

Research Article

Assay of *NPR1* gene expression in wheat under powdery mildew stress

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Abstract: One of the effective plant disease management strategies is based on the employment of resistance inducers. In the present study, to assay, the effects of Salicylic acid (chemical inducer) and Piriformospora indica (biological inducer) on wheat powdery mildew (Blumeria graminnis f. sp. tritici), the expression rate of Non-expresser of pathogenesis-related genes1 (NPR1) gene was evaluated using qPCR. For this purpose, Falat and Tajan cultivars were selected as susceptible and resistant genotypes to powdery mildew, respectively. To evaluate the rate of gene expression, the P. indica colonized Falat along with mock plants were inoculated with Powdery mildew. In another experiment, Falat treated with SA and control plants were inoculated with Powdery mildew 48 h after treatment with SA. Gene expression was assayed in Falat compared with resistant cv. Tajan. Sampling was carried out at 0, 6, 12, 24 and 48 hours after infection. Comparisons of gene expression patterns showed that after infection, the expression levels of NPRI increased in induced and non-induced Falat and Tajan cultivars. The maximum gene expression levels were observed at 24 hours post infection. But the expression levels of the gene at this time were much higher in induced treatments compared with control. The current study showed that NPR1 can be involved in resistance strategy. Thus, using NPR1 gene as a desired gene in genetic engineering for increasing the potential of plant resistance to pathogens can be considered. Moreover, the high response of NPR1 gene in induced plants indicated that both SA and P. indica play a critical role in inducing resistance.

Keywords: Wheat, Resistance inducer, *Piriformospora indica*, Salicylic acid, *NPR1* gene

Introduction

Powdery mildew (PM) is an economically important disease in wheat (*Triticum aestivum*) caused by the obligate biotrophic fungus *Blumeria graminis* f. sp. *tritici* (*Bgt*). Under low

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infestation, the disease damage ranges from 13 to 34% but 50 to 100% loss could be recorded in a field when a severe infestation occurs (Li *et al.*, 2011; Alam *et al.*, 2013).

Plants have evolved the sophisticated immune system in challenge with microbial pathogens. Each plant genome encodes hundreds of so-called resistance (R) proteins that allow the plant to recognize specific pathogen-derived molecules known as avirulence (avr) factors (Dong, 2001). Among these resistance proteins,

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Non-expresser of pathogenesis-related genes1 (NPR1) is essential for transduction of the salicylic acid (SA) signal to activate Pathogen Related (PR) genes and induction of systemic acquired resistance (SAR), which confers longlasting broad-spectrum disease resistance in plants (Dong, 2001). The antagonistic effects of SA on Jasmonic acid (JA) signaling require also NPR1 expression (Spoel et al., 2007). Moreover. homologues of NPR1 have been cloned and characterized in several crop plants (Yuan et al., 2007; Zhao et al., 2009; Zhang et al., 2008). Several studies also indicated overexpression of AtNPR1 in Arabidopsis, and its expression in rice, tomato, and wheat caused enhancement in pathogen resistance via increasing PR genes expression (Fitzgerald et al... 2004; Makandar et al., 2006). It has been reported that silencing of NPR1 gene in rice exhibited enhanced susceptibility to pathogens and over-expression of this gene conferred disease resistance to bacterial blight in rice (Yuan et al., 2007). In addition, researches conducted by Stein et al. (2008) and Molitor et al. (2011) showed that NPR1 gene has an important role in resistance to Powdery mildew in Arabidopsis. Makandar et al. (2006) also confirmed that increasing NPR1 gene in wheat causes resistance to Fusarium graminearum.

Several reports indicated that SA and its analogues such as Benzothiadiazole (BTH) act as inducers for *NPR1* (Durrant and Dong, 2004). This leads to induction of both local and SAR in a wide range of plant species when challenged with invaders (Durrant and Dong, 2004). SAR is an inducible defense mechanism that confers resistance to a broad spectrum of pathogens (Durrant and Dong, 2004; Makandar *et al.*, 2006).

Moreover, some scholars have observed that root colonization of various plants by the endomycorrhizal fungus *Piriformospora indica* protected symbiotic plants, including cereals, against abiotic and biotic stresses (Deshmuckh *et al.*, 2006; Stein *et al.*, 2008). The *P. indica* confer resistance reminiscent induced systemic resistance (ISR) in plants (Stein *et al.*, 2008; Molitor *et al.*, 2011). ISR strategies based on *P. indica* could provide an approach to enhance

resistance against a broad range of pathogens without high metabolic cost (Molitor *et al.*, 2011).

In spite of several results with *NPR1* gene in diverse plants, there is little information on the function of *NPR1* homologue in wheat. Therefore, it was necessary for isolation, characterization, and identification of *NPR1* gene from wheat. In the present study *NPR1*gene, designated as *TaNPR1*, was isolated and its partial length characterized in wheat cultivar 'Falat', due to importance of *NPR1* gene in disease resistance induction. Then, *NPR1* gene expression profiles under treatments with inducers SA and endomycorrhizal fungus *P. indica* on the susceptible plant were also examined.

Materials and Methods

Experimental materials

Seeds of the two wheat cultivars, including Falat and Tajan were obtained from seed and plant improvement institute. These two wheat genotypes were selected as susceptible and resistant cultivars to powdery mildew, respectively. These genotypes were grown in a greenhouse at 20 °C/18 °C (day/night) with 60% relative humidity and a photoperiod of 16/8 h light/dark with 240 μmolm²s⁻¹ photon flux density in 3 replicates. The biotrophic causal agent of powdery mildew, *Blumeria graminis* f. sp. *tritici* (*Bgt*), race Karaj, was propagated in the same conditions on susceptible wheat cultivar 'Bolani'.

Salicylic acid treatment

Two weeks later, susceptible plants were treated with SA (3 mM) that was dissolved in 30% ethanol. Only 30% (v/v) ethanol with no SA was used in control (Muchembled *et al.*, 2006). All plants were inoculated with *Bgt* spores (50 conidia per mm²) at 48 hours post treatment and incubated in a growth chamber (Renard-Merlier *et al.*, 2009).

Piriