Metabolic activity of *Hordeum vulgare*, *Brassica napus* and *Vicia faba* in Worm and Root type Biopore Sheaths

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**Abstract**

Aims Biopores offer favorable chemical, biological and physical properties for root growth in untilled soil layers. There they are considered as nutrient “hot-spots” with preferential root growth. However, the literature lacks a quantification of metabolic activity due to nutrient acquisition of main crops while growing in the biopore sheath.

Methods A pot experiment was performed to map the metabolic activity of roots, as indicated by pH change. The roots of spring barley (*Hordeum vulgare* L.), spring oilseed rape (*Brassica napus* L.) and faba bean (*Vicia faba* L.) were growing through the biopore sheath influenced by an earthworm (*Lumbricus terrestris* L.) or a taproot (*Cichorium intybus* L.), in comparison to subsoil without a pore (bulk soil). pH sensitive planar optodes were applied in order to image a planar section of the sheath, while preserving an intact biopore sheath during the experiment.

Results Roots were first found in the field of view in worm biopore then root biopore and bulk soil. At time of the first measurement the pH value was highest in worm biopore sheath (LS-Mean±SEM: 7.16a±0.11), followed by root biopore sheath (6.99ab±0.12) and bulk soil (6.61b±0.12). In spring oilseed rape a significant alkalization (+0.80 Δ pH) was found over time in bulk soil. Faba bean significantly acidified the root biopore sheath (-0.73 Δ pH). Spring barley showed no significant pH changes.

Conclusions The results of the current study reveal a trend of faster root growth through biopores and a higher initial pH value in the biopore sheaths compared to the bulk soil. Biopores serve not only as an elongation path for roots, but their sheaths also provide an environment for root activity in the subsoil.

**Keywords** biopore sheath · non-invasive imaging · pH planar optodes · metabolic activity · rhizosphere · root-soil interaction

**Introduction**

Biopores are continuous large pores that can be found in compacted soil and are mainly formed by roots and earthworms. Root growth and earthworm activity result in different chemical, biological (Tiunov and Scheu 1999; Uksa et al. 2015) and physical...
properties (Pagenkemper et al. 2015) of biopores and their sheath. Root growth in un-tilled deeper soil layers can be facilitated by physical access paths through biopores (Passioura 1991; Logsdon and Linden 1992). Biopore surroundings provide elevated nutrient concentrations, particularly in the first two millimeters of the sheath due to earthworm linings and root debris (Lavelle 1988; Athmann et al. 2017; Petzoldt et al. 2020). Generally, crops can respond to different nutrient availabilities in their root growing environment by changing their metabolic activity as reviewed by, e.g. Hinsinger et al. (2009), Hodge et al. (2009), Walter et al. (2009). Therefore it can be assumed, that the metabolic activities of the crop species differ in heterogeneous soil structures, such as worm or root type biopore sheaths and bulk soil, due to the specific environmental conditions.

Root growth in the biopore sheath in subsoil can be considered as relevant to overall nutrient acquisition, particularly in low input cropping systems. Substantial amounts of nutrients originate from subsoil, e.g. the subsoil contribution to the total N uptake of winter wheat was reported to be on average 33% across 22 different sites, and even 75% when mineral fertilization to the topsoil was omitted (Kuhlmann et al. 1989). Athmann et al. (2016) showed in a 15N tracer field study that the quantity of N derived from the subsoil amounted to at least about 40% for mallow and 32% for wheat. Positive correlations were found between the number of biopores and N uptake of wheat (Volkmar 1996). However, it has not yet been demonstrated that roots growing through biopores effectively take up nutrients from the biopore sheath and that the inner layer of the biopore sheaths are truly relevant “hotspots” for nutrient acquisition. In fact, nutrient uptake from the biopore sheath can be limited by lack of root-soil contact and restricted root growth due to high bulk densities in the biopore sheath. Beneficial effects of biopores can also be based on provision of preferential pathways for root elongation and general facilitation of root growth.

