoot production is clearly incorrect at least in some times of the year dependent on seasonal and/or climatological conditions. High resolution measurements are necessary to capture such events and advancements in minirhizotron technology including autonomously operational, frequent image capture (matching above-ground proximal remote sensing) and methods to analyze these data, will allow much greater understanding of seasonal cycles in root production and their link to above-ground productivity.

5 Conclusions

Much of our ecosystem-level understanding of plant seasonality and responses to global change is drawn from above-ground measurements but it is not clear how well this understanding holds belowground. In the highly seasonal system of this study, from both minirhizotron and direct soil measurements, we found a seasonal cycle of root dynamics broadly matching both the above ground ‘growing’ season and seasonal patterns of above-ground biomass as inferred from near-surface remote sensed measurements. However there was a notable delay in root production in the early growing season with most root growth not commencing until well after the regreening of the system. Nutrient treatments (N and NP addition)
**N addition** increased root biomass and root shoot ratios in productive periods for understory vegetation, indicating that there was not an effect of altering ecosystem level stoichiometry on root production, and changes were a result of increased N availability. However, NP treatments in grassland areas did show increased RLD over other treatments indicating a herbaceous layer vegetation. +NP (with similar N:P availability to control) showed increases in RLD but not increases in root biomass. This indicates that nutrient availability-driven changes and stoichiometry-driven shift in root traits or plant community following fertilization—changes are not necessarily the same in our system. The expected gradual shift towards P limitation with N deposition could therefore drive increased plant allocation belowground on the community level but also alter traits and function in ways dependent on other environmental conditions, including N:P stoichiometry.

Further work should focus on i) understanding community and plant trait responses to ecosystem stoichiometry, especially in highly diverse and seasonal communities with a large pool of species able to exploit changes and ii) **ii) understanding the consequences of these responses to ecosystem function and iii) increasing temporal density of belowground observations.** Phenology responses to global change factors may be on the scale of days and hence high frequency sampling is essential to understand the belowground response to such forcings and the mechanistic effects of global change on such communities.

*Data availability.*

The authors are eager to share the minirhizotron imagery used in this study with groups working on the critical problem of interpreting minirhizotron imagery. However the large size of the datasets makes permanent hosting of these images difficult. Please contact the authors for distribution of these data.

*Author contributions.* RN designed the experiment, performed the root-associated field, lab and data analysis work, and wrote the manuscript. KM performed all soil nutrient extractions and respective sample collection. MH designed the minirhizotron system. YL provided the processed proximal remote sensed data and assisted with their interpretation. GM provided both above-ground data, and fieldwork assistance. MS and MM helped design the experiment, all authors helped in manuscript preparation.

*Competing interests.* No competing interests are present.
Acknowledgements. This work was funded by a Marie Skłodowska-Curie Individual Fellowship (MsParts) to RN and the via the Alexander von Humboldt Foundation Max Planck Research Prize (Markus Reichstein to MR). We also acknowledge the invaluable field support of Kendal Morris, Enrique Juarez, Andrew Durso, and Jinhong Guan as well as Mónica Lorente, Fatima Khalid, and Rolf Rödiger who assisted in laboratory processing of root samples. We are also especially grateful to Gianluca Fillipa for sharing his expertise with interactive GUIs for graphics processing in R.
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Ziter, C. and Macdougall, A. S.: Nutrients and Defoliation Increase Soil Carbon Inputs in Grassland, Ecology, 94, 106–116,
Figure 1. Schematic diagram of experiment. In panel a), we show conceptual release from N limitation in our +N treatment but promotion of P limitation which is alleviated by NP additions. In panel a) we show 12 minirhizotrons per treatment, split between three trees into ‘OP’ (open pasture) and ‘UC’ (under canopy) areas. This is a small subset of the total trees in each footprint. Median distance between trees within treatment is 116 metres and minimum distance between individual minirhizotrons is 5 metres.
Figure 2. Ratio of extractable inorganic N (2M KCL) to extractable P (Olsen method) in 0-5 cm soil during the experiment. Over all measurements, the +N treatment has a borderline significant ($P = 0.06$) greater N:P ratio and the +NP treatment a significantly lower ($P < 0.001$) ratio than control. Letters show Tukey HSD groupings within campaign-habitat combinations and errorbars show standard error of the mean.
Figure 3. Root Cover Index (RCI) against root length density (RLD) for a random sample of images (n = 300) from three imaging sessions campaigns at our site. A simple model 3rd order polynomial (indicated) fit to these data with an $R^2$ of 0.77 was used to predict RLD for all other figures.
Figure 4. Seasonal cycle of minirhizotron-calculated root length density (RLD) between treatment at two selected depths (vertical position of camera). +NP treatments tended to diverge, especially in spring 2017. Error bars show value ± SE. A mixed effect model revealed that these data revealed a significant effect of the +NP treatment ($P < 0.005$) but N did not differ from control on this metric.

