Metabolite profiling of rhizosphere soil of different allelopathic potential rice accessions

Yingzhe Li 1,2, Lining Xu 1,2, Puleng Letuma 1,2,3 and Wenxiong Lin 1,2,4*

Abstract

Background: Identification of the allelopathy-interrelated metabolites from the allelopathic rice rhizosphere is crucial to understand the allelopathic mechanism of rice, which in turn can promote its applications to farming. In this study, the metabolites from the rhizosphere soil of five different rice lines, including allelopathic rice accession PI312777 (PI) and non-allelopathic rice accession Lemont (Le) as well as their genetic derivatives (e.g., phenylalanine ammonia-lyase (PAL) gene overexpression transgenic lines of PI and Le, namely, PO and LO respectively, and PAL RNA interference line of PI, namely, PR) were identified and comparatively analyzed to explore the positive compounds that are involved in the process of rice allelopathy.

Results: The results showed that 21 non-polar compounds and 21 polar compounds differed in content in the rhizosphere soil of PI and Le, which include several volatile fatty acids and long-chain fatty acids. The relative contents of fatty acids also differed between PAL overexpressing or RNA interference (RNAi)-silenced line and their wild-type respectively. Acetic acid content also differed among groups, i.e., it is higher in the high allelopathic potential rice. Further analysis showed that different metabolites from the ADS8 resin-extracted phase were more abundant than that those from the ADS21 resin-extracted phase, suggesting that the allelochemicals in root exudates of allelopathic rice are mainly non-polar substances. KEGG annotation of these differential metabolites revealed that these compounds were related to nutrient metabolism, secondary metabolite synthesis, signaling substance synthesis, and toxin degradation.

Conclusions: Rice allelochemicals deposited in the ADS8 resin-extracted phase were more abundant than those in the ADS21 resin-extracted phase. Allelochemicals in root exudates of allelopathic rice are mainly non-polar substances, and long-chain fatty acids are considered as allelopathy interrelated metabolites.

Keywords: Fatty acids, Metabolomics, Rhizosphere, Allelopathy, Resin extraction
Background

Allelopathy is a chemical ecological phenomenon in which donor plants release their chemical substances, called allelochemicals, into the environment through the process of secretion, volatilization, leaching, and residual degradation and affect other recipient organisms [1], which could be used to control harmful organisms, such as weeds and pests in the field. Therefore, it has become the focus of research studies [2–7]. There are two main viewpoints on the categories of allelochemicals in rice. Phenolic acids are considered as a group of allelochemicals, and the others are terpenoids and flavonoids. Rice and Chou et al. believed that phenolic acids, such as p-coumaric acid, ferulic acid, p-hydroxybenzoic acid, and oxalic acid, which are produced after the decomposition of rice residues, could be fixed by soil aggregate structure substances or humic acid and stored in the rhizosphere soil to inhibit the growth of rice seedlings and weeds [1, 8, 9]. Rimando and Seal et al. isolated allelochemicals from root extracts of rice, and they identified several major phenolic allelochemicals with inhibitory effects on the target plants [10, 11]. All of this evidence strongly supports the idea that phenolic acids are allelochemicals. However, Olofsdotter et al. were skeptical of this issue and suggested that phenolic acid compounds might not be the allelochemicals that inhibit the target weeds using 4-aminoantipyrine spectrophotometry [12].

Another argument was that phenolic acids impart inhibitory effects on the target grasses, but the dosage used for the bioassay was much higher than that detected in rhizosphere soil of allelopathic rice accession. Therefore, terpenoids, such as momilactone B, were considered as promising allelochemicals in rice because these exhibited the higher inhibitory effect on the target weeds at very lower dosages of 3–30 μmol/L [13–16]. However, all of the above results were obtained under laboratory conditions. Furthermore, the effective concentrations of phenolic acids and terpenoids used in laboratory bioassays are always higher than those of allelopathic rice released and detected in the field, which therefore has been often questioned by some scholars.

Previous studies have shown that plant allelopathy is mainly a complex rhizosphere biological process that is mediated by allelochemicals (secondary metabolites) secreted by their roots. Allelopathic plant accessions have different potentials in inhibiting target weeds, which is mainly due to differences in the functions of allelopathic genes in the synthesis of secondary metabolites and the results of the interactions between plant, soil, and rhizosphere microorganisms mediated by the allelochemicals, which in turn require a more in-depth study to elucidate the allelopathic mechanism underlying this particular rhizosphere biological process [17–20].

Previous studies have documented that several genes, including copalyl diphosphate synthase 4 (OsCPS4), kaurene synthase-like 4 (OsKSL4), and phenylalanine ammonia lyase (OsPAL), play roles in the process of rice allelopathy. Knocking out or silencing OsCPS4, OsKSL4, and OsPAL results in weaker allelopathic activity from the donor rice to the barnyardgrass. This weakened activity is attributed to reduced momilactones or phenolic acid content in root exudates [21, 22]. Among these genes, OsPAL is the first key gene in the phenylpropanoid metabolism and functions in the regulation of phenolic acid synthesis. Numbers of studies have documented that gene expression level of OsPAL from allelopathic rice accession PI312777 was higher that from non-allelopathic rice Lemont [23–26], of which these two rice accessions were widely taken as donor plant to investigate the underlying mechanism of allelopathy, PI312777 was first reported by Dilday in the field experiments to identification of allelopathic rice germplasm [27], while the Lemont rice is the contrary line with non-allelopathic activity [23–26]. Silencing of OsPAL gene expression in PI312777 results in a 50% reduction of weed-suppress capacity, whereas its overexpression increases the inhibition ratios of the weed by 13% [19]. The diversities of microbial community from the rhizosphere of these genetic modified rice have also changed compared to that of the rhizosphere of WT of PI312777, which are possibly correlated with changes in root exudates secretion. These metabolites are thus regarded to play crucial roles in the process of weed suppression.

Scientists have gradually realized that allelopathy is an ecogenetic trait that is involved in an extremely complex chemico-biological process in the rhizosphere ecosystem. However, more recent studies have shown that plant

![Fig. 1](https://example.com/figure1.png)
allelopathy includes direct allelopathic effects caused by allelochemicals, and indirect allelopathic effects mediated by allelochemicals through microbial utilization, transformation, and resynthesis. Therefore, it has been suggested that allelopathy in rice might result from the interaction of allelochemicals with specific microorganisms in rhizosphere soil [28–31]. Therefore, extensive studies have focused on assessing the interaction between allelopathic rice and rhizosphere microorganisms [19, 22, 32], including the separation and identification of allelochemicals. Thus, what other substances, besides phenolic acids and terpenoids, are allelochemicals? This requires the application of metabonomics, a bioanalytical approach that has recently emerged.

Table 1 Comprehensive comparison of ADS8 resin extracted fatty acids from the rhizosphere soil of rice

| VIP Name | PI: Le | VIP Name | PI: PR | VIP Name | PO: PI | LO: Le |
|----------|--------|----------|--------|----------|--------|--------|
| Acetic acid | U | 1.90713 | U | 1.70646 | Le | Benzoic acid |
| Pentanoic acid | U | 1.59614 | U | 1.57475 | Le | Acetic acid |
| Heptanoic acid | U | 1.25548 | U | 1.25174 | PR | Propenoic acid |
| Cyclopropanecarboxylic acid | D | 1.25499 | U | 1.09064 | PO | Azelaic acid |
| Methylphosphonic acid | D | 1.36746 | D | 1.49533 | PO | Cyclohexanecarboxylic acid |
| Octanoic acid | D | 1.37916 | D | 1.48714 | PR | 3-Amino-2,3-dihydrobenzoic acid |
| Fumaric acid | D | 1.0572 | D | 1.44648 | PR | Succinic acid |
| 8,11,14-Eicosatrienoic acid | U | 1.20765 | U | 1.14348 | Le | 2,2-Bis(4-hydroxyphenyl)-propanoic acid |
| Note: U upregulated expression. D downregulated expression |

Metabolites, which are the end products of cell metabolism regulation, are the material basis of biological phenotypes, and can directly and effectively reflect biological processes and contribute to the analysis of their mechanisms [33]. Changes in metabolites, including species and quantity, are the ultimate responses of biological systems to internal or external stimulus, such as gene mutations or environmental stress [34]. Qualitative and quantitative analyses of low-molecular weight metabolites have been conducted to investigate metabolic pathways or metabolic networks, to compare and analyze the metabolic differences in macroscopic phenotypic phenomena among different biological individuals, and to study the metabolic response mechanism of

Table 2 Comprehensive comparison of ADS21 resin extracted fatty acids from the rhizosphere soil of rice

| VIP Name | PI: Le | VIP Name | PI: PR | VIP Name | PO: PI | LO: Le |
|----------|--------|----------|--------|----------|--------|--------|
| Heptadecanoic acid | U | 1.95187 | U | 1.25069 | Le | 1,2-Benzenedicarboxylic acid |
| Hexanoic acid | U | 1.14989 | U | 1.28803 | PR | Cyclohexanecarboxylic acid |
| Pentanoic acid | D | 1.15034 | D | 1.17727 | PO | Hexanoic acid |
| 8,11,14-Eicosatrienoic acid | D | 1.50193 | D | 1.33051 | PO | Cyclopropanecarboxylic acid |
| 9-octadecanoic acid | D | 1.30172 | D | 1.17800 |
| Hexadecanoic acid | D | 1.08017 | D | 1.08017 | PO | Decanoic acid |
| Note: U upregulated expression. D downregulated expression |
substances after different induction and stress [33]. Metabolomics is usually used in combination with transcriptome and proteomics to study how changes in the physiological pathway of DNA → mRNA → protein → metabolite elicit responses to various environmental stimuli [33]. Due to the “high-dimensional and massive” nature of metabolomics data, the integration of statistical analysis of differential metabolites has facilitated accurate mining and metabolic pathway annotation, and other studies have been conducted to elucidate the mechanism underlying sample differences. Our previous studies have shown that when plant extracts or root secretions from allelopathic rice are treated with polar resin at the five-leaf stage, the inhibition rate of target weeds significantly increased, and the reverse was observed when plant extracts or root secretions were treated with a non-polar resin [35]. We also found that allelopathic rice accessions had much more microbial diversity than its counterparts because allelopathic rice synthesize and release higher amounts of allelochemicals, such as phenolic acids into rhizosphere soil environments, implying that the composition of metabolites in the rhizosphere soil ecosystem is a key factor influencing microbial community structure. In this study, allelopathic rice and non-allelopathic rice accessions as well as their genetic derivatives, namely, transgenic lines with major allelopathic gene PAL2–1-inhibited and PAL2–1-overexpressed, were used as research materials. Metabolomics was used to analyze differences in the metabolites extracted from the rhizosphere soil of different allelopathic potential rice accessions using different resins at the five-leaf rice seedling stage to improve our understanding of the chemoeocological characteristics of rhizosphere soil in allelopathic rice.