It is well documented that plant roots respond to the environment they grow in, e.g. soil structure, nutrient availability, acidic or calcareous soils (Schubert et al. 1990) by actively altering the biochemical transfer in their rhizosphere (Hinsinger et al. 2003; Hinsinger et al. 2009; Hodge et al. 2009). In order to mobilize and forage on nutrients, the rhizosphere pH value varies along the root axis (Marschner et al. 1986). Hence, pH alterations can arise temporal and are dependent on root age and order (Hinsinger et al. 2009; Rudolph-Mohr et al. 2014). Besides this, heterogeneity in pH values in soils can occur due to organic material originated by diverse flora and fauna, such as earthworms or decaying plant roots. Earthworm casts were found with increased pH values in acidic soils (Tiunov et al. 2001; Horn et al. 2003), and characterized with higher contents of available nutrients, as nitrate and ammonia (Parkin and Berry 1999), or available phosphate, sulfur, enzyme activity and microbial biomass (Athmann et al. 2017). These advantageous properties were also found in root pores, but with lower contents or different microbial communities compared to worm pore, yet higher contents and activities than in bulk soil (Athmann et al. 2017). It can be suggested that the form of the nitrogen influences the metabolic activity of crop species in worm pores, root pores and bulk soil. Earthworm casts, older than two weeks, are reported to be enriched in nitrate (Parkin and Berry 1999), while the experimental bulk soil in this study contains clay minerals with non-exchangeable ammonia. For instance, nitrate leads to physiological alkalization in the rhizosphere and non-exchangeable ammonia to acidification. The chosen crop species in the current study have diverse strategies to acquire nitrogen. Faba bean is nitrogen independent since N2-fixing-symbiosis is established, whereas oilseed rape can mobilize, e.g. non-exchangeable ammonia more successfully by acidification than barley, due to a higher potassium-ion uptake and proton release (Beuters et al. 2014).

Beyond that, non-invasive techniques with pH sensitive planar optodes have previously been reported as having been successfully deployed for observing pH alteration in the rhizosphere on single roots of an individual (e.g. Blossfeld et al. 2013; Rudolph-Mohr et al. 2017). It is possible to quantify the proton-fluxes via colored pH indicator precisely with a relative error of 0.03 pH units (Jaillard et al. 1996). Also mapping differences in pH value along root axis and the influence of the form of N on the proton release were shown (Plassard et al. 1999), as well as the exchange of macro- and trace elements in flooded rice land (Williams et al. 2014). Yet, to the author’s knowledge, there are scarcely articles, where pH sensitive planar optodes were performed in a root-soil interaction experiment with several treatments and replicates and there is no study focused on structured
subsoil of arable field. For instance, a related main experiment was performed by Kreuzeder et al. (2018), who observed the pH change induced by wheat, buckwheat and lupine after ammonia or nitrate supply in a calcareous and non-calcareous soil. A recent article using such technique showed in a main experiment the spatial heterogeneity along roots of lupine from apical to basal in case of P mobilization accompanying different pH values (Ma et al. 2021). Other related studies using pH sensitive planar optodes observing burrow walls were however carried out in marine sediment (Hulth et al. 2002; Stahl et al. 2006).

Considering this, the current study focuses on measuring dynamic pH changes induced by root activity of three crop species. It is assumed that the roots, in turn, are influenced by biologically formed soil structures, which is in this study related to biopores formed by taproot or earthworm. It is hypothesized, that roots of spring barley, spring oilseed rape and faba bean i) induce pH change in biopore sheaths, and ii) a higher pH change is expected for the worm than for the root biopore sheath and bulk soil.

Material and methods

Experimental design

For each of the three species, spring barley (Hordeum vulgare L.), spring oilseed rape (Bassica napus L.) and faba bean (Vicia faba L.), the following three treatments were tested: (1) “Worm type biopore”: column with one worm pore formed by anecic earthworm (Lumbricus terrestris L.), (2) “Root type biopore”: column with one root pore formed by chicory (Cichorium intybus L.), and (3) “Bulk soil”: column with subsoil without a pore.

Per treatment four columns as replicates were prepared, a total of 36 columns were randomized together, resulting in a completely randomized design. The preparation of the soil columns in brief: plastic pipe columns (diameter 10 cm, height 50 cm) were filled and repacked (soil density 1.5 g cm⁻³) with air-dried, sieved (<2 mm) and homogenized subsoil of a haplic luvisol (see below). In the center of each column of treatment (1) and (2) a pore with a diameter of 6 mm was formed with an iron rod. Then one adult earthworm was inserted and removed after two months. For the root type biopore one seedling of chicory was planted as precrop and cultivated for three months. The decaying time lasted ten months and large portions of the taproot were removed from the pore before main crop cultivation (further details in Petzoldt and Kautz 2021). Before the main cultivation started, the rhizotrones were built by cutting the subsoil columns vertically into a 40 cm high section. The rhizotrone was built with a subsoil layer of 30 cm and above a topsoil layer of 10 cm (Fig. 1). The experimental soil texture of the topsoil is classified as silty loam with 15-20% clay, 74-77% silt and 5-8% sand and the subsoil as silty clay with 25-30% clay, 65-70% silt and 4-5% sand (details of the reference soil profile in Pätzold et al. 2006). A vertical half was tightly sealed with a plexiglas panel with a pH sensitive planar optode (Type SF-HP5R, PreSens, Regensburg, Germany) at the inside. The optode was placed at a soil depth of 30-34 cm at time of sowing the main crops (Fig. 1). The plexiglas side was darkened with a fleece, tightened with armor tape at the sides, and three iron clamps per column. The main crop cultivation lasted from the 16th of March to the 5th of May 2019. Thus, germinated spring barley ‘Barke’, spring oilseed rape ‘Ability’ and faba bean ‘Fanfare’ were
cultivated for seven weeks. During the cultivation, the glass faced columns were mounted at an angle of 20° to allow root growth at the soil-glass interface. The rhizotrones were irrigated with deionized water daily from the top and once a week from the bottom. Therefore the bottom was kept closed with a plastic cap within a moist layer of cotton. When the experiment was finished, spring oilseed rape was in developmental code (DC) 15-17, spring barley in DC 39-50 and faba bean in DC 50-60 (Meier et al. 2009).

The planar optode had a size of 4 x 4 cm and the measurement field of view amounted 3.5 x 3.2 cm. Fluorescent molecules were embedded in the planar optode, emitting a characteristic pattern of red, green and blue light. The ratiometric approach allows the detection of changes in the ratio between green and red color (R values) caused by acidification or alkalinization. The variation is detectable by a detector camera (Principle of this planar optode technique, e.g. Blossfeld and Gansert 2007; Blossfeld et al. 2013). Their pH range was 5.5-7.5. The planar optodes had been soaked and equilibrated in a buffer solution for 4 h before fixation at the plexiglas.

Calibration

A calibration was done in the experimental set-up at the beginning and the end of the experiment by using a specific tool called “CaliPlate” (PreSens, Regensburg, Germany), which contains the SF-HP5R sensor foils in a 10-well plate. Six wells were filled with a buffer solution of different pH values from 4.5 to 9. Two stock solutions of phosphate buffers with a buffer capacity of 40 mM and an ionic strength of 140 mM were used for the calibration: Solution A: 5.5 g NaH₂PO₄·H₂O + 5.8 g NaCl, Solution B: 7.1 g Na₂HPO₄·2 H₂O + 1.2 g NaCl. The buffer solutions were prepared by mixing different volumes of the stock solutions.

Determination of pH values

The columns were controlled daily, and if it was apparent that a root had grown into the field of view, measurements were taken with a 2D-detector unit USB-Microscope device (PreSens, Regensburg, Germany) with 470 nm wavelength LED light, an exposure time of 0.2 ms and the fluorescence images had a resolution of 0.2 mm per pixel. The microscope is sensitive within the emission range of the optode. The measurements were calibrated with the imaging software VisiSens AnalytiCal VA 2.11 (PreSens, Regensburg, Germany). The calibrated images were further processed with the open source software Fiji, Image J (Schindelin et al. 2012). All measurements of a rhizotron were bound into a stack, so that the regions of interest (ROI) had a defined size (20 x 2 mm) besides the pore (Figure 2), and were fitted to the pore diameter. The pore diameters ranged between 6 and 10 mm. The pH value was determined as the mean value from 2 consecutive measurements within 10 minutes. ROI’s were defined, because roots especially of spring barley and oilseed rape were usually not visible through the planar optode during the experiment. Differences in pH values were determined as mean values with parts of roots, rhizosphere and soil. However, visual inspection of the area behind the...
planar optode after the experiment revealed, that a ROI devoid of roots rarely occurred.

Determination of total C- and N-content

After finishing the pH measurements, soil samples were taken in a soil depth of 25-35 cm in three lateral distances from pore surface: 0-2, 2-4 and 4-8 mm for analyzing the total content of C and N ($C_t$, $N_t$). The scraping sampling device and the proceeding of the samples, as well as the measurement of the elements are described in detail in Petzoldt et al. 2020.

Statistical analysis

The three treatments were evaluated at the same number of days after a root appeared behind the planar optode as "Day after root entry (DAE)", separately for each species. The root entry was completed when root(s) were grown into the field of view.

First, the treatments were compared in the number of days until first root appearance behind the planar optode within the field of view (DAE=0), using a one-factorial model with pore type as fixed treatment factor. The same procedure was done to compare the pH values of the treatments at DAE=0, without separating the species to refer to the treatment effect.

The main focus was on measuring the intensity ($\Delta$ pH) of the metabolic activity of the plants. The change in pH ($\Delta$ pH) was calculated as difference between the pH value of a subsequent point in time to the pH value on DAE=0 (first appearance of the root). The calculated $\Delta$ pH were related to the following time intervals: 1-2, 3-4, 5-9, 10-14, 15-19 day after root entry. For the analysis of $\Delta$ pH, we used a mixed model for split-plot design with pore type as fixed treatment factor and DAE as fixed repeated factor. The rhizotrones served as whole plot for the repeated measurements in time, this resulted in an additional random error term (pore type x column) for whole-plot. For faba bean and barley, heterogeneous variances for DAE were additionally considered due to better model fit. Since our greenhouse did not allow precise temperature control, the temperature at the time of a pH measurement was included in the model as a co-factor (temperature range: 26±5 °C), with the result of no statistical effect on $\Delta$ pH response.

The $C_t$- and $N_t$-content were analyzed, separately for species, using a mixed model for split-plot design with pore type and lateral distance (0-2, 2-4 and 4-8 mm) from each macropore as two fixed treatment factors. The rhizotrones served as whole plot for the lateral measurements leading to pore type x column effect as additional random whole plot error term.

Pairwise comparisons between Least Square Means were made by t-Test with Bonferroni correction to account for the multiple testing situation ($\alpha=0.05$). Statistical analysis was performed using MIXED procedures by SAS 9.4 software (SAS Institute Inc., Cary, NC, USA). The figures were done using the software package Microsoft Office 2013 (Microsoft Cooperation, Redmond, WA, USA).

Results

Root growth, nodulation, pore diameter and initial pH value

The roots of faba bean reached the field of view first in pots with worm pores, followed by pots with root pores and bulk soil (Figure 3a). The diameter of the worm pore was significantly larger than the root pore, however both pores were several times larger than the root diameters (Figure 3a, Supplement 1). Overall, at the time, when spring oilseed rape reached the planar optode, a comparatively fast root growth with the formation of plenty of roots was observed. That resulted in a comparatively small number of measurements. In the case of the bulk soil treatment in faba bean less measurements were taken due to delayed root growth until the experiment had to be stopped.

The average root diameter of spring oilseed rape (range: 0.19-0.37 mm) and faba bean (range: 0.39-0.60 mm) did not differ between the treatments, as well as between the biopore and biopore sheath in the worm or root type treatment (Supplement 1; no data available for spring barley).

In faba bean there was a significantly higher nodulation in the topsoil than in subsoil in worm type and bulk soil treatment, which was equal in root type treatment (Supplement 2). At the end of the experiment the number of nodules pot$^{-1}$ in the field of view was 5.00±1.21 (LS-Mean±SEM) in root type, 0.75±1.21 in worm type and 0±0 in bulk soil treatment.

The pH value of the treatment at time of the first measurement (DAE=0) were decreasing in the
following order: worm biopore sheath (pH 7.16) > root biopore sheath (pH 6.99) > bulk soil (pH 6.61) with a significant difference between worm type and bulk soil (Figure 3b).

**Metabolic activity as indicated by pH change**

In spring barley, the Δ pH varied between -0.19 and +0.32 in the bulk soil, -0.10 and +0.01 in worm type pore and +0.05 and +0.29 in root type pore (Figure 4). In spring oilseed rape, the Δ pH raised in root pore from time interval 1-2 to 5-9 DAE with 0.67 unit (p-value = 0.062). In the bulk soil a significant increase was found since time interval 1-2 to 5-9 DAE with 0.80 pH unit. While in worm pore sheath a maximum was reached at the time interval 1-2 DAE with 0.22 Δ pH. In faba bean, a significant difference was found in root type treatment over time from the time interval 1-2 to 15-19 DAE, with a strong decrease of -0.95 until 10-14 DAE or -1.05 Δ pH unit until 15-19 DAE. In worm biopore sheath the pH change reached its maximum in 0.33 Δ pH at time interval 3-4 DAE and in bulk soil with 0.29 Δ pH at time interval 1-2 DAE. In the three crops, the Δ pH was comparatively small in the worm pore sheath, whereas in root pore sheath and bulk soil higher Δ pH were found. The observed pH values for all measuring days can be found in Supplement 3.

