IAA Producing Endophytic Fungus *Fusarium oxysporum* wlx Colonize Maize Roots and Promoted Maize Growth Under Hydroponic Condition

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Abstract

Indole-3-acetic acid (IAA) is known for their role in plant root interactions with microbial partners. Current study was focused on role of IAA as signal for colonization between endophytic fungus *Fusarium oxysporum* wlx and maize roots. Culture filtrate of the strain contained 31 μg/mL of IAA. Addition of tryptophan concentration ranging from 500 to 1000 μg/mL in culture medium significantly enhanced production of IAA by *Fusarium oxysporum* wlx. The strain effectively colonized the roots of maize and subsequently enhanced the growth and proliferation of host plant. In order to demined the role of IAA in root colonization by *F. oxysporum* we inhibited the biosynthesis of IAA by using IAA biosynthesis inhibitor which efficiently reduced te colonization of *F. oxysporum* in maize roots by 46% (foliar application) and 62% (root application) of the seedlings without yucasin (IAA biosynthesis inhibitor) treatment suggesting an IAA crosstalk between the two partners. Exogenous application of IAA restored the ability of endophyte *F. oxysporum* to colonize maize roots and significantly improved different growth parameters of maize seedlings. It is concluded that a molecular crosstalk of maize roots and endophytic *F. oxysporum* wlx is necessary for subsequent endophytic association between them.

Keywords: IAA; *Fusarium oxysporum*; Yucasin; Maize

Introduction

Colonization of endophytic fungi enhances the ecological adaptability of the plant by improving its tolerance against the biotic and abiotic stresses [1]. Endophytic fungi colonize the living plant tissue without causing damage, and include a very diverse group of fungi with different life history. Mutualistic endophytic fungi have been the focus of increasing interest worldwide, mainly because of their beneficial effects on plant suitability, such as tolerance to biotic and abiotic stresses, nutrient availability and increased growth [2]. Endophytic fungi provide a number of benefits to the host plants in various ways, such as promotion of growth, protection against diseases and pests, and augmenting absorption of minerals [3]. Fungal endophytes produce bioactive metabolites that promote the plant endophyte interaction [4]. Promoting plant growth is the most important effect of fungal symbiosis [5], however endophytic fungi promote plant growth by producing various secondary metabolites, including ammonia and plant hormones, particularly IAA [3,6]. Phytohormones are flexible signaling molecules affecting many physiological processes of plants development even when present at low concentrations [7]. The whole catalog of plant developmental process controls by hormones auxin and it is conceivable that some microbes also produce auxin [8]. Plants and microorganisms, including bacteria, algae and fungi, can produce IAA [9]. The role of microbial IAA in plant-microorganism interactions has received increasing attention recently. In addition, some studies show that IAA is a signaling molecule in microorganisms because it affects gene expression in several microorganisms [10]. Plants and their interactions with rhizosphere-related microbes such as fungi have always been an area of concern, as the knowledge of these processes may lead to environmentally-friendly farming practices. Roots produce a variety of organic compounds, including sugars, organic acids and vitamins [11]. These are then used as nutrients or signals for the fungal population. In contrast, fungi release iron carriers, volatile compounds, and plant hormones that may enhance plant growth, either directly or indirectly, by increasing the nutrient availability of their host [12]. IAA produced by fungi can induce lateral root formation and root hair development [13]. In addition, IAA produced by fungi can defeat pathogenic strains and disease progression by enhancing the plant’s immune response [14]. Under in vitro conditions, it has recently been recognized that various endophytic fungi produce GAs and indole-3-acetic acid (IAA). The biosynthesis of indole-3-acetic acid (IAA) has not been well developed in fungi like bacteria [15]. However, it has recently been established that certain fungi, including endophytes,
harbor them in their growth medium [16]. Indole-3-acetic acid (IAA) is considered to be a biologically active form of auxin its degree has been strongly regulated in plants through the interaction of several alleys, involving the biosynthesis, oxidation and hydrolysis of IAA, as well as the binding of IAA to macromolecules such as carbohydrates and amino acids [17]. Auxin is often suggested to play a role in the cross-talk between plant and fungal signaling during ectomycorrhizal establishment [18]. IAA is a plant hormone known to play an important role in cell elongation, division and cell enlargement [19]. We have published an article regarding in vitro growth promotion by endophytic Fusarium oxysporum wlw. Keeping the above discussion in view the current study was aim to identify the role of IAA in maize root colonization by endophytic Fusarium oxysporum wlw and host growth promotion under hydroponic condition.