Fig. 2 Analysis of differential metabolites between PI and Le samples absorbed by ADS-8 resin from rhizosphere soil. a The relative content of the metabolites between PI and Le displayed in the heat map, and the differential substances are arranged according to the VIP values from small to large. Each row represents a compound; of which the value of each compound represents the relative content directly normalized on the scale of the graph. Bluish color represents low content while reddish color represents high content. b The possible metabolic pathway distribution of single metabolic substance between PI and Le. c The left side represents the number of single differential substance in the same pathway where PI is lower than Le, and the right side is the opposite.
Results

**PAL gene and protein expression on the rice**

To validate the positive transgenic rice line of PI and Le, we assessed the PAL gene and protein expression levels in relation to the WT of PI and Le, which were higher, whereas the RNA interference transgenic line of PI showed a decrease in PAL gene and expression in rice relative to the WT of PI. Further comparison of the PAL protein expression on these rice lines also showed that the protein expression levels of PAL were higher in the PAL overexpression rice line than the WT, and the protein expression levels of PAL from the PAL RNAi interference of PI were lower than the WT of PI (Fig. 1). In addition, the changes in PAL expression in rice did not substantially influence the phenotypes and growth period traits.

**Principal component analysis (PCA) of different polar metabolites from rice rhizosphere soil**

PCA was performed on the samples to assess the variability of each sample within the group. For the non-polar metabolites from rhizosphere soil of PI, PR, PO, Le, and LO, and those metabolites from the same soil without rice plants (blank soil), the results showed that six groups of samples were distinctly clustered, indicating that the metabolites from different rice rhizospheric soil were clearly different, whereas the three repeats from the same group only showed slight differences (Additional file 2). Specifically, metabolites from the samples of PI and Le were deposited in the first principal component (t [1]), whereas PI was distinguished from Le and similar to the blank control CK (Additional file 2). For the three transformed rice lines, PO was distinguished from PR and LO among transgenic lines using principal component II (t [2]).

PCA was also performed on six samples extracted from the ADS-21 values. The results showed that although the samples of PO, Le, and LO were roughly close to each other, these were well separated among samples of the six groups, indicating that the rhizosphere soil samples from different rice accessions could be clearly distinguished, but the samples within the same group also exhibited slight differences (Additional file 3). The samples from the PR and PO transgenic lines and Le non-allelopathic rice were separated from the sample of PI allelopathic rice, Le was isolated from its

| Name | KEGG  | VIP | PI | Le | Participating in metabolic pathways |
|------|-------|-----|----|----|-------------------------------------|
| Acetic acid | C00033 | 1.90713 | U | D | ko01120 Microbial metabolism in diverse environments (7) |
| 5-Cholesterol-3,26-diol | C17336 | 1.71897 | U | D | cpd:C00037 Glycine |
| Cyclohexanol | C00854 | 1.63707 | U | D | cpd:C0122 Fumarate |
| Pentanoic acid | C00803 | 1.59614 | U | D | cpd:C00469 Ethanol |
| 1,25-Dihydroxyvitamin D3, TMS derivative | C01673 | 1.58096 | U | D | cpd:C00854 Cyclohexanol |
| Ethanol | C00469 | 1.50588 | D | U | cpd:C16267 Cyclopropanecarboxylate |
| Androsta-1,4-dien-3-one, 17-hydroxy-17-methyl-, (17. alpha.)- | D00389 | 1.49721 | D | U |
| Benzenamine | C00292 | 1.4247 | D | U | |
| Semicarbazide | C02077 | 1.39372 | D | U | ko04961 Endocrine and other factor-regulated calcium reabsorption (1) |
| Pseudosmilagenin | C19650 | 1.37739 | D | U | cpd:C01673 Calcitriol |
| Cyclopropanecarboxylic acid | C16267 | 1.30204 | D | U | ko01040 Biosynthesis of unsaturated fatty acids (1) |
| Heptanoic acid | C17714 | 1.27283 | U | D | cpd:C03242 (8Z,11Z,14Z)-Icosatrienoic acid |
| Methylphosphonic acid | C20396 | 1.24085 | U | D | ko00061 Fatty acid biosynthesis (1) |
| 8,11,14-Eicosatrienoic acid | C03292 | 1.20765 | U | D | cpd:C06425 Octanoic acid |
| Deoxyxypogualin | D08032 | 1.18815 | U | D | |
| 2-Butanone | C02845 | 1.17795 | U | D | |
| Ethyl 3-hydroxybutyrate, TMS derivative | C03499 | 1.17133 | D | U | |
| Octanoic acid | C06423 | 1.17 | U | D | |
| Glycine | C00037 | 1.09679 | U | D | |
| Ether | C13240 | 1.09528 | D | U | |
| Fumaric acid | C00122 | 1.0572 | D | U | |

Note: _U_ upregulated expression, _D_ downregulated expression
counterpart, LO, and PI was distinguished well from Le by principal component I (t[1]). Furthermore, the samples of PO and LO were closer to those of Le, especially PO and Le samples (Additional file 3).

**Metabolites in the ADS8 and ADS21 resin-extracted phase**

 ADS8 and ADS21 resins mainly adsorb non-polar and polar compounds, respectively, compared to the metabolites from the rhizosphere soil, which were respectively deposited onto the ADS8 resin and ADS21 resin, showing that the compounds from ADS8 resin, which were classified as non-polar compounds, were more abundant than the compounds from the ADS21 resin, which were polar compounds. Among the five rice lines, there were relatively higher quantity and abundance of non-polar substances, such as beta-n-butylerther, 9-octadecenoic acid, and pentadecane, than polar substances, such as 2-butanolone, d-mannitol, and nitrile. Acetic acid was detected in the rhizosphere soil of PI, Le, PR, PO, and LO, using the ADS8 resin. Based on the ADS21 resin-extracted phase, d-glucose, propanal, glutaric acid, and polar compounds were detected in the rhizosphere soil of PI, Le, PR, PO, and LO. The results indicate that rhizosphere soil metabolites with non-polar properties were more abundant than those with polar properties.

![Fig. 3](image-url)

**Fig. 3** Analysis of differential metabolites between PI and Le samples absorbed by ADS-21 resin from rhizosphere soil. a The relative content of the metabolites between PI and Le is shown in the heat map, and the differential substances are arranged according to the VIP values from small to large. Each row represents a compound. The value of each compound represents the relative content directly normalized on the scale of the graph. Blue represents low content while red represents high content. b The possible metabolic pathway distribution of single metabolic substance between PI and Le. c The left side represents the number of single differential substance in the same pathway where PI is lower than Le, and the right side is the opposite.
Changes in ADS8 resin-extracted fatty acids from the rhizosphere soil of rice

Previous studies have mainly focused on changes and differences among phenolic acids and terpenoids, two kinds of allelochemicals from different allelopathic rice accessions. Furthermore, the fatty acids, e.g., long-chain fatty acids and volatile fatty acids, were also regarded to play potential roles in allelopathy [1, 36, 37]. In the ADS8 resin-extracted phase from the rhizosphere soil of allelopathic rice PI and Le, eight kinds of fatty acids were detected at different concentrations in the two samples, with acetic acid, pentanoic acid, and heptanoic acid showing higher relative contents from PI than Le, whereas cyclopropanecarboxylic acid, methylphosphonic acid, octanoic acid, and fumaric acid exhibited higher concentrations in the rhizosphere soil of Le. Silencing PAL gene expression in PI rendered levels of hexanoic acid, acetic acid, heptanoic acid, and pentanoic acid, lower than in WT PI. It also rendered the concentration of hexanedioic acid higher than in WT PI. When the PAL gene was overexpressed in PI, the levels of five fatty acids, including benzoic acid, propenoic acid, cyclopropanecarboxylic acid, and azelaic acid, increased and levels of pentanoic acid and thioacetic acid decreased. Changes in fatty acid levels were also detected in the PAL overexpressing line of Le. This showed that butanoic acid and acetic acid levels were higher in the LO than in the WT Le. In addition, there were eight fatty acids, namely, 9-octadecenoic acid, heptanoic acid, cyclohexanecarboxylic acid, 3-amino-2,3-dihydrobenzoic acid, succinic acid, 2,2-bis(4-hydroxyphenyl)-propanoic acid, octanoic acid, and decanoic acid which exhibited lower concentrations in the LO compared to Le (Table 1).