Vertical root profiles (mean of minirhizotron measurements) for four dates in 2017. NP treatments tended to diverge in the grassland, especially in shallow soil where RLD was highest. Treatment effects were most evident in the shallow soil depths where root biomass was highest. As accurately annotating minirhizotron measurements is very time consuming, the full depths for September and December 2017 were not fully analysed for this graph.
Figure 5. **Root Vertical root profiles (mean of minirhizotron measurements) for the four sampling dates in 2017, representing one complete annual cycle (peak RLD in March of this year was also higher than 2018).** NP treatments tended to diverge in the grassland, especially in surface soils where RLD was highest. Treatment effects were most evident in the shallow soil depths where root biomass was highest.
Figure 6. Fine root biomass in top 13 cm from direct soil cores (Nov. 16, Nov. 17) and ingrowth cores (installed Dec. 2016 and removed then in Mar. 17, Nov. 17, then installed Nov. 17, removed Mar. 18, May 18) and direct soil cores to obtain standing living root biomass during the sampling period. Direct measured root biomass showed a seasonal cycle and also differences between treatments, with more roots in general under canopies and in fertilized plots. Error bars show value ± SE and letters indicate Tukey-HSD groupings for most parsimonious models within treatment for both Open Pasture (OP) and Under Canopy Tree Covered (TC) locations combined in individual sessions. Both +N and +NP treatments tended to have more root biomass than control treatments. Across the whole dataset, +N had significantly (P < 0.05) more fine roots than the other two treatments.
Figure 7. Control Treatment root length density (RLD) response to rain pulse in May 2017 (Section ??). Repeat MR sampling showed increases in 2017. RLD throughout the soil profile (P < 0.001 for both locations) indicating short-term proliferation of root growth.
Comparison of minirhizotron root length density (RLD, mm mm⁻²) dynamics for open pasture with site-level grassland Normalized Difference Vegetation Index (NDVI) and grassland Greenness Colour Coordinate (GCC). After the rain pulse in May 2017 (indicated on graph, light blue), minirhizotron measurements could detect the similar response to above-ground (shown in more detail figure ??). Desynchronization was evident in both autumn periods where proximal-remote sensed metrics reached near-peak levels while RLD remained low.