Materials and Methods

The endopysts wlw was isolated from leaves of Withania sominfera and was identified as Fusarium oxysporum wlw [20]. The experiment was conducted in plant microbe interaction laboratory Abdul Wali Khan University Mardan, Pakistan. Maize variety GAUHER was obtained from national agriculture research center (NARC) Islamabad. The seedlings were allowed to grow for 14 days inside growth chamber (DAIHN Lab Tech) under controlled conditions (photoperiod 16/8 h and temperature 25°C at midday and 15°C at night; 390 ppm CO2; 40% humidity). The F. oxysporum was applied to seedlings after two days in Hoagland’s solution in the form of 1 mL of fungal spores suspension prepared by the hemacytometer and adjusted to an inoculum density of 106 mL⁻¹ spores.

Determination of Indole-3-acetic acid in culture filtrate

The isolated strain was screened for IAA production by subjecting the CF to calorimetric assay using Salkowski reagent (150 mL concentrated H₂SO₄, 250 mL distilled water, 7.5 mL 0.5 M FeCl₃.6H₂O) [21]. The strain was cultured in czapek broth media containing different concentrations of tryptophan (100, 500, 1000 µg/mL) at 28°C and 120 rpm for seven days. For comparison, endopysts were also cultured in czapek broth having no tryptophan. After seven days, the culture were harvested and filtered through Whatman No 1 filter paper. The culture filtrate was centrifuged at 11963 g for 2 min and to 1 mL of CF 2 mL of salkowski reagent was added. The salkowski reagent was allowed to react with the IAA (in CF) at room temperature for 30 min in the dark. After incubation period, intensity of the pink color was determined by taking optical density of the mixture against the blank (2 mL blank media and 2 mL salkowski reagent) at 540 nm using a uv-vis spectrophotometer (PerkinElmer Lambda 25). Straight line equation was derived from the standard curve produced by plotting standard IAA (Sigma Aldrich) concentrations (10, 25, 50, 100, 150, 200, 250, 300 µg/mL) against OD. The derived equation Y = 0.0081X-0.0151 was used to quantify IAA in the CF of endopysts.

Effect of yucasin on release of IAA in culture filtrate and biomass production

In order to examine the effect of yucasin (-S-(4-chlorophenyl)-1,2-dihydro-1,2,4-triazole-3-thione) auxin biosynthesis inhibitor, the isolated endophyte was grown in czapek medium containing (50 µM) of yucasin under conditions mentioned above. One week later, the cultures were collected and sieved with Whatman No 1 filter paper. The production of IAA in the filtrate was screened as previously described, and the biomass was collected and stored in a 50 mL Falcon tube at 60°C in an oven for 5 days. After five days, dry biomass was determined in the fractions.

Maize growth in hydroponic system

For the root colonization assay, 40 mL of half-strength Hoagland’s solution [22,23], was taken in a 100 mL beaker. The truncated micropipette tip was held in sterile cardboard and sealed with a pumpkin faucet as Hydroponics set up [24]. Maize seed was surface sterilized using HgCl₂ solution (0.1%) and washed three times with distilled water. After surface sterilization, 5 seeds were placed in an autoclave petri plates with 2x filter paper and incubated at 28°C for 5 days and allowed to germinate. The uniform seedlings were transferred to a hydroponic setting and the seedlings were inoculated with endophyte spores suspension in water adjusted to an inoculum density of 106 mL⁻¹ spores. After two weeks of growth, seedlings were harvested and various growth parameters i.e shoot and root length, and dry weights were measured.

Root colonization assay

To determine fungal colonization in maize root plants were collected after 14 days, washed with tap water, cut into small pieces and stained with lectophenol cotton blue dye (Cotton blue 0.05g, phenol crystals 20 g, glycerol 40 mL, lactic acid 20 mL, distilled water 20 mL) for 20 mins. After 20 min, the segments were washed and observe under light microscope. The level of colonization was quantified by plotting the root segments on PDA. Shortly, the root were surface sterilized with 0.1 % HgCl₂ and washed with distill water and cut into 1 cm fragments. Six root segments for each plant, 2 from upper part (near to inoculum) two from middle part and two from lower part (near to stem) were kept on clean filter paper and inoculated on PDA plates and incubated for one week. After incubation for a week, colonization percentage was determined by dividing number of root segments positive for colonization by total number of root segments studied and multiplied by 100. For comparison and validation of surface sterilization procedure, non-injured root segments were plated on PDA. Also, the root exudates of inoculated and non-inoculated seedlings were collected and screened for the determination of IAA, according to standard procedure as discussed previously.