**C₅- and N₅-content in the biopore sheath**

For spring oilseed rape and faba bean the C₅- and N₅-content in the biopore sheath of different lateral distances were measured (Figure 5). A significantly higher C₅-content was found in 0-2 mm than in 4-8 mm lateral distance from macropore in the worm type sheath in spring oilseed rape. In faba bean a significant increase was found in the C₅- and N₅-content in worm type sheath of 0-2 mm. No differences were found in root type sheath for both elements and crops. Data for spring barley is not available.

**Discussion**

Roots of an individual plant grow through different structures along a soil profile with temporal and spatial heterogeneity. While growing they adapt the growing path, e.g. due to nutrient attraction or to bypass physical impedance (e.g. Correa et al. 2019). In the current study faster root growth, in terms of reaching the planar optode, was observed for worm than for root type pore and least for bulk soil, particularly in faba bean. Faster root growth through biopores, also reported by Gaiser et al. (2013), especially during dry spells (Gaiser et al. 2012) can be facilitated by unimpeded physical access pathway
Even if large-sized biopore allowed for unimpeded root elongation, there was still a comparatively high variance between the individuals, which pointed out the heterogeneous nature of root growth. Moreover, in this experiment faba bean roots were comparatively early in time and highly present in the biopores, which is contrary to other results (Athmann et al. 2019a; Petzoldt and Kautz 2021). That can be caused by the inclined position of the rhizotrone, the placement of the seedling above the pore and faba as a species, which was found to have less root plasticity than graminaceous species (Li et al. 2014). In addition, for the bulk soil, it cannot be ruled out that the construction of the rhizotrones had an influence on the delayed root growth, since thickened faba bean roots were observed partly. However, there was no significant difference in the average root diameter between the treatments (Supplement 1). Contrary to faba bean, there were no significant differences in the elongation rate between biopores and bulk soil in spring oilseed rape, even though oilseed rape showed an attraction to biopores in a previous study (Athmann et al. 2019a). This could also be caused by the rhizotrone, due to small spaces between the bulk soil and plexiglas, which could have influenced the root growth in the bulk soil treatment. However, it appeared that roots were more attracted by worm than by root type biopores. This can be a result of the larger sized worm pores and the highly available nutrients in earthworm casts, rather than in root debris or bulk soil (Athmann et al. 2017). Hence, biopores, particularly coated with worm linings, supported faster root elongation towards deeper soil layers with the potential of a more rapid access to resources in the subsoil.

Comparatively higher microbial activity and nutrient availabilities for worm pore sheath, but also for root pore sheath in comparison to bulk soil (Athmann et al. 2017) was considered facilitating higher metabolic activity, as observed by pH change as indicator. However, in all three crops in the worm type sheath the pH value varied at the slight alkaline pH range within the first 20 days after root entry, without distinct effects in alteration over time. Although, in the root type and bulk soil treatment the intensity of the metabolic activity differed between the three experimental crop species. Homorhizous spring barley showed slight pH changes in the biopore sheaths and bulk soil, whereas both allorhizous crops showed a distinct pH change in root type sheath, as well as spring oilseed rape in the bulk soil. This can be due to a higher ability of oilseed rape and faba bean to alter the pH value (Beuters et al. 2014; Schubert et al. 1990). In spring oilseed rape an alkalization effect over time was found in root pore sheath and bulk soil, which had a neutral to slightly acidic initial pH value. While faba bean showed the same pattern of pH change in all treatments until day 9 after root entry. Thus, young faba bean roots slightly increased the pH value until day 4 after root entry followed by
a decrease. The same pattern of the pH change of first slightly increasing and then decreasing, was previously found for white clover (Marschner et al. 1986). The decrease was particularly found for root type biopore with a strong acidifying effect since day 9 after root entry. At that time nodulation in root type biopore in the field of view occurred. Presumably, less nodulation in worm than in root type pore can be caused by a higher nitrate content in worm type sheath, because higher nitrate contents were found to inhibit the formation of the infection tube (Munns 1968). However, it is apparent that the roots showed activity when growing through the biopore sheaths. Even though pH alteration in the worm biopore sheath was less intensive than in root type sheath and bulk soil. Moreover, homorhizous spring barley showed overall no distinct effects in the ability to alter the pH value, conversely to both allorhizous crops.