A comprehensive comparison among the soil samples from PI and Le, PI and PR, PO and PI, as well as LO and Le indicated that acetic acid is one of the potential compounds from the non-polar phase that is involved in the allelopathic process.

Changes in ADS21 resin-extracted fatty acids from the rhizosphere soil of rice

Identification of the ADS21 resin-extracted phase showed that several fatty acids were also deposited in this resin. Comparison of fatty acids deposited in the resins, which were used for the extraction of PI and Le rhizosphere soil showed that there were seven fatty acids occurring at different concentrations in the two rice samples. Among these seven compounds, heptanediolic acid, hexanoic acid, and pentanedioic acid showed relatively higher levels in PI than Le, whereas 8,11,14-eicosatrienoic acid, 9-octadecanoic acid, hexadecanoic acid and dodecanoic acid exhibited relatively lower levels in the LO compared to Le (Table 1).

| Name | KEGG | VIP | PI | Le | Participating in metabolic pathways |
|------|------|-----|----|----|-----------------------------------|
| Phenol | C00146 | 2.74924 | D | U | ko01100 Metabolic pathways (4) cpd:C00249 Hexadecanoic acid cpd:C00829 Naphthalene cpd:C02656 Pimelate cpd:C03242 (8Z,11Z,14Z)-Icosatrienoic acid |
| 8,11,14-Eicosatrienoic acid | C03242 | 2.0113 | D | U | |
| Heptanediolic acid | C02656 | 1.95187 | U | D | |
| Hexanoic acid | C01585 | 1.71039 | U | D | |
| 9-octadecanoic acid | C01530 | 1.7027 | D | U | |
| Naphthalene | C00829 | 1.6561 | U | D | ko00627 Aminobenzoate degradation (2) cpd:C00146 Phenol cpd:C00292 Aniline |
| Pentanedioic acid | C00489 | 1.63059 | U | D | |
| Hexahydropyridine | C01746 | 1.61662 | U | D | |
| Hexadecanoic acid | C00249 | 1.58059 | D | U | ko00982 Drug metabolism - cytochrome P450 (1) cpd:C01471 Acrolein |
| Chromane-3-carbonitrile | C11697 | 1.47238 | U | D | |
| Propenal | C01471 | 1.39017 | D | U | ko00310 Lysine degradation (1) cpd:C00489 Glutarate |
| Trimethylamine | C00565 | 1.37453 | D | U | |
| Acetamide | C06244 | 1.24142 | D | U | ko02010 ABC transporters (1) cpd:C00392 Mannitol |
| d-Mannitol | C00392 | 1.21903 | D | U | |
| Pyrrolidine-2,5-dione | C07273 | 1.18807 | U | D | |
| Glycylglycine | C02037 | 1.13566 | D | U | |
| alpha-D-Galactopyranose | C00738 | 1.1085 | U | D | |
| Heptanal | C14390 | 1.08526 | U | D | |
| Dodecanoic acid | C02679 | 1.06824 | D | U | |
| Benzenamine | C00292 | 1.01478 | U | D | |
| 1,2,4-Triazol-3-amine | C11261 | 1.0102 | U | D | |

Note: U upregulated expression. D downregulated expression
concentrations in PI than Le. In the rhizosphere soil of the PR rice line, four fatty acids showed differential concentrations in the PI and PR, of which pentanedioic acid and cyclopropanecarboxylic acid were higher in the PI than PR, but the concentrations of tetradecanoic acid and 10-undecenoic acid were lower in PI than PR. When \textit{PAL} was overexpressed in the PI, the 1,2-benzenedicarboxylic acid, cyclohexanecarboxylic acid, and hexanoic acid exhibited higher levels of accumulation in PI than PR, and the concentration of cyclopropanecarboxylic acid in these two rice samples showed the opposite trend. In addition, there were eight fatty acids that were differentially generated in Le than in its transgenic line LO. The concentrations of isobutyric acid, cis-2-dodecenoic acid, hexadecanoic acid, decanedioic acid, and cyclopropanecarboxylic acid were higher in LO than in Le, whereas that of 9-octadecanoic acid, 2-butenolic acid, and dodecanoic acid were lower in the soil sample of LO than Le (Table 2).

Differences in metabolites from the rhizosphere soils of PI and Le
A full comparison of the ADS8-extracted phase of rhizospheric soil of PI and Le showed that there were mainly 21 compounds that were differentially expressed across

![Fig. 4](attachment:fig4.png)

Fig. 4 Analysis of differential metabolites between PI and PR samples absorbed by ADS-8 resin from rhizosphere soil. a The relative content of the metabolites between PI and PR is presented in the heat map, and the differential substances are arranged according to the VIP values from small to large. Each row represents a compound. The value of each compound represents the relative content directly normalized on the scale of the graph. Blue represents low content while red represents high content. b The possible metabolic pathway distribution of single metabolic substance between PI and PR. c The left side represents the number of single differential substance in the same pathway where PI is lower than PR, and the right side is the opposite.
these two samples (Fig. 2, Table 3), and 8 compounds—acetic acid, cyclohexanol, pentanoic acid, heptanoic acid, 8,11,14-eicosatrienoic acid, deoxyspergualin, 2-butanone, and glycine, more abundant in the extraction from rhizosphere soil of PI than Le. Acetic acid showed the most pronounced difference (VIP = 1.90713). In contrast, there were 13 compounds that showed lower content in the PI rhizosphere soil than the Le rhizosphere soil, which include 5-cholesten-3,26-diol, 1,25-dihydroxyvitamin D3, ethanol, androsta-1,4-dien-3-one, 17-hydroxy-17-methyl-, benzenamine, semicarbazide, pseudosmilagenin, cyclopropanecarboxylic acid, methylphosphonic acid, ethyl 3-hydroxybutyrate, ether, and fumaric acid (Fig. 2a and b). These compounds participated in microbial metabolism in diverse environments (acetate, glycine, fumarate, aniline, ethanol, cyclohexanol, and cyclopropanecarboxylate), endocrine (calcitriol), and other factor-regulated calcium reabsorption, and biosynthesis of fatty acids (octanoic acid) and unsaturated fatty acids [(8Z,11Z,14Z)-icosatrienoic acid involved].

Identification and comparative analysis of the ADS21-extracted phase also showed that there were 21 compounds that were differentially expressed in the rhizosphere soil of PI and Le. Among the 21 compounds, 11 compounds were relatively higher in the PI than Le, whereas 10 other compounds were higher in the Le. Phenol was the most significantly differentially expressed metabolite (VIP = 2.74924),

![Fig. 5 Analysis of differential metabolites between PI and PR samples absorbed by ADS-21 resin from rhizosphere soil.](image)

- **A** The relative content of the metabolites between PI and PR is shown in the heat map, and the differential substances are arranged according to the VIP values from small to large. Each row represents a compound. The value of each compound represents the relative content directly normalized on the scale of the graph. Blue represents low content while red represents high content.
- **B** The possible metabolic pathway distribution of single metabolic substance between PI and PR. The left side represents the number of single differentia substance in the same pathway where PI is lower than PR, and the right side is the opposite.
which was higher in Le than PI. The differential content compounds between these two rice lines might participate in the metabolic pathways (four compounds, namely, hexadecanoic acid, naphthalene, pimelate and \((8Z,11Z,14Z)\)-icosatrienoic acid, involved on the pathway), aminobenzoate degradation (phenol and aniline involved), drug metabolism-cytochrome P450 (acrolein involved), lysine degradation (glutarate involved), and ABC transporters (mannitol involved) (Fig. 3a and b, Table 4).

The results of the analysis of the metabolic pathways in which each of the 21 differential metabolites were intricate, as shown in Fig. 3c and d, illustrated that all of the differential metabolites involved in xenobiotics biodegradation and metabolism and metabolism of cofactors and vitamins were higher in content in PI than in Le. In contrast, the differentially expressed metabolites involved in lipid metabolism, energy metabolism and carbohydrate metabolism were downregulated in PI.

**Differences in the expression of metabolites from the rhizosphere soils of PI, PR, and PO**

The samples from the rhizosphere soil of PI and PR were analyzed according to the above methods, and the results indicated that there are 26 key differential

---

**Fig. 6** Analysis of differential metabolites between PI and PO samples absorbed by ADS-8 resin from rhizosphere soil. a The relative content of the metabolites between PI and PO is shown in the heat map, and the differential substances are arranged according to the VIP values from small to large. Each row represents a compound. The value of each compound represents the relative content directly normalized on the scale of the graph. Bluish color represents low content while reddish color represents high content. b The possible metabolic pathway distribution of single metabolic substance between PI and PO. c The left side represents the number of single differentia substance in the same pathway where PI is lower than PO, and the right side is the opposite.
metabolites between PI and PR samples (Fig. 4a and b, Table 5), of which hexanoic acid was significantly different. All of these might be involved in the metabolic pathways: Microbial metabolism in diverse environments (six substances are involved in the process, i.e., acetate, L-aspartate, cyclohexanone, adipate, fluoren-9-one, and benzamide), tropane, piperidine, and pyridine alkaloid biosynthesis (piperidine involved in the pathway), fructose and mannose metabolism (mannitol takes part in the process) and neomycin, kanamycin, and gentamicin biosynthesis (paromomycin involved).

