Transportation and localization of phenanthrene and its interaction with different species of arsenic in *Pteris vittata* L.

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**Highlights**

- Different species of arsenic absorption by *Pteris vittata* were inhibited by phenanthrene.
- Phenanthrene absorption by *P. vittata* was reduced by arsenic in the following order: DMA > As(V) > As(III).
- Phenanthrene mainly located in the cell membrane or membrane structure.

**Abstract**

The interaction between arsenic (As) and phenanthrene (PHE) in *Pteris vittata* L. was investigated in this study. The migration and occurrence of PHE in *P. vittata* were determined by two-photon laser scanning confocal microscopy. Data indicated that PHE supplementation lowers the As concentration in *P. vittata*, decreasing As levels by 16.8–39.9% in the pinnae, 30.0–49.0% in the rachis, and 45–51.5% in the roots, respectively. Different arsenic species inhibited *P. vittata* PHE absorption. The most significant effect was observed using dimethylarsenic acid (DMA), which decreased PHE accumulation by 20.73%. With the exception of elevated As(V) concentrations in As(III)-treated plants, PHE treatment significantly reduced inorganic As concentrations in *P. vittata*. However, PHE elevated root DMA concentrations by 9%. According to *in situ* visualization, PHE is primarily found in the upper and lower epidermis and stomatal cells, particularly the stomata guard cells.

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**1. Introduction**

The hyperaccumulator, *Pteris vittata* L., which can absorb and accumulate high-arsenic (As) levels, is used to treat As-polluted soil or water (Sun et al., 2011). As absorbed by *P. vittata* roots is transported aboveground via the xylem and stored in the fronds. The distribution of As varies in *P. vittata* generally following the order: pinnae > rachis > roots (Srivastava et al., 2010). The maximum As concentration accumulated in *P. vittata* pinnae can reach 22,630 mg/kg. Interestingly, 93% of the As was found in the aboveground parts of the plant, which accounts for 2.3% of the plant dry weight. Furthermore, the As concentration in the aboveground portions can be up to 25-fold higher than in the roots (Ma et al., 2001; Tu and Ma, 2002).

Arsenate [As(V)] is transported to the aboveground portions of *P. vittata* via the vascular bundles, and can undergo biochemical reduction within the plant, primarily being transformed to trivalent As [As(III)], which is stored in the fronds (Pickering et al., 2006; Hokura et al., 2006). Additionally, other arsenic species, including As(III)-O and As(III)-S compounds, are found in *P. vittata* (Webb et al., 2003). Energy dispersive X-ray (EDXA) revealed that the As concentration in *P. vittata* pinnae decreased from base to apex, with significantly lower levels in the vein than in the pinna. As was primarily distributed in the vacuoles of the epidermal cells on the upper and lower sides of the pinnae (Lombi et al., 2002). Similarly, Yang et al. (2009) found that 91% of As was present in the vacuoles, as revealed by subcellular classification. Thus, As reduction and vacuole uptake are important pathways for As detoxification by *P. vittata*, and the major mechanism of As hyperaccumulation.

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As in the environment often coexists with polycyclic aromatic hydrocarbons (PAHs), such as phenanthrene (PHE), which are carcinogenic organic pollutants that degrade slowly. Combined PAH and As pollution has been found in industrial sites, including coke plants, smelters, and coal mines. As-PAH pollution exceeded the standard rate by 40.6% in a smelting industrial site and 68.8% at a coal mining site in Chenzhou, Hunan Province, China (Zhu et al., 2012). Our previous study revealed that P. vittata is capable of absorbing both As and PAHs, which can be used to treat soil with combined As and PAH pollution. In a field investigation conducted at an industrial site in Chenzhou, P. vittata was shown to tolerate high As and PAHs levels (1276 mg/kg As and 1.48 mg/kg PHE) (Sun et al., 2014). Further experimental studies revealed that P. vittata tolerates As-PHE co-contamination. Moreover, the co-existence of As and PHE significantly affected As speciation in P. vittata and altered PHE absorption in the roots by significantly elevating PHE root concentrations in the plants (Sun et al., 2011).