Exogenous application of IAA

For the exogenous application of IAA, plants were grown in 40 mL half strength of Hoagland's solution in 100 mL beaker. After two days of growth in liquid solution, standard IAA at a final
concentration of 10 µg/mL was applied through ariel spray on plants [25]. The seedlings were then inoculated with fungi as described earlier and uninoculated seedlings were used for comparison. After 14 days of growth the seedlings were harvested and various growth constraints via shoot length, root length, and dry weights were measured and roots exudates were screened for IAA.

Inhibition of IAA biosynthesis by yucasin

Biosynthesis and exudation of IAA by the root was inhibited by applying yucasin (50 µM) [26], in the form of foliar spray. Root colonization by the endophyte was assayed in the presence of yucasin to assess the role of IAA in root-endophyte association.

Statistical analysis

Data was analyzed using SPSS for windows 16.0 (SPSS Inc., Chicago, IL, USA). An analysis of variance was performed to determine the effect of the treatment and the errors associated with the experiment. To identify significant differences between treatments, average comparisons were performed using the Duncan multiple-range test ($P=0.05$), where the error of the mean squared error was used to estimate the standard error of the difference between the means.

Results

Determination of IAA in culture filtrate of *F. oxysporum*

The strain *F. oxysporum* wlv was able to secrete significant quantity (31.6 µg/mL) IAA in their culture medium. The culture of endophyte wlv exposed to yucasin (IAA biosynthesis inhibitor) had significantly lower amount of IAA than their culture grown in the absence of this inhibitor. The amount of IAA released by endophyte was reduced by up to 37% in yucasin containing media than the control culture. Addition of different concentrations of tryptophan (500 and 1000 µg/mL) in the media affect the amount of IAA released by the endophyte wlv. Increasing tryptophan concentration from 500 to 1000 significantly enhanced the amount of IAA released in the culture by 17% and 25% respectively (Figure 1).

Endophyte growth response to exogenously applied yucasin

Application of yucasin was inhibitory to endophyte growth as indicated by lower dry biomass (Figure 2). Yucasin exposed culture had significantly lowered (30%) the biomass than in culture having no inhibitor.

Effect yucasin, IAA and endophyte *F. oxysporum* on maize growth

Exogenous application of yucasin in the form of foliar spray or in Hogland’s media (Y-R) retarded shoot length by 28% and 12% in comparison to control. As was expected, exogenous application of phytohormone IAA significantly enhanced shoot length by 40% as compared to control. Also the shoot growth of maize seedlings was enhanced significantly in media containing endophyte wlv. Endophyte wlv improved shoot length by 26% as compared to the control. As noted above, shoot growth declined in seedlings exposed to exogenously applied yucasin. However, endophyte wlv associated seedlings were resistant to this inhibitor and showed normal growth even in the presence of yucasin. As renowned above, seedlings growth was promoted by exogenous application of IAA. However growth of maize seedlings was enhanced further in media containing endophyte in binary combination with IAA. Exogenously applied IAA synergistically interacted with endophyte wlv in promoting shoot length as evident from up to 13% greater shoot length in media containing a binary combination of IAA and endophyte than the media containing IAA without endophyte (Figure 3a).

Root growth of maize seedlings was also negatively influenced by exogenously applied yucasin either as foliar spray or root entries. For instance, exogenous application of yucasin in the form of foliar spray caused 30% reduction in root length whereas root treatment of yucasin reduced root length by 31% in comparison to control. Exogenous application of IAA significantly improved root length by 22% as compared to control.

Root length was also significantly enhanced by endophyte wlv, where an increase of 60% over control was noticed. As noted above, exogenous application of yucasin declined seedlings root growth. However, endophyte associated seedlings were resistant to this inhibitor and showed normal growth even in the presence of yucasin. Similarly endophyte associated seedlings treated with foliar IAA had more roots than seedlings treated with foliar IAA alone (Figure 3b). Application of both foliar and root entries of yucasin reduced (69%) seedlings dry biomass in comparison to control. Likewise exogenously applied IAA interacted with endophyte wlv and enhanced dry biomass significantly. For instance, seedlings that received binary combination of endophyte wlv and foliar IAA had 56%, greater biomass than seedling receiving foliar IAA alone. Dry biomass was also enhanced significantly by the endophytes as compared to the control. Likewise seedling hosting endophyte *F. oxysporum*

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in binary combination with foliar yucasin had 27% greater biomass than foliar yucasin alone (Figure 3c).