The relative content of 26 kinds of differential metabolites across PI and PR were analyzed, and possible metabolic pathways were established for each substance involved (Fig. 4c and d). Results showed that the levels of acids, such as hexanoic acid, acetic acid, and heptanoic acid were higher in PI than in PR. The mainly metabolic pathway that these differential metabolites involved were microbial metabolism in diverse environments. Substances involved in microbial metabolism in diverse environments and biosynthesis of secondary metabolites were up-regulated in PI.

In the comparison of the ADS21-extracted phase from PI and PR samples, 12 key differential metabolites were identified between the PI and PR samples, and di-n-octyl phthalate was most significantly different (VIP = 5.38322). Among the 12 differential metabolites, six were upregulated and six downregulated in the PI sample, and the same trend was also observed in the PR samples (Fig. 5a and b, Table 6). These metabolites may be
### Table 5
The biomarkers between PI and PR samples absorbed by ADS-8 resin in rhizosphere soils

| Name                                      | KEGG   | VIP     | PI   | PR   | Participating in metabolic pathways                                                                 |
|-------------------------------------------|--------|---------|------|------|-----------------------------------------------------------------------------------------------------|
| Hexanoic acid                             | C01585 | 2.70187 | U    | D    | ko01120 Microbial metabolism in diverse environments (6)                                           |
| Deoxyxpergualin                           | D08032 | 1.69418 | U    | D    | cpd:C00033 Acetate                                                                                    |
| Acetic acid                               | C00033 | 1.67186 | U    | D    | cpd:C00414 Cyclohexanone                                                                             |
| Carbominal                                | D02619 | 1.62782 | D    | U    | cpd:C06104 Adipate                                                                                    |
| 2-Phenoxethanol                           | D08359 | 1.59034 | D    | U    | cpd:C06712 Fluoren-9-one                                                                             |
| Hexahydropyrindone                        | C01746 | 1.55401 | D    | U    | cpd:C09815 Benzamide                                                                                  |
| Acetamide                                 | C06244 | 1.54784 | D    | U    |                                                                                                      |
| Cucurbitacin b                            | C08794 | 1.54166 | U    | D    | ko00960 Tropane, piperidine and pyridine alkaloid biosynthesis (1)                                      |
| Methyl 3-O-mesyl-5-O-methoxybenzoyl-d-xyluronic acid | C00049 | 1.52048 | U    | D    |                                                                                                      |
| Pyridine                                  | C00747 | 1.41385 | D    | U    | ko00051 Fructose and mannose metabolism (1)                                                           |
| Thymol                                    | C09908 | 1.4007  | D    | U    |                                                                                                      |
| 3-Buten-2-one                             | C20701 | 1.37027 | D    | U    | ko00524 Neomycin, kanamycin and gentamicin biosynthesis (1)                                             |
| Hexanedioic acid                          | C06104 | 1.36746 | D    | U    |                                                                                                      |
| Cyclohexanone                             | C00414 | 1.3182  | U    | D    |                                                                                                      |
| 9-Fluorenone                              | C06712 | 1.27628 | D    | U    |                                                                                                      |
| Pentanoic acid                            | C00803 | 1.25499 | D    | U    |                                                                                                      |
| Paromomycin                               | C00832 | 1.20995 | D    | U    |                                                                                                      |
| d-Mannitol                                | C00392 | 1.20081 | D    | U    |                                                                                                      |
| (+−)-2-(4′-Isobutylphenyl) propionitrile   | C04469 | 1.13964 | D    | U    |                                                                                                      |
| Heptanoic acid                            | C17714 | 1.11581 | U    | D    |                                                                                                      |
| Acetohydroxamic acid                      | C06808 | 1.04682 | D    | U    |                                                                                                      |
| phenylamino                               | C01302 | 1.03838 | D    | U    |                                                                                                      |
| Butane                                    | D03186 | 1.034   | U    | D    |                                                                                                      |
| 2-Cyclohexenone                           | C02395 | 1.03148 | D    | U    |                                                                                                      |
| Ascaridole epoxide                        | C09836 | 1.01456 | U    | D    |                                                                                                      |
| benzamide                                 | C09815 | 1.00777 | D    | U    |                                                                                                      |

Note: U upregulated expression. D downregulated expression

### Table 6
The biomarkers between PI and PR samples absorbed by ADS-21 resin in rhizosphere soil

| Name                                      | KEGG   | VIP     | PI   | PR   | Participating in metabolic pathways                                                                 |
|-------------------------------------------|--------|---------|------|------|-----------------------------------------------------------------------------------------------------|
| Di-n-octyl phthalate                       | C14227 | 5.38322 | D    | U    | ko01100 Metabolic pathways (2)                                                                      |
| Tetradecanoic acid                        | C06424 | 1.55034 | D    | U    | cpd:C00829 Naphthalene                                                                             |
| Naphthalene                               | C00829 | 1.50684 | U    | D    | cpd:C06424 Tetradecanoic acid                                                                       |
| 10-Undecenoic acid                        | C13910 | 1.50193 | D    | U    | ko00310 Lysine degradation (1)                                                                      |
| Pentanedioic acid                         | C00489 | 1.48364 | U    | D    | cpd:C00489 Glutarate                                                                               |
| Ledol                                     | C09698 | 1.26306 | D    | U    | ko00627 Aminobenzoate degradation (1)                                                                |
| 3-Amino-2,3, dihydrobenzoic acid          | C12110 | 1.25025 | U    | D    | cpd:C16267 Cyclopropanecarboxylate                                                                 |
| Anhydro-5-mercapto-2-methyl-1,3,4-thiadiazolium | C08159 | 1.15087 | U    | D    |                                                                                                      |
| Cyclopropanecarboxylic acid               | C16267 | 1.14989 | U    | D    |                                                                                                      |
| Phosphorodithioic acid, O-(2,4-dichlorophenyl) O-ethyl S-propyl ester | C18405 | 1.11633 | D    | U    |                                                                                                      |
| Pyrrolidine-2,5-dione                     | C07273 | 1.081   | U    | D    |                                                                                                      |
| Crystalline Antibiotic                    | C15751 | 1.06066 | D    | U    |                                                                                                      |

Note: U upregulated expression. D downregulated expression
involved in processes such as lysine degradation (glutamate involved) and aminobenzoate degradation (cyclopropane carboxylate involved).

Similarly, the metabolic pathways of each differential metabolite in PI and PR were analyzed (Fig. 5c and d). The contents of pentanedioic acid and cyclopropanecarboxylic acid were higher in PI than in PR, except for several differential metabolites of unknown function, which were mainly involved in xenobiotic biodegradation and metabolism.

OPLS-DA analysis identified 18 and 17 differential metabolites in the 2 rhizosphere soils of the isogenic lines PI and PO by ADS8 and ADS21, respectively. The higher VIP value was methoprene (VIP = 1.85117) in ADS8 and di-n-octyl phthalate (VIP = 4.48732) in ADS21. The metabolites that were extracted by both resins possibly participate in microbial metabolism in diverse environments. At the same time, the nonpolar substances extracted by the ADS8 resin may also participate in steroid hormone biosynthesis and biosynthesis of antibiotics, whereas the polar substances extracted by ADS21 may participate in neomycin, kanamycin, and gentamicin biosynthesis (Figs. 6 and 7, Tables 7 and 8).

The relative content of 18 kinds of differential metabolites between PI and PO and the possible metabolic pathway with a single substance involved were analyzed (Fig. 6c and d), and the results showed that the levels of

**Fig. 8** Analysis of differential metabolites between Le and LO samples absorbed by ADS-8 resin from rhizosphere soil. a The relative content of the metabolites between Le and LO is shown in the heat map, and the differential substances are arranged according to the VIP values from small to large. Each row represents a compound. The value of each compound represents the relative content directly normalized on the scale of the graph. Blue represents low content while red represents high content. b The possible metabolic pathway distribution of single metabolic substance between Le and LO. c The left side represents the number of single differentia substance in the same pathway where Le is lower than LO, and the right side is the opposite.
pentanoic acid, paromomycin, and thioacetic acid in PI were higher than those in PO. There were more differential metabolites involved in microbial metabolism in diverse environments and xenobiotics biodegradation and metabolism, and these were down-regulated in PI. We compared with the metabolic pathways in which each differential metabolite extract by ADS21 resin might be involved in PI and PO (Fig. 7c and d) indicated that the content of d-glucose in PI was lower than that in PO. D-glucose is mainly involved in carbohydrate metabolism. Other differential metabolites and their possible metabolic pathways in PI and PO showed a similar trend.

Compared with the metabolites extracted by ADS8 of allelopathic rice PI and its transgenic rice PR and PO, we found that the unique metabolites of PR were paromomycin and benzamide, whereas that of PO were naphthalene and thiourea. When the PAL gene was overexpressed, the levels of seven substances decreased, which included dodecane, cucurbitacin b, and butane, whereas those of cyclopropanemethanol, 2-isopropylidene-alpha-methyl-, acetic acid, and pentadecane increased. At the same time, after PAL gene interference, the levels of five substances decreased (including cyclohexanone, paromomycin, pentanoic acid, thioacetic acid, and prenan-3,20-dione), whereas those of ascaridole epoxide and benzenemethanol increased.

Comparison of the ADS21-extracted phase from the PI, PR, and PO samples showed that the concentrations of 8 polar metabolites decreased after interference or overexpression of the PAL gene, e.g., methoprene, pyrrolidine-2,5-dione, and cyclohexane. However, the concentrations of tetradecanoic acid and 10-undecenoic acid increased with interference of the PAL gene. However, when the PAL gene was overexpressed, the levels of seven metabolites increased.

**Differences in metabolites from the rhizosphere soils of Le and LO**

Figure 8a and b and Table 9 indicated that of the differentially expressed metabolites identified, 14 were up-regulated and 10 were down-regulated in the Le samples, and a more significant differential expression was observed for metabolites 9-octadeconoic acid (VIP = 2.48717) and o-acetyl-L-serine (VIP = 2.03914). Analysis of the contents of 24 differential metabolites in Le and LO and the metabolic pathways in which single differential metabolites may be involved (Fig. 8c and d) showed that the contents of acids (such as 9-octadecanoic acid, heptanoic acid, and succinic acid), ethenone, and octane in Le were higher than in LO. The differential metabolites involved in the biosynthesis of plant secondary metabolites, xenobiotics biodegradation and metabolism, and lipid metabolism were upregulated in Le.

Figure 9a and b and Table 10 show that on the basis of metabolomics analysis, 26 differential metabolites were identified in the Le and LO samples, of which phenol and 9-octadecanoic acid compounds showed the most significant differences. The two metabolites were upregulated in

### Table 7 The biomarkers between PI and PO samples absorbed by ADS-8 resin in rhizosphere soils

| Name                                      | KEGG     | VIP     | PI   | PO   | Participating in metabolic pathways                                                                 |
|-------------------------------------------|----------|---------|------|------|-----------------------------------------------------------------------------------------------------|
| Methoprene                                | C14308   | 1.85117 | D    | U    | ko01120 Microbial metabolism in diverse environments (5)                                              |
| Cyclopropanemethanol, 2-isopropylidene-alpha-methyl- | C08159   | 1.55323 | D    | U    | cpdC00180 Benzoate                                                                                     |
| Abietate                                   | C06087   | 1.49409 | D    | U    | cpdC00511 Acrylic acid                                                                                |
| Pantaoic acid                              | C00803   | 1.47045 | U    | D    | cpdC00556 Benzyl alcohol                                                                              |
| Paromomycin                                | C00832   | 1.39737 | U    | D    | cpdC00829 Naphthalene                                                                                 |
| Naphthalene                                | C00829   | 1.3827  | U    | D    | cpdC16267 Cyclopropanecarboxylate                                                                      |
| Thioacetic acid                            | C01857   | 1.37916 | U    | D    | ko00140 Steroid hormone biosynthesis (1)                                                               |
| Benzoic acid                               | C00180   | 1.25548 | U    | D    | cpdC03681 Salpha-Pregnane-3,20-dione                                                                    |
| Propenooic acid                            | C00511   | 1.25174 | D    | U    | ko01130 Biosynthesis of antibiotics (1)                                                                 |
| Dodecane                                   | C08374   | 1.21753 | U    | D    | cpdC00832 Paromomycin                                                                                |
| Cyclopropanecarboxylic acid                | C16267   | 1.18536 | D    | U    |                                                                                                       |
| Pentadecane                                | C08388   | 1.12324 | D    | U    |                                                                                                       |
| 2,4-Dichlorophenethyl alcohol, n-propyl ether | C13240  | 1.11501 | U    | D    |                                                                                                       |
| Azelaic acid                               | C08261   | 1.09064 | D    | U    |                                                                                                       |
| Ascaridole epoxide                         | C09836   | 1.05321 | U    | D    |                                                                                                       |
| Pregnane-3,20-dione                        | C03681   | 1.02972 | U    | D    |                                                                                                       |
| Benzenemethanol                            | C00556   | 1.02626 | U    | D    |                                                                                                       |
| Thiourea                                   | C14415   | 1.01138 | D    | U    |                                                                                                       |

Note: U upregulated expression. D downregulated expression
the Le samples, but downregulated in the LO samples. Based on the analysis of the data of Le and LO (Figs. 9c and d), it was found that the content of differential metabolites involved in lipid metabolism, energy metabolism, biosynthesis of other secondary metabolites, biosynthesis of plant hormones, and amino acid metabolism in Le was higher than in LO, but the content of differential foreign bodies involved in biosynthesis of secondary metabolites and cellular community-prokaryotes was lower than in LO.

Comparison of possible pathways of metabolites between the two resins extracted showed that the substances extracted by ADS8 were more likely involved in the four pathways (biosynthesis of plant secondary metabolites, biosynthesis of secondary metabolites, energy metabolism, and lipid metabolism) than in ADS21, whereas the involvement of the two pathways (microbial metabolism in diverse environments and xenobiotics biodegradation and metabolism) decreased.

**Discussion**

Previous studies have shown that plants gather specific microbial communities and form unique rhizosphere microbial community structure through the mediation of root exudates [28–30]. The unique microecological environment is composed of plants, root exudates, and root microorganisms, which cause allelopathy [36–42]. Previous studies have demonstrated that the community and population of bacteria, fungi, and actinomycetes significantly differ in the rhizosphere soil of allelopathic and non-allelopathic rice under the mediation of different exudates [22, 32]. Accordingly, phenolic acids and terpenes could specifically aggregate certain microorganisms [43, 44]. Most of the existing studies have focused

![Fig. 9](image-url)

**Fig. 9** Analysis of differential metabolites between Le and LO samples absorbed by ADS-21 resin from rhizosphere soil. **a** The relative content of the metabolites between Le and LO is shown in the heat map, and the differential substances are arranged according to the VIP values from small to large. Each row represents a compound. The value of each compound represents the relative content directly normalized on the scale of the graph. Blue represents low content while red indicates high content. **b** The possible metabolic pathway distribution of single metabolic substance between Le and LO. The left side represents the number of single differentia substance in the same pathway where Le is lower than LO, and the right side is the opposite.
on the differences and functions of phenolic acids and terpenes, as well as their relationship with microorganisms in rice. Limited research has referred to other potential allelopathic substances, such as aromatic acid, benzene, derivatives, long-chain hydrocarbons, fatty acids, etc. [45].

Among these compounds, some long-chain fatty acids are known phytotoxins [1]. For example, nonanoic and decanoic acids have phytotoxic effects on algae as indicated by bioassays [46], and pelargonic acid (nonanoic acid) is regarded as a natural herbicide based on its particular phytotoxicity [47].

In the present study, the concentrations of several fatty acids varied among allelopathic potential rice lines. The resin adsorbed phase extracted from the rhizosphere soil of PI and Le at the five-leaf stage showed that the metabolites extracted from different allelopathic potential rice accessions were significantly different. The metabolites of allelopathic rice PI and non-allelopathic rice Le were analyzed and indicated that 21 different metabolites were extracted from rice rhizosphere soil by ADS-8 resin, of which acetic acid was the most significantly differentially expressed metabolite. The rhizosphere soil of allelopathic rice PI contained more acetic acid than non-allelopathic rice Le. Acetic acid is a weak carboxylic acid; in wheat, the anaerobic decomposition of wheat straw released acetic acid and this compound exhibited phytotoxicity [48].

The relative content of acetic acid was higher in the high allelopathic rice than the low allelopathic rice, which may be associated with rice allelopathy, and the compound could be generated from the anaerobic reaction and play a role in inhibiting weed growth. It is also regarded as a precursor in the process of biosynthesizing phenolic acids, terpenes and other metabolites [49].

In addition, long-chain fatty acids have also been documented as allelochemicals, with maize straw decomposed products showing allelopathic promotion or inhibition of the soil-born microorganisms, and several fatty acids were identified from these decomposed compounds, which included hexanoic acid (1.73%), 8-octadecenoic acid (1.06%), and 3-(4-hydroxy-3-methoxyphenyl)-2-propenoic acid (1.04%) [50]. The content of fatty acids were differed in rhizosphere soil of PI and Le, which would lead to the variations in allelopathic inhibition activity on weeds. Since PI and Le have different genetic backgrounds, when it turns to a same rice line with PAL overexpressed or silenced, the fatty acids from these transformed rice lines also differed to the WT, which indicated that changes in gene expression of PAL in rice results in changes in fatty acid contents in their rhizosphere and suggesting a putative link between PAL expression and fatty acids. The fatty acids would impact on the soil environment, since hexadecanoic acid from the root exudates of peanut (Arachis hypogaea L.) has close relationship with its soil sickness, high

Table 8 The biomarkers between PI and PO samples absorbed by ADS-21 resin in rhizosphere soil

| Name                        | KEGG     | VIP     | PI     | PO     | Participating in metabolic pathways |
|-----------------------------|----------|---------|--------|--------|------------------------------------|
| Di-n-octyl phthalate        | C14227   | 4.48732 | D      | U      | ko01120 Microbial metabolism in diverse environments (6) cpdC00033 Acetate |
| 1,2-Benzenedicarboxylic acid| C01606   | 2.85069 | D      | U      | cpdC00218 Methylamine               |
| Hexamethylphosphoramide     | C19250   | 1.50382 | D      | U      | cpdC01606 Phthalate                 |
| Anhydro-5-mercapto-2-methyl-3-phenyl-1,3,4-thiadiazolium | C08159 | 1.33164 | U      | D      | cpdC09822 Cyclohexane-1-carboxylate |
| Cyclopropanecarboxylic acid | C16267   | 1.33051 | D      | U      | cpdC11249 Cyclohexane               |
| Cyclohexanecarboxylic acid  | C09822   | 1.28803 | D      | U      | cpdC16267 Cyclopropanecarboxylate   |
| Pyrrolidine-2,5-dione       | C07273   | 1.2508  | U      | D      |                                    |
| d-Glucose                   | C00031   | 1.22743 | D      | U      | ko00524 Neomycin, kanamycin and gentamicin biosynthesis (1) cpdC00031 D-Glucose |
| Hexanoic acid               | C01585   | 1.17727 | D      | U      |                                    |
| Heptanal                    | C14390   | 1.16578 | D      | U      |                                    |
| Monothioglycerol            | D05075   | 1.12844 | U      | D      |                                    |
| 1,4-Methanoazulen-7-ol      | C09631   | 1.08899 | D      | U      |                                    |
| Methoprene                  | C14308   | 1.0807  | U      | D      |                                    |
| Acetic acid                 | C00033   | 1.03811 | D      | U      |                                    |
| Methanamine                 | C00218   | 1.03708 | U      | D      |                                    |
| Geldanamycin                | C11222   | 1.02144 | D      | U      |                                    |
| Cyclohexane                 | C11249   | 1.00451 | U      | D      |                                    |

Note: U upregulated expression. D downregulated expression
concentration (160 mg/kg soil and 240 mg/kg soil) of hexadecanoic acid would suppress the soil enzyme activity, which in turn to reduce root activity and the chlorophyll content in peanut leaves [51]. The hexadecanoic acid in the rhizosphere soil of PI and LO extracted by ADS21 resin was higher than that in Le. Fischer et al. also believed that long-chain fatty acids may interact with plant lipids (sterols etc.) to form micelle as allelochemicals [52]. Therefore, it was speculated that long-chain fatty acids act roles in directly impact on the soil microorganism, or interacting with other metabolites to form mixture with alleloapthic activity.

In terms of other metabolites, except for fatty acids, many are also involved in microbial or plant metabolism. Phenol, for example, is the metabolic substrate of various microorganisms, such as Pseudomonas [53], Acinetobacter [54], and Rhodococcus [55]. Wild watermelon and Cucurbita maxima could produce cucurbitacin b, which could inhibit cancer cells and Meloidogyne species population densities [56, 57]. Glomerella cingulata could transform ledol [58]. The differences in these substances may lead to different allelopathic potentials in rice. However, the relationship between these substances and rice allelopathy requires further investigation.

However, the exact role of long-chain fatty acids and acetic acid on the diversities of microbial community in the rhizosphere and their correlation with weed suppression still needs in-depth study.

**Conclusions**

Our study showed that the rhizosphere soil of allelopathic rice accumulates more non-polar metabolites than polar metabolites, and fatty acids are regarded as vital compounds, which is different from the well-known allelochemicals, including phenolic acids and terpenoids. The fatty acids might be indispensable for the activity of rhizospheric microorganisms, which helps in constructing the micro-environment to assist in maintaining allelopathic activity.
Methods
Plant growth conditions
This experiment was conducted in 2016 at Fujian Agriculture and Forestry University, Fuzhou, China, where temperature varied from 25 °C to 30 °C, while humidity varied from 65 to 78% during the experiment. The site is the outdoor network room experimental field. The globally known PI312777, the stronger allelopathic rice (PI, introduced from the USA), its transgenic lines with \( \text{PAL2}^{-1} \) overexpression (PO) and with \( \text{PAL2}^{-1} \) interfered expression (PR), non-allelopathic rice Lemont (Le, introduced from the USA), and its transgenic line with \( \text{PAL2}^{-1} \) overexpression (LO) were used for this study. All of the tested materials were genetically stable ones selected for multiple generations provided by Institute of Agroecology, Fujian Agriculture and Forestry University.

The dry-raised seedlings were transplanted following the method as described by Li et al. [59]. The field soil was sandy loam, and there was 1.72 g·kg\(^{-1}\) total nitrogen, 60.67 mg·kg\(^{-1}\) alkali hydrolysable nitrogen, 0.67 g·kg\(^{-1}\) total phosphorus, 30.45 mg·kg\(^{-1}\) available phosphorus, 1.53 g·kg\(^{-1}\) total potassium, 206.48 mg·kg\(^{-1}\) available potassium, and 21.3 g·kg\(^{-1}\) organic matter, with pH 6.13 in the tillage layer. The growth of rice seedlings at seedling stage was observed in random plants, and the soil sampling was taken at the five-leaf stage.

Rice seedlings were transplanted on March 25, 2016 with appropriate basal fertilizers (70% of total N as basal dressing and top dressing, in total N = 225 kg·hm\(^{-2}\), N: P: K = 1:0.5: 0.8). Each variety was planted separately in different plots (3 m \( \times \) 1 m) with a seeding rate of 150 g/m\(^2\). In addition, CK was set as the blank control soil in the same area without rice

| Name                                | KEGG     | VIP      | Le  | LO  | Participating in metabolic pathways                                                                 |
|-------------------------------------|----------|----------|-----|-----|-----------------------------------------------------------------------------------------------------|
| Phenol                              | C00146   | 2.82987  | U   | D   | ko01120 Microbial metabolism in diverse environments (8)                                              |
| 9-octadecanoic acid                 | C01530   | 2.27918  | U   | D   | cpd:C00049 L-Aspartate                                                                                |
| Isobutyric acid                     | C02632   | 2.19198  | U   | D   | cpd:C00084 Acetaldehyde                                                                                 |
| Carboxic acid                       | C01563   | 2.07028  | U   | D   | cpd:C00146 Phenol                                                                                      |
| Cyclopentanol                       | C02020   | 1.96481  | D   | U   | cpd:C00180 Benzoate                                                                                     |
| m-Cresolic acid                     | C14103   | 1.94407  | U   | D   | cpd:C00472 2-Methylpropanoate                                                                          |
| 3,5-Cyclohexadiene-1,2-dione        | C02351   | 1.67809  | U   | D   | cpd:C02632 2-Methylpropanoate                                                                          |
| cis-2-Dodecenoic acid               | C21202   | 1.6433   | D   | U   | cpd:C16267 Cyclopropanoic acid                                                                          |
| Hexadecanoic acid                   | C00249   | 1.4779   | U   | D   | cpd:C00249 Hexadecanoic acid                                                                            |
| Decane dioic acid                   | C08277   | 1.43455  | U   | D   | cpd:C01530 Octadecanoic acid                                                                            |
| Propenal                            | C01471   | 1.43094  | U   | D   | cpd:C01471 Acrolein                                                                                     |
| Hexahydropyridine                   | C01746   | 1.41484  | U   | D   | cpd:C01471 Acrolein                                                                                     |
| Ethanone                            | C00084   | 1.39731  | U   | D   | cpd:C00193 Benzaldehyde                                                                                 |
| Benzaldehyde                        | C01913   | 1.36669  | U   | D   | cpd:C00193 Benzaldehyde                                                                                 |
| Butane                              | D03186   | 1.344    | U   | D   | cpd:C00193 Benzaldehyde                                                                                 |
| Purine-2-acetamide, trans-2,7-Di-  | C01500   | 1.29879  | U   | D   | Carbonate                                                                                               |
| methyl-3,6-octadien-2-ol            |          |          |     |     | cpd:C00249 Hexadecanoic acid                                                                            |
| Acetamide                           | C06244   | 1.27783  | U   | D   | cpd:C00249 Hexadecanoic acid                                                                            |
| Cyclobarbital                       | D07323   | 1.22403  | D   | U   | cpd:C00249 Hexadecanoic acid                                                                            |
| 2-Butenolic acid                    | C01771   | 1.20318  | D   | U   | cpd:C00249 Hexadecanoic acid                                                                            |
| Glycol glycolate                    | C02037   | 1.16897  | U   | D   | cpd:C00249 Hexadecanoic acid                                                                            |
| 2,5-cyclohexadiene-1,4-dione        | C00472   | 1.12428  | U   | D   | cpd:C00249 Hexadecanoic acid                                                                            |
| 5-Cholesterol-3,26-diol             | C17336   | 1.10082  | D   | U   | cpd:C00249 Hexadecanoic acid                                                                            |
| Dodecanoic acid                     | C02679   | 1.09957  | D   | U   | cpd:C00249 Hexadecanoic acid                                                                            |
| Benzenene                           | C01407   | 1.05894  | D   | U   | cpd:C00249 Hexadecanoic acid                                                                            |
| Cyclopropanoic acid                 | C16267   | 1.05361  | D   | U   | cpd:C00249 Hexadecanoic acid                                                                            |
| Benzoic acid                        | C00180   | 1.04824  | D   | U   | cpd:C00249 Hexadecanoic acid                                                                            |

Note: U upregulated expression, D downregulated expression
planting. After the emergence of rice seedlings, the recommended field management practices were followed.

**Soil sample collection**
Random sampling was conducted in field at the five-leaf stage of rice seedlings. The sample collection method was followed by the method with slightly improvement [60]. Briefly, the rhizospheric soil was collected by shaking the roots to remove soil lumps sticking to the roots, and the remaining soil clinging to the roots was removed with forces and put into a 50 mL centrifuge tube. Then 50 mL of PBS solution (pH = 7.0) was added and ultrasonically oscillated for 10 min. Subsequently, the plant materials were removed from the resulting solution after centrifugation. Lastly, the control samples were collected from uncultivated soil at the depth of 0–5 cm.

**Extraction and identification of metabolites from rhizosphere soils of different rice accessions**
Six samples were taken from the rhizosphere soil of each treatment. Ten grams of each sample was added with 30 mL of sterile water, and oscillated at 200 rpm at 20°C for 3 h, then centrifuged three times at 12,000×g, for 10 min. Finally, the supernatants were pooled, mixed well, and divided evenly into two portions.