Root activity is influenced by the source of nutrients (Marschner et al. 1986), the soil’s buffer capacity and initial pH value (Nye 1981; Schubert et al. 1990; Kreuzeder et al. 2018). Earthworm linings were reported to have increased pH values in comparison to the bulk soil (Tiunov et al. 2001; Horn et al. 2003) and increased nitrate contents (Parkin and Berry 1999). Nitrate or ammonia nutrition led to a physiologic acidic or alkaline response dependent on the initial soil pH value (Kreuzeder et al. 2018). In the current study the earthworm’s modification of the inner layer of the biopore sheath led to a higher initial pH value (pH 7.16), higher $C_t$- and $N_t$-contents, except $N_t$-content in spring oilseed rape, and moderate pH changes induced through the roots. These results of the $C_t$- and $N_t$-contents analysed at the end of the experiment might indicate an N-uptake at the inner layer of the sheath of spring oilseed rape. However, the comparatively moderate metabolic activity in worm type sheath can be assumed as an effect of i) higher nutrient availabilities, which resulted in less root activity, or ii) a less measurable pH change of the planar optodes caused by a higher buffering capacity (Mengel 1991; Hinsinger et al. 2003). A higher buffering capacity was found in earthworm casts due to an increasing calcite-content (Lambkin et al. 2011). Although, in a recent study with pH sensitive planar optodes a higher P availability lead to less acidification in the rhizosphere of maize and lupine (Ma et al. 2021), which would be in alignment with the assumption of less activity due to highly available nutrients in earthworm casts. Instead, in the root type sheath

![Fig. 5](image)

**Fig. 5** $C_t$- and $N_t$-content in the biopore sheath in spring oilseed rape and faba bean. Lowercase letter show significant differences if there is no letter in common between lateral distances within the respective biopore type “Worm” (left) and “Root” (right), the red dashed line shows the value of the bulk soil treatment, t-test with Bonferroni-correction, $\alpha=0.05$, ns=not significant.
no distinct enrichment with $C_t$ and $N_t$ was found, which was contrary to a former study (Petzoldt and Kautz 2021). In the current study, this could be due to removing the scarcely decayed taproot of the pre-crop before the experiment started. The N-content of the root type sheath was not different to the bulk soil, but small effects of root deposits or debris left from the pre-crop are likely. That was indicated by a slightly higher initial pH value (pH 6.99) in comparison to the bulk soil (pH 6.61) and the observed appreciable pH change in spring oilseed rape and faba bean. A higher initial pH value in the worm type than root type sheath is in line with results from the same experimental soil (Haas and Horn 2018). The earthworm and pre-crop treatment modified initial chemical conditions at the inner layer of the biopore sheath. That presumably resulted in less observable effects in the earthworm treatment with this method or less activity due to a higher availability of nutrients. The use of pH sensitive planar optodes for comparative observations of the metabolic activity of roots in environments with different initial pH values are clearly limited.

Overall, high variations of the pH value within the treatments were found (Fig. 4), which can be a result of (in)active roots, root orders and root maturity over time. Other studies using pH planar optode techniques showed, that metabolic activity changed in response to the form of phosphor (Ma et al. 2021), nitrogen and initial pH value (Kreuzeder et al. 2018), but also by root aging and differentiation (Blossfeld et al. 2013; Rudolph et al. 2013; Rudolph-Mohr et al. 2017). Complementary observations were made for temporal, permanent (Blossfeld et al. 2010) or diurnal (Blossfeld et al. 2011) differences in the metabolic activity. Our study reveals that structured subsoil contribute to the large temporal and spacial variation of root activity in the subsoil.

Conclusions

The methodological approach with pH sensitive planar optodes made it possible to measure root activity in the biopore sheath. However, earthworms’ modification of the chemical properties resulted in less pH change, which might not be necessarily a result of less root activity and would need additional investigation. Our hypothesis, that roots induce pH change in biopore sheaths was confirmed for oilseed rape and faba bean. This clearly indicates that biopores are not only preferential pathways for root growth but also hotspots of root activity. Higher pH changes around roots in worm pores were not confirmed which was probably caused by increased buffer capacity in the worm pore sheath. In conclusion, faster root growth of spring crops through deeper untilled soil layers was facilitated by biopores. Such a beneficial effect can be increased by nutrient uptake at the biopore sheath especially in periods with increased drought.

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Declarations

Conflicts of Interest The authors declare that they have no conflict of interest.

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