**Figure 2.** Effect of yucasin on the biomass of endophytic fungi *F. oxysporum* w1w. Data are mean of 9 replicates from 3 independent experiments with standard error bars. Bars (same group) labelled with different letters are significantly different (Duncan test; p<0.05).

**Figure 3.** Effect of endophyte association (*F. oxysporum*), IAA (foliar), yucasin (foliar, F and root applied, R) on the (a) shoot length, (b) root length and (c) dry biomass of maize seedlings grown hydroponically in Hogland’s solution for 2 weeks. Data are mean of 9 replicates from 3 independent experiments with standard error bars. Bars labelled with different letters are significantly different (Duncan test; p<0.05).

**Effect of exogenously applied IAA and yucasin on exudation of IAA by *F. oxysporum* associated maize roots**

Highest amount of IAA was found in root exudates collected from endophyte associated maize seedling (10.5 ± 1.6 to 12.5 ± 1.8), no matter whether they received exogenous IAA or not. On exposure to yucasin (foliar or root), the seedlings ability to exude IAA was drastically reduced in comparison to the control. Interestingly, both foliar and root application of yucasin was inhibitory to the release of IAA by maize roots. In yucasin treated seedling, endophyte *F. oxysporum* restore the amount of IAA to the level of control seedlings (Table 1).

**Table 1.** Effect of IAA and yucasin (foliar, F and root applied, R) on the exudation of a) IAA by maize roots associated with endophyte *F. oxysporum*. Seedlings inoculated with isolated strain and in control were left for growth for 2 weeks. Absorbance was compared with standard curve to find out concentration of IAA and sugars in samples. Data are mean of 9 replicates ± standard error of mean. Similar bars labelled with different letters are significantly different (Duncan multiple range test; p<0.05).

<table>
<thead>
<tr>
<th>Treatments</th>
<th>IAA (µg/mL)</th>
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</thead>
<tbody>
<tr>
<td>Control</td>
<td>6.9 ± 0.8c</td>
</tr>
<tr>
<td>Yucasin (Foliar)</td>
<td>1.8 ± 0.0a</td>
</tr>
<tr>
<td>Yucasin (Root)</td>
<td>2.0 ± 0.0a</td>
</tr>
<tr>
<td>IAA</td>
<td>10.1 ± 1.6d</td>
</tr>
<tr>
<td>w1w</td>
<td>10.5 ± 1.6d</td>
</tr>
<tr>
<td>w1w+Y (F)</td>
<td>6.6 ± 0.7c</td>
</tr>
<tr>
<td>W1w+Y (R)</td>
<td>6.4 ± 0.4c</td>
</tr>
<tr>
<td>w1w+IAA</td>
<td>12.5 ± 1.8e</td>
</tr>
<tr>
<td>w1w+IAA+Y</td>
<td>11.3 ± 1.0e</td>
</tr>
</tbody>
</table>

**Effect of IAA, yucasin on maize root colonization by endophyte w1w**

Colonization frequency of w1w in maize root varied among different root zones (Figures 4 and 5). Root zone of division attracted the endophyte with highest efficiency showing a colonization frequency of 66.6%. However, colonization frequency dropped in the root above this zone and zone of elongation had a colonization frequency of 60.2%. Area of root in the proximity of stem i.e. zone of maturation was the least colonized showing a colonization frequency of 25% only. Application of IAA in combination with the endophyte had even more pronounced effect on root colonization by the endophyte enhancing colonization frequency to 75% in the zone of cell division, 57.6% in the zone of elongation and 25.5% in the zone of maturation. Application of yucassin (IAA inhibitor) negatively influenced the root colonization of w1w strain reducing its colonization frequency to 46% (foliar application) and 62% (root application) of the seedlings receiving no yucasin.
Figure 4. Effect of (a) yucasin (Y) and IAA on the mean percent colonization of maize roots by endophyte *F. oxysporum* wlw. The inhibitor were applied in the form of foliar spray (F) or in the culture media (R). Other treatments included exogenously applied IAA. The level of colonization was quantified by plotting the root segments on PDA after 14 days of co-culturing. Six root bricks for each plant, 2 from superior part (near to inoculum) two from middle part and two from inferior part (near to stem) were kept on clean filter paper and inoculated on PDA plates and incubated for one week. After a week colonization percentage was recorded according to the expression. Percent colonization of endophytic fungi=Number of positive segments/total number of root segments studied*100.