The resin ADS-8 (weakly polar resin) and ADS-21 (polar resin) (purchased from Tianjin Nankai Hecheng Co., Ltd.), respectively, were weighed with 100 g each. After pretreatment, these were added to the above water extracts at 20°C and then oscillated at 200 rpm for 24 h. Later, the resin was removed, 100 mL of methanol was added for elution, and then oscillated at 200 rpm at 25°C for 2 h with three repeats, then the methanol phase was combined and vacuum concentrated to dry. Then, 1 mL of methanol was added to dissolve resin, filtered across a 0.22 mm mixed cellulose membrane, and then the metabolites were analyzed.

A Shimadzu GCMS-TQ8040 Gas Chromatography/Triple four-stage bar mass spectrometer was used to analyze the above processed samples using the following conditions: The four-stage bar mass spectrometer was used to analyze the compounds in the absorption phase by ADS-8 resin from the rhizosphere soils of different allelopathic potential rice accessions and OPLS-DA evaluation model was established. The OPLS-DA score of the compounds in the absorption phase by ADS-8 resin from the rhizosphere soils of different allelopathic potential rice accessions. Note: The numbers 1, 2, and 3 in the figure represent three parallel duplicates of the same sample. PI stands for allelopathic rice PI312227 (PI); PAL for PAL2–1 inhibited transgenic line PR; O for the transgenic line PO of PAL2–1 overexpressed in allelopathic rice PI312227 (PI); Le for non-allelopathic rice Lemont (Le); LOP for the transgenic line LO of PAL2–1 overexpressed in non-allelopathic rice Le.

**Additional file 3: Figure S3.** Principal component analysis of the compounds absorbed by ADS-21 resin from the rhizosphere soils of different potential allelopathic rice accessions. Note: The numbers 1, 2, and 3 in the figure represent three parallel duplicates of the same sample. PI stands for allelopathic rice PI312227 (PI); PAL for PAL2–1 inhibited transgenic line PR; O for the transgenic line PO of PAL2–1 overexpressed in allelopathic rice PI312227 (PI); Le for non-allelopathic rice Lemont (Le); LOP for the transgenic line LO of PAL2–1 overexpressed in non-allelopathic rice Le. Picture order from top to next: PI-Le, PI-PR, PI-PO, Le-LO, PO-LO.

**Additional file 4: Figure S4.** Metabolomics analysis was conducted to identify the soil substances in the extracted phase by ADS-8 resin from the rhizosphere soils of different allelopathic potential rice accessions and OPLS-DA evaluation model was established. The OPLS-DA score of the compounds in the absorption phase by ADS-8 resin from the rhizosphere soils of different allelopathic potential rice accessions. Note: The numbers 1, 2, and 3 in the figure represent three parallel duplicates of the same sample. PI stands for allelopathic rice PI312227 (PI); PAL for PAL2–1 inhibited transgenic line PR; O for the transgenic line PO of PAL2–1 overexpressed in allelopathic rice PI312227 (PI); Le for non-allelopathic rice Lemont (Le); LOP for the transgenic line LO of PAL2–1 overexpressed in non-allelopathic rice Le. Picture order from top to next: PI-Le, PI-PR, PI-PO, Le-LO, PO-LO.

**Additional file 5: Figure S5.** The OPLS-DA score of the compounds absorbed by ADS-21 resin from the rhizosphere soils of different allelopathic potential rice accessions. It shows that the predictive parameters of the OPLS-DA model, R2X, R2Y and Q2, were both greater than 0.9, suggesting that the model is excellent and suitable for further analysis of differential metabolites. Note: The numbers 1, 2, and 3 in the figure represent three parallel duplicates of the same sample. PI stands for allelopathic rice PI312227 (PI); PAL for PAL2–1 inhibited transgenic line PR; O for the transgenic line PO of PAL2–1 overexpressed in allelopathic rice PI312227 (PI); Le for non-allelopathic rice Lemont (Le); LOP for the transgenic line LO of PAL2–1 overexpressed in non-allelopathic rice Le. Picture order from top to next: PI-Le, PI-PR, PI-PO, Le-LO, PO-LO.

**Abbreviations**
PAL: Phenylalanine ammonia-lyase gene; PAL2–1: Phenylalanine ammonia-lyase (PAL) gene 2–1; WT: Wild-type; PI: PI312777, the stronger allelopathic rice; PR: Transgenic line with major allelopathic gene PAL2–1 inhibited of PI; PO: Transgenic line with PAL2–1 overexpression of PI; Le: Non-allelopathic rice Lemont; LO: Transgenic line with PAL2–1 overexpression of Le; CK: The blank control soil; KEGG: Kyoto Encyclopedia of Genes and Genomes; PBS solution: Phosphate buffer saline; PCA: Principal Component Analysis; OPLS-DA: Orthogonal partial least squares discriminant analysis; VIP: Projection of variable importance of OPLS-DA model

**Acknowledgements**
Thanks to Dr. Fang Changxun from Fujian Agriculture and Forestry University for the methodological guidance and suggestions for the revision of this article. Thanks to Dr. Li Zhou for his guidance and help on drawing figures in this article.
Authors' contributions
WXL, and YZL conceived the study; YZL and WXH wrote the manuscript; YZL performed the experiments; YZL and LNX conducted statistical analyses; PL has revised the manuscript. All of the authors discussed the results and commented on the manuscript, and approved its submission for potential publication.

Funding
This work was supported by the National Key Research and Development Program of China (2016YFD0300508), the National Natural Science Foundation of China (81573530, 31271670), Fujian-Taiwan Joint Innovative Center for Germplasm Resources and Cultivation of Crop (FJ 2011 Program, no. 2015–75), Foundation for the Science and Technology Innovation of Fujian Agriculture and Forestry University (KFX2015043, CXZX2018042, CXZX2017309). The funders had no role in study design, data collection and analysis, data interpretation, or in writing of the manuscript.

Availability of data and materials
All data generated during this study are included in this published article and its supplementary information files, and the raw data used or analysed during the current study available from the corresponding author on reasonable request.

Ethics approval and consent to participate
during the current study available from the corresponding author on reasonable request.

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Author details
1 Fujian Provincial Key Laboratory of Agroecological Processing and Safety Monitoring, College of Life Sciences, Fujian Agriculture and Forestry University, Fuzhou 350002, Fujian, People’s Republic of China. 2 Key Laboratory of Ministry of Education for Genetics, Breeding and Multiple Utilization of Crops, College of Agriculture, Fujian Agriculture and Forestry University, Fuzhou 350002, P. R. China. 3 Crop Science Department, National University of Leshoto, Maseru 100, Leshoto. 4 Key Laboratory of Crop Ecology and Molecular Physiology, Fujian Agriculture and Forestry University), Fujian Province University, Fuzhou 350002, P. R. China.