Figure 8. Comparison of minirhizotron root length density (RLD, mm mm⁻²) dynamics at 9 cm depth for open pasture with site-level grassland, Normalized Difference Vegetation Index (NDVI) and grassland Greenness Colour Coordinate (GCC). After the rain pulse in May 2017 (indicated on graph, light blue dashes), minirhizotron measurements could detect the similar response to above-ground (shown in more detail figure 7). Desynchronization was evident in both autumn periods where proximal-remote sensed metrics reached near-peak levels while RLD remained low.
Table 1. Summary of pretreatment measures within EC footprints ('treatments'). All measurements were made in Nov 2013 for soil and in spring 2014 for plants. All soil properties are measured on surface (0-5 cm) soil and open pasture leaf N:P are a biomass weighted aggregate of samples of grass and forbs (98% of herbaceous layer biomass). All errors are standard deviations. Soil nutrient contents (C,N, and Olsen-P) were always significantly higher in UC than OP locations (P < 0.005, P < 0.005, P < 0.05 respectively). There were no significant effects due to footprint except soil pH (*, p < 0.05) but this did not result in a difference of weighted mean vegetative N:P ratios, so we considered this unimportant for the overall design.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Control</th>
<th>+N</th>
<th>+NP</th>
<th>Control</th>
<th>+N</th>
<th>+NP</th>
</tr>
</thead>
<tbody>
<tr>
<td>C (mg g⁻¹)</td>
<td>4.87 ± 1.4</td>
<td>5.33 ± 1.2</td>
<td>6.56 ± 1.6</td>
<td>13.59 ± 0.5</td>
<td>10.06 ± 1.2</td>
<td>12.57 ± 3.8</td>
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<tr>
<td>N (mg g⁻¹)</td>
<td>0.86 ± 0.1</td>
<td>0.93 ± 0.1</td>
<td>1.04 ± 0.1</td>
<td>1.54 ± 0.1</td>
<td>1.30 ± 0.2</td>
<td>1.49 ± 0.4</td>
</tr>
<tr>
<td>Soil C:N</td>
<td>5.7:1</td>
<td>5.7:1</td>
<td>6.3:1</td>
<td>8.9:1</td>
<td>7.8:1</td>
<td>8.4:1</td>
</tr>
<tr>
<td>Olsen-Extract P (µg g⁻¹)</td>
<td>2.3 ± 0.6</td>
<td>3.68 ± 1.4</td>
<td>3.38 ± 2.0</td>
<td>5.44 ± 0.9</td>
<td>5.45 ± 3.8</td>
<td>3.77 ± 1.4</td>
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<tr>
<td>Soil pH</td>
<td>5.42 ± 0.1</td>
<td>5.56 ± 0.5</td>
<td>4.93 ± 0.3 *</td>
<td>5.50 ± 0.6</td>
<td>5.58 ± 0.4</td>
<td>4.93 ± 0.3 *</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Control</th>
<th>+N</th>
<th>+NP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Herbaceous Layer Leaf N:P</td>
<td>13:1</td>
<td>13:1</td>
<td>13:1</td>
</tr>
</tbody>
</table>
Table 2. Absolute (Herbaceous layer) Biomass Measurements and Root:Shoot Ratios at two points in 2017. Root shoot ratio increased later in the growing season but tended to decrease in nutrient added treatments, with the exception of N:OP in March and N:TC in May which exhibited unusually high root:shoot ratio.

**March 2017**

<table>
<thead>
<tr>
<th>Nutrient</th>
<th>Vegetation Aboveground Biomass</th>
<th>Belowground Biomass</th>
<th>Root:Shoot Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control TC</td>
<td>290 ± 20</td>
<td>5770 ± 320</td>
<td>20 ± 2 : 1</td>
</tr>
<tr>
<td>Control OP</td>
<td>290 ± 20</td>
<td>4510 ± 250</td>
<td>15 ± 1 : 1</td>
</tr>
<tr>
<td>N TC</td>
<td>340 ± 20</td>
<td>6890 ± 390</td>
<td>21 ± 2 : 1</td>
</tr>
<tr>
<td>N OP</td>
<td>290 ± 20</td>
<td>7590 ± 420</td>
<td>26 ± 3 : 1</td>
</tr>
<tr>
<td>NP TC</td>
<td>380 ± 30</td>
<td>6380 ± 350</td>
<td>17 ± 2 : 1</td>
</tr>
<tr>
<td>NP OP</td>
<td>300 ± 30</td>
<td>4900 ± 270</td>
<td>16 ± 2 : 1</td>
</tr>
</tbody>
</table>

**May 2017**

<table>
<thead>
<tr>
<th>Nutrient</th>
<th>Vegetation Aboveground Biomass</th>
<th>Belowground Biomass</th>
<th>Root:Shoot Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control TC</td>
<td>260 ± 40</td>
<td>6900 ± 700</td>
<td>26 ± 5 : 1</td>
</tr>
<tr>
<td>Control OP</td>
<td>370 ± 60</td>
<td>4280 ± 330</td>
<td>12 ± 2 : 1</td>
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<tr>
<td>N TC</td>
<td>220 ± 30</td>
<td>8250 ± 640</td>
<td>38 ± 6 : 1</td>
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<tr>
<td>N OP</td>
<td>400 ± 50</td>
<td>5960 ± 580</td>
<td>15 ± 2 : 1</td>
</tr>
<tr>
<td>NP TC</td>
<td>360 ± 40</td>
<td>7850 ± 900</td>
<td>22 ± 3 : 1</td>
</tr>
<tr>
<td>NP OP</td>
<td>440 ± 30</td>
<td>6670 ± 700</td>
<td>15 ± 2 : 1</td>
</tr>
</tbody>
</table>