The absorption, transport, and storage of As in P. vittata have been well-studied (Xie et al., 2009). However, the transportation and localization of PHE in P. vittata remains unclear. In this study, the interaction between different arsenic species and PHE in P. vittata was evaluated. The transport and storage patterns of PHE were observed using two-photon laser scanning confocal microscopy (TPLSCM) to better understand the biochemical behaviors and interaction mechanisms of As and PHE in P. vittata.

2. Materials and methods

2.1. Experiment setup

The spores of P. vittata were collected from Chenzhou, Hunan Province, China and evenly spread on a seed tray loaded with medium (flower cultivation soil:vermiculite = 1:1). The medium was kept moist by watering and covering with a plastic sheet to prevent evaporation. The spores were germinated and developed several young pinnae, the seedlings were isolated and transplanted into 1/5-strength Hoagland solution (pH 6.0). Healthy and uniform P. vittata seedlings were selected for experimental use.

Eight treatments were included in the experiment: control check, PHE, arsenite [As(III)], As(III) + PHE, arsenate [As(V)], As(V) + PHE, dimethylarsenic acid (DMA), and DMA + PHE. There were three replicates for each treatment. For each group, As was prepared at a final concentration of 10 mg/L, and PHE (dissolved in acetone) was prepared at a final concentration of 2 mg/L in nutrient solution. Arsenate was added in the form of Na2HAsO4·7H2O, arsenite was added in the form of Na2AsO2. The nutrient solution was changed every 3 d, and the plants were harvested after 12 d. The harvested plants were washed with clean water and air-dried. The pinnae, rachis, and roots were then separated to measure the fresh weight. A portion of the plant was deactivated at 105 °C for 30 min and oven-dried at 65 °C to obtain the dry weight and determine the total As. The rest of the material was freeze-dried for 24 h, fragmented in an ice bath, and maintained at −80 °C to determine the As speciation and PHE concentration.

2.2. Total arsenic and arsenic speciation analysis

The oven-dried plant samples were digested with HNO3-HClO4 (US EPA 3050), and then diluted to a constant volume. Total As was determined using an atomic fluorescence spectrometer (AFS-9130). The national standard (GBW-07603) was used as a reference for analytical quality control.

The plant samples were subjected to ultrasonic extraction with methanol and water (v/v, 1:1). The extract was diluted to a constant volume after rotary evaporation of the methanol. Arsenic speciation was analyzed by liquid chromatography coupled with atomic fluorescence spectroscopy (AFS-LC-9130) (Sun et al., 2011).

2.3. Phenanthrene analysis

The plant samples underwent 3 rounds of ultrasound extraction with a mixed solution of acetone and methylene chloride (v/v, 1:1). The upper layer of the extract was concentrated by rotary evaporation to 0.5 mL, and then transferred to a silica gel column for purification. The column was eluted with a mixture of n-hexane and methylene chloride. The eluted solution was collected and concentrated by rotary evaporation to 1 mL, followed by the addition of 7–8 mL n-hexane. The solution was concentrated by rotary evaporation to 0.5 mL, mixed with an internal standard, and adjusted to a constant volume of 1 mL. PHE was determined using gas chromatography-mass spectrometry (Agilent 6890, USA) (n = 4, RSD < 9.67). The recovery rate was 96–105%.

2.4. Detection of PHE fluorescent signal in fresh samples

The plant samples were washed with distilled water and prepared as live plant sections. The specimens were mounted with neutral gum and observed using TPLSCM (Carl Zeiss LSM 780) equipped with an objective lens (W plan-Apochromat 20 × 1.0 DIC [UV] VIS-IR M27 55 mm). The laser wavelength was set to 710 nm, the emission wavelength was 740 nm, and the detection wavelength was 371–495 nm. Microscopic photographs were analyzed using ZEN 2012 (blue edition) imaging software. The PHE fluorescent signal was blue, and the fluorescent signal of the plant was green.

2.5. Statistical analysis

Data were analyzed using SPSS 21.0 (SPSS Inc., Chicago, IL, USA). One-way analysis of variance was used to compare the effects of As on PHE concentration in plant and growth media. Multiple comparisons were made by the least significant difference (LSD) test, and P-values less than 0.05 were considered significant. Two-way analysis of variance was used to compare the interaction of arsenic and PHE. Plots were made using Origin 8.0 and ChemBioOffice.

3. Results

3.1. Concentration of As and PHE in P. vittata

PHE treatment significantly altered the total As concentration in the pinnae, rachis, and roots of P. vittata (Table 1). Compared to As treatment alone, the As(III) + PHE, As(V) + PHE, and DMA + PHE treatments decreased pinnae As concentrations by 16.8%, 39.9%, and 38.7%, respectively. Furthermore, compared to As treatment alone, PHE significantly inhibited As concentrations in the rachis and roots by 30–49.0% and 45–51.5%, respectively.

Different arsenic species exhibited a significant effect on PHE absorption by P. vittata. Compared to PHE treatment, DMA reduced pinnae PHE concentrations, whereas As(III) or As(V) treatment had minimal effects. Treatment with different arsenic species decreased PHE concentrations in the roots and rachis of P. vittata.

Compared with As treatment alone, PHE decreased 50% As accumulation of pinnae (Fig. 1). The corresponding rachis and root As decreased by 68.9–117.2 and 51.2–149.4 μg/plant, respectively. PHE-induced inhibition of As accumulation varied between different arsenic species. As(V) showed the greatest decrease in As.
accumulation, followed by DMA, whereas As(III) caused a relatively small decrease.

Among the different arsenic species, DMA significantly inhibited PHE accumulation in *P. vittata*, decreasing accumulation by 295 ng/plant. As(III) and As(V) also inhibited PHE accumulation by 147 and 56.53%, respectively, whereas the proportion of As(III) among total As decreased. This is probably because As(V) concentrations were greatly decreased by the former two treatments, but slightly increased by the latter. Compared with As treatment alone, PHE and DMA treatments decreased As(V) concentrations by 56.51% and 78.83%, respectively. DMA was only detected in the pinnae of plants after DMA treatment. Following PHE treatment, the proportion of DMA increased by 2.5%, while its concentration decreased by 30.94 mg/kg dry weight (DW).

In the rachis, PHE had little effect on As speciation following As(III) treatment. However, PHE inhibited the absorption and transformation of different arsenic species following As(V) and DMA treatments. PHE promoted root absorption of DMA, resulting in an elevation of DMA proportion (9%) and concentration (4.3 mg/kg DW). However, PHE inhibited root absorption of inorganic As (Fig. 2-root).

### 3.3. Microscopic distribution of PHE in *P. vittata*

PHE signals were observed in different parts of *P. vittata* by TPLSCM (Fig. 3). Following PHE treatment alone, PHE signal was observed in the roots, primarily near the cell wall and in the cytoplasm (Fig. 3A-root). PHE was evenly distributed in the cytoplasm, with individual storage sites in the cells. The intensity of PHE signals in the roots descended as follows: near the cell wall > individual storage sites in the cytoplasm > cytoplasm. In the rachis, PHE was primarily distributed in the U-vascular bundle cells, the walls of the outer cells, and the cytoplasm of certain inner cells.
Fig. 2. Concentrations of arsenic speciation (mg/kg) and proportion (%) in pinnae (A), rachis (B) and roots (C) of *P. vitata* upon various treatments with different arsenic species with or without PHE. Data are means ± SE (n = 3).
Fig. 3. The microscopic distribution of PHE signal in *P. vittata* under four different medium containing PHE treatment (A, B, C, D represent PHE, As(III) + PHE, As(V) + PHE, and DMA + PHE, respectively. Blue is the PHE fluorescent signal, green is the plant fluorescent signal). The red arrows point to the spot which PHE existed. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
(Fig. 3A-rachis). The intensity of PHE signal in the rachis varied, as follows: wall of U-vascular bundle cells > wall of outer cells > inner cells. In the pinnae, PHE was observed in the pinnae veins, epidermal cells, and a small number of mesophyll cells (Fig. 3A-pinnae). The strongest PHE signal in the pinnae was observed in the epidermal cell walls and guard cells. In the epidermal cell cytoplasm, a strong signal was observed on the surface of some organelles (i.e., organelle membranes). The intensity of the PHE signal was as follows: epidermal cells > pinnae veins > mesophyll cells (palisade and spongy tissues).

Different arsenic species did not alter the absorption or localization of PHE, however, they altered the intensity of the PHE signal. After treatment with different arsenic species, PHE signal in the cytoplasm of the root cells was significantly weakened. Following As(V) or DMA treatment, PHE signal was weakened in the outer cell membrane of the rachis. Additionally, the signal intensity of PHE was relatively lower in U-vascular bundles after treatment with different arsenic species.

4. Discussion

4.1. Study on the absorption and translocation of PHE in P. vittata

In this study, the absorption and transport pathway of PHE in P. vittata was speculated in Fig. 4 based on the in situ visualization by TPL SCM. PHE absorbed by root epidermal cells, and then transported to the aboveground portions of the plant through apoplast or symplast. The majority of PHE is transported to the fronds through U-vascular bundles in the rachis, while a small portion is transported upward through rachis epidermis cells. PHE is transported to the mesophyll, epidermis, and stomatal guard cells in the fronds through the pinnae veins (Fig. 4).

With the TPL SCM, the absorption and transport of PHE in the roots were observed in Fig. 3. After being cultured in contaminated solutions, PAHs can also be absorbed through the roots and transported to the aboveground parts of the stems and leaves in various plant species, such as Oryza sativa L. (Jiao et al., 2007), Lolium multiflorum L. (Gao et al., 2010) and Helianthus annus L. (Sneath et al., 2013). Transpiration drives the absorption and transport of PAHs from the underground part to the aboveground part (Tao et al., 2006; Mattina et al., 2003; Gao and Zhu, 2004; Wu et al., 2009). For instance, Wang et al. (2012) proposed that PAHs are absorbed in the root epidermis before entering the pith and lateral roots. They then enter the xylem and phloem of the taproot through transpiration. Moreover, the PAHs in the ambient air could deposit to the surface cuticular wax of the leaves (Howsam et al., 2000, 2001).

4.2. Study on the localization of PHE in P. vittata

The localization of PAHs in plants is associated with its biochemical behaviors, as well as the structure and function of plant cells. PAHs have a relatively low solubility in water and low Henry’s law constants, but are highly lipophilic and have high octanol-water partition coefficients (Kow), while the phospholipid bilayer of the cell membrane or cytoplasm in different parts of the plant. In the pinnae, PHE is found in the cell membrane or cytoplasm in different parts of the plant. In the pinnae, PHE is primarily stored in the upper and lower epidermal cells, and the stomatal guard cells of the fronds. The retention of PAHs in the pinnae surface cuticular wax and the cuticle may facilitate the photodegradation and volatilization of the compounds (Wild et al., 2005). Presumably, PHE is exchanged to the external environment or photodegraded in stomatal guard cells, which contain large amounts of PHE, to reduce PHE-induced damage to the plants.

Interestingly, the concentration or signal intensity of PHE in P. vittata was influenced by different arsenic species, while the adsorb to the cell wall of the roots, then enter the organelles within the cell via the cell membrane. Ren et al. (2010) compared the transmembrane transport of four typical lipophilic organic compounds, and proposed that PAHs in the extracellular medium enter the cytoplasm in a steady flow through the partition-inverse release-binding mechanism. Keyte et al. (2009) indicated that PHE was primarily concentrated in the cytoplasm and vacuoles in spinach (Spinacia oleracea), while in the cell membrane and adjacent cells in mosses (Hypnum cupressiforme). In P. vittata, PHE is found in the cell membrane or cytoplasm in different parts of the plant. In the pinnae, PHE is primarily stored in the upper and lower epidermal cells, and the stomatal guard cells of the fronds. The retention of PAHs in the pinnae surface cuticular wax and the cuticle may facilitate the photodegradation and volatilization of the compounds (Wild et al., 2005). Presumably, PHE is exchanged to the external environment or photodegraded in stomatal guard cells, which contain large amounts of PHE, to reduce PHE-induced damage to the plants.
migration pathways and storage sites of PHE in the plant had minor effects (Fig. 3). We speculated that PHE and As was regulated by different transporter in *P. vittata* and PHE was stored in cell membrane owing to its biochemical behaviors (e.g. high Kow). Whether this is the reason for the appearance which showed in Fig. 3 should be explored in future research.

4.3. Effect of PHE on As migration and transformation in *P. vittata*

PHE inhibited As absorption in the roots, rachis and pinnae of *P. vittata*. Additionally, the concentrations of inorganic and organic arsenic species decreased by PHE treatment, with the exception of As(V) in aboveground portion of the plant and DMA in the root after As(III) and DMA treatment (Table 1). It was inferred that PHE inhibits As absorption for two reasons. Firstly, the activity of the arsenic transporter or arsenate reductase affected by PHE, thereby inhibiting the absorption and transformation of As. The As(V) reductase gene in *P. vittata* has been coded, and other As(V) reducing proteins or As(V) reductases, such as TPI and Grx5, were also involved in reduction (Rathinasabapathi et al., 2006; Sundaram et al., 2008). PHE may affect the normal transport and transformation of As by binding to As transport proteins on the cell membrane or As reductase in the cytoplasm, leading to decrease As(III) and As(V) concentrations. Secondly, the transport pathway of different arsenic species, particularly inorganic arsenic, was influenced by PHE. The research of Pickering et al. (2006) demonstrated that, As(III) and As(V) are transported through the xylem and phloem in the rachis of *P. vittata*, respectively. As(V) primarily exists as a bundle in the central vein of the pinnae or the center of the rachis, and primarily plays a role in storage and transportation. In contrast, there was weak signal of As(III) in the central vein of the pinnae; in fact, As(III) presents an interlaced distribution. Moreover, Kachenko et al. (2010) showed that As(III) is present in the vascular bundles of the roots. Sixty percent of As(V) is absorbed and reduced to As(III) in the roots, meanwhile, all different speciation of As are transported to the aboveground part of the plant through this vascular bundle tissue. In the present study, it was also revealed that PHE is transported into the cells of the rachis and pinnae through the vascular bundle tissue. Different speciation of As share the channel with PHE for upward transport and migration. PHE likely affects the inorganic As transport capacity of vascular bundle tissue and reduces the As accumulation capacity.

5. Conclusions

(1) As absorption in the roots, rachis, and pinnae of *P. vittata* were inhibited by PHE. PHE treatment lowered inorganic and organic As concentrations following different arsenic species treatment, with the exception of the elevated concentrations of As(V) in the aboveground parts after As(III) treatment and DMA in the roots after DMA treatment.

(2) Different species of As exhibited an inhibitory effect on PHE absorption in *P. vittata*. DMA had the greatest inhibitory effect, followed by As(V), and finally, As(III).

(3) PHE mainly occurs in the cell membrane or membrane structure (organelle membrane) and cytoplasm. The specific sites include the membrane and cytoplasm of root cells; epidermis cell membranes, U-vascular bundle cells and the cytoplasm of inner cells in the rachis; veins, epidermal cells and stomatal guard cells in the pinnae.

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References