Received: 17 October 2019 Accepted: 26 May 2020

References
1. Rice EL. Allelopathy (second edition). Physiological Ecology. Orlando Academic Press; 1984.
2. Callaway RM, Maron JL. What have exotic plant invasions taught us over the past 20 years? Trends Ecol Evol. 2006;21(7):369–74.
3. Fu BJ, Li SG, Yu XB, Yang P, Yu QR, Feng QG, Zhuang XL. Chinese ecosystem research network: progress and perspectives. Ecol Complex. 2010;7(2):225–33.
4. Lin WX, Fang CX, Chen T, Lin RX, Xiong J, Wang HB. Rice allelopathy and its properties of molecular ecology. Front Biol. 2010;5(3):255–62.
5. Mallik AU. Challenges and opportunities in allelopathy research: a brief overview. J Chem Ecol. 2000;26(9):2007–9.
6. Olofsdotter M, Jensen LB, Courtois B. Improving crop competitive ability using allelopathy—an example from rice. Plant Breed. 2002;121(1):1–9.
7. Olofsdotter M. Rice—a step toward use of allelopathy. Agron J. 2001;93(1):3–8.
8. Chou CH, Jiang YC, Chfng HH. Phenolic compounds in a series of allelopathic and non-allelopathic rice root exudates. J Plant Physiol. 2004;160(8):1647–62.
9. Olofsdotter M, Rebulanan M, Madrid A, Wang D, Navared D, Olk DC. Why phenolic acids are unlikely primary Allelochemicals in Rice. J Chem Ecol. 2002;28:229–42.
10. Kato-Noguchi H. Allelopathic substance in rice root exudates: rediscovery of momilactone B as an allelochemical. J Plant Physiol. 2004;161(3):271–6.
11. Kato-Noguchi H, Ino T. Concentration and release level of momilactone B in the seedlings of eight rice cultivars. J Plant Physiol. 2005;162(9):965–9.
12. Kato-Noguchi H. Molecular evolution of momilactone a and B: potent allelochemicals, momilactones have only been found in rice and the moss Hyphnum plumifforme; 2011.
13. Kato-Noguchi H. Convergent or parallel molecular evolution of momilactone a and B: potent allelochemicals, momilactones have been found only in rice and the moss Hyphnum plumifforme. J Plant Physiol. 2011;168(13):1511–6.
14. Lin WX, Fang CX, Wu LX, Lin S. Research on and application of rice allelopathy and crop allelopathic autotoxicity in China. In: Luo SM, Glessner SR, editors. Agroecology in China: science, practice, and sustainable management; 2017. p. 161–96.
15. Zhang Q, Li L, Li JF, Wang HB, Fang CX, Yang XY, He HB. Increasing rice allelopathy by induction of barnyard grass (Echinochloa crus-galli) root exudates. J Plant Growth Regul. 2018;37(3):745–54.
16. Fang CX, Li YZ, Li CX, Li BL, Ren YJ, Zheng HP, Zeng XM, Shen LH, Lin WX. Identification and comparative analysis of microRNAs in barnyardgrass (Echinochloa crus-galli) in response to rice allelopathy. Plant Cell Environ. 2015;38(7):1368–81.
17. Niklas S, Claude B. Allelopathic Plants: Models for Studying Plant-Interkingdom Interactions. Trends Plant Sci. Author links open overlay panelNiklasSchandy1ClaudeBecker2. 2020;25(2):176–85.
18. Xu MM, Galhano R, Wielmann P, Bueno E, Tiernan M, Wu W, Chung IM, Gershenzon J, Tudzynski B, Sesma A, Peters R. Genetic evidence for natural product-mediated plant-plant allelopathy in rice (Oryza sativa). New Phytol. 2012;193(3):570–5.
19. Fang CX, Zhuang YE, Xu TC, Li YZ, Li Y, Lin WX. Changes in Rice Allelopathy and Rhizosphere microflora by inhibiting Rice phenylalanine ammonia-lyase gene expression. J Chem Ecol. 2013;39(2):204–12.
20. Song BX, Xiong CJ, Fang CX, Li YL, Lin RX, Liang YY, Lin WX. Rice enhanced differential and gene expression in rice under low nitrogen treatment. J Chem Ecol. 2008;34(5):688–95.
21. Fang CX, Xiong J, Qiu L, Wang HB, Song BX, He HB, Lin RX, Lin WX. Analysis of gene expressions associated with increased allelopathy in rice (Oryza sativa L) induced by exogenous salicylic acid. Plant Growth Regul. 2009;57(2):163–72.
22. He HB, Wang HB, Fang CX, Wu HW, Guo XQ, Liu CH, Lin ZH, Lin WX. Barnyard grass stress up regulates the biosynthesis of phenolic compounds in allelopathic rice. J Plant Physiol. 2012;169(17):1747–53.
23. Dilday RH, Lin J, Yan W. Identification of allelopathy in the USDA-ARS rice germplasm collection. Aust J Exp Agric. 1994;34(7):907–10.
24. Lin RX, Rong H, Zhou JJ, Yu CP, Ye CY, Chen LS, Lin WX. Impact of allelopathic rice seedlings on rhizospheric microbial populations and their functional diversity. Acta Ecol Sin. 2007;27(6):3644–54.
25. Kong CH, Wang P, Zhao H, Xu XH, Zhu YD. Impact of allelochemical exuded from allelopathic rice on soil microbial community. Soil Biol Biochem. 2008;40(7):1862–9.
26. Qu XH, Wang JG. Effect of amendments with different phenolic acids on soil microbial biomass, activity, and community diversity. Appl Soil Ecol. 2008;39(2):172–9.
27. Vivanco JM, Bais HP, Sternitz FR, Thelen GC, Callaway RM. Biogeographical variation in community response to root allelochemistry: novel weapons and exotic invasion. Ecol Lett. 2004;7(4):285–92.
28. Xiong J, Lin HF, Li ZF, Fang CX, Han QD, Lin WX. Analysis of rhizosphere microbial community structure of weak and strong allelopathic rice varieties under dry paddy field. Acta Ecol Sin. 2012;32(19):5009–9 (in Chinese with English abstract).
29. Ahwood JW, De Vos RCH, Moing A, Deborde C, Erban A, Kopka J, Gisdace R, Hall RD. Chapter sixteen - plant metabolomics and its potential for systems biology research: background concepts, Technology, and Methodology. Methods Enzymol. 2011;500:299–36.
35. He HB, Wang HB, Chen XX, Lin WX, Jia XL, Fang CX, Gan QF, Ni NN, Wu WX. Allelopathic effects of aqueous extracts from different parts and root exudates of rice on barnyardgrass. Chin J Eco-Agri. 2007;02:14–7 (in Chinese with English abstract).

36. Weston LA, Duke SO. Weed and crop allelopathy. Crit Rev Plant Sci. 2003;11:367–89.

37. Alsadawi IS, Rice EL, Kars TKB. Allelopathic effects of Polygonum aviculare L. III. Isolation, characterization, and biological activities of phytoxins other than phenols. J Chem Ecol. 1983;9:761–74.

38. Berendt RL, Pleterse CJ, Bakker PM. The rhizosphere microbiome and plant health. Trends Plant Sci. 2012;17(8):478–86.

39. Doorbos RS, van Loon LC, Bakker PM. Impact of root exudates and plant defense signaling on bacterial communities in the rhizosphere. A review. Agronomy Sustainable Development. 2012;32(1):227–43.

40. Kamlova F, Kravchenko LV, Shaposhnikov AI, Azarova T, Makarova N, Lugtenberg B. Organic acids, sugars, and L-tryptophan in exudates of vegetables growing on stonewool and their effects on activities of rhizosphere bacteria. Mol Plant-Microbe Interact. 2006;19(3):250–6.

41. Li JY, Lin SX, Zhang Q, Li L, Hu WW, He HB. Phenolic acids and terpenoids in the soils of different weed-suppressive circles of allelopathic rice. Arch Agron Soil Sci. 2019;1–13.

42. Zhang SS, Zhu WJ, Wang B, Tang JJ, Chen X. Secondary metabolites from the invasive Solidago canadensis L. accumulation in soil and contribution to inhibition of soil pathogen Pythium ultimum. Appl Soil Ecol. 2014;83:280–6.

43. Bacilio-Jiménez M, Aguilar-Flores S, Ventura-Zapata E, Pérez-Campos E, Bouquelet S, Zenteno E. Chemical characterization of root exudates from rice (Oryza sativa) and their effects on the chemotactic response of endophytic bacteria. Plant Soil. 2003;249(2):271–7.

44. Kong CH, Xu XH, Zhou B, Hu F, Zhang CX, Zhang MX. Two compounds from allelopathic rice accession and their inhibitory activity on weeds and fungal pathogens. Phytochemistry. 2004;65(8):1123–8.

45. Rimando AM, Duke SO. Studies on rice allelochemicals. Rice: origin, history, technology and production. 2003. p. 221–44.

46. McCraken MD, Middaugh RE, Middaugh RS. A chemical characterization of an algal inhibitor obtained from Chlamydomonas. Hydrobiologia. 1980;70:271–6.

47. Izyk GP, Zomer P, Kern A. A new contact herbicide based on naturally-occurring pelargonic acid. Weed Science Society America, Abstract Book, vol. 37. 1997. p. 260.

48. Lynch JM. Phytotoxicity of acetic acid produced in anaerobic decomposition of wheat straw. The J Appl Bacteriol. 1977;42:81–7.

49. Wang L, Shi LL, Zhang YX, Liu YJ. Biosynthesis and regulation of the secondary metabolites in plants. J Wuhan Botanical Res. 2007;25(5):500–8 (in Chinese with English abstract).

50. QI YZ, Zhen WC, Li HY. Allelopathy of decomposed maize straw products on three soil-borne diseases of wheat and the analysis by GC-MS. J Integr Agric. 2015;14(1):88–97.

51. Liu P, Liu ZH, Wang CB, Guo F, Wang M, Zhang YF, Dong L, Wan SB. Effects of three long-chain fatty acids present in peanut (Arachis hypogaea L.) root exudates on its own growth and the soil enzymes activities. Allelopath J. 2012;29(1):13–24.

52. Fischer NH, Quijano L. Allelopathic agents from common weeds: Amaranthus palmeri, Amaranthus antirrhinifolius, and related weeds. Chem Allelopathy. 1985;9:133–47.

53. Hill GA, Robinson OW. Substrate inhibition kinetics: Phenol degradation by Pseudomonas putida. Biotechnol Bioeng. 1975;17(11):659–615.

54. Hao OJ, Kim MH, Seagren EA, Kim H. Kinetics of phenol and chlorophenol utilization by Acinetobacter species. Chemosphere. 2002;46(6):797–807.

55. Liu XH, Liu ZP, Liu SJ. Functional identification of the gene locus ngc12319 and characterization of catechol 1,2-dioxogenase in Corynebacterium glutamicum. Biotechnol Lett. 2004;26(7):575–80.

56. Shadung KG, Mashela PW, Mphosi MS. Response of cucurbitacin B concentration in Nemafric-BL phytonematicide to increasing storage period. J Stored Prod Postharvest Res. 2016;32:6–2.

57. Ferguson JE, Metcalf RL, Fischer DC. Disposition and fate of cucurbitacin B in five species of diabroticites. J Chem Ecol. 1985;11(9):1307–21.

58. Miyazawa M, Uemura T, Kameoka H. Biotransformation of sesquiterpenoids,-(-)-globulol and (+)-ledol by Glomerella cingulata. Phytochemistry. 1994;37(4):1027–30.

59. Li Z, Li Z, Letumra P, Zhao H, Zhang ZY, Lin WW, Chen HF, Lin WX. A positive response of rice rhizosphere to alternate moderate wetting and drying irrigation at grain filling stage. Agric Water Manag. 2018;207:26–36.

60. Edwards J, Johnson C, Santos-Medellin C, Lurie E, Podishetty NK, Bhatnagar S, Eisen JA, Sundaresan V. Structure, variation, and assembly of the root-associated microbiomes of rice. Proc Natl Acad Sci U S A. 2015;112(8):E911–20.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Ready to submit your research? Choose BMC and benefit from:
- fast, convenient online submission
- thorough peer review by experienced researchers in your field
- rapid publication on acceptance
- support for research data, including large and complex data types
- gold Open Access which fosters wider collaboration and increased citations
- maximum visibility for your research: over 100M website views per year

At BMC, research is always in progress.

Learn more biomedcentral.com/submissions