Figure 5. Visualization of wlw colonization (blue stained hyphae with cotton blue dye) in the maize root (a) control, (b) endophyte (c) endophyte+ yucasin.

Discussion

In present study an endophyte wlw was isolated from leaves of *Withania somnifera* previously and was identified as *Fusarium oxysporum* by comparative homology of ITS region near the 18 S rDNA gene [20]. The strain *Fusarium oxysporum* was also previously isolated as endophyte from *Juniperus recurva* [27]. The isolated fungus *F. oxysporum* possess plant growth promoting activities like ammonia and IAA production [20,28]. Fungal endophytes yield bioactive metabolites that promote the plant endophyte interaction [4]. Promotion of plant growth is the most important effect of fungal symbiosis [5]. Endophytic fungi promote plant growth by producing various secondary metabolites, including ammonia and plant hormones, particularly IAA [29,30]. Indole acetic acid as an essential compound for the growth and development of shoot and roots, many microorganism including plant growth promoting rizobacteria (PGPR) produce IAA [31]. Soil fungi and endophytes produce plant growth promoting substances IAA and gibberellin [7]. Culture filtrate of the endophyte *F. oxysporum* contained (31.6 µg/mL) of IAA and was greater than IAA recorded in culture filtrate of *P. glomerata* and *Penicillium sp* [16]. The endophyte *F. oxysporum* ability to produce IAA was tryptophan dependant demonstrating the presence of IPA (indole-3-pyruvic acid) pathway for the biosynthesis of IAA [32], described IPA, IAM (indole-3-acetamide), and TRA (tryptamine), pathways in *Fusarium delphinooides* [31], reported IAM, IPA pathways in *Colletotrichum gloeosporioides*. Tryptophan is considered as a precursor for IAA biosynthesis and its addition of in culture medium enhances IAA production [33,34]. According to our result increasing tryptophan concentration from 500 and 1000 µg/mL significantly enhanced amount of IAA by 17% and 25% greater than the control. Our results are in agreement with the work of Khalid et al. [35], who studied the effect of L-tryptophan concentration for the production of IAA and observed that L-tryptophan-derived auxin biosynthesis was enhanced several folds. Production of IAA has been observed in several endophytic fungi during the recent years [7,16,36]. In present study increasing tryptophan concentration lead to significant increase in production of IAA by *F. oxysporum*. Indole-3-acetic acid (IAA) with or without precursor L-tryptophan, is produced by most genus of *Trichoderma* [37]. Production of IAA is thought to be a unique character enabling endophytes to promote plant growth [38]. In present study *F. oxysporum* was able to colonize maize roots and significantly enhanced shoot and root length and seedling dry biomass (P>0.05). Increase in shoot and root
length in fungal inoculated plants had been reported previously by many authors [39,40]. Also previously Mehmood et al. [20] reported increase in shoot and root length and seedling dry biomass by culture filtrate of endophytic Fusarium oxysporum. Previously Machungo et al., [41] reported that endophytic F. oxysporum increase plant height 11% which was lower than increase in shoot length caused by our isolate. Enhanced plant growth through endophytes may be the result of phytohormones produced by fungal endophytes [42]. Application of IAA in the form of aerial spray on corn seedlings enhanced the ability of roots to attract endophytes and their subsequent colonization in the roots. IAA contents increased in root exudates sprayed with IAA and inoculated with F. oxysporum. Previously, it was shown that exogenous application of IAA restored in vitro growth as well as IAA biosynthesis was inhibited in plant by foliar application of yucasin [26]. Seedlings exposed to yucasin release significantly low amount of IAA confirming the inhibition of its biosynthesis. Interestingly, low IAA exuding maize roots were least colonized by the endophyte F. oxysporum pointing a role of this phytohormone in plant-endophyte interaction.

Conclusions

The endophytic fungus F. oxysporum wlv promoted the growth of maize seedlings under under hydroponic condition. Its ability to generate IAA is responsible for colonizing the roots of corn and establishing beneficial symbiotic association. Inhibition of IAA limits their ability to interact and carry out symbiotic associations.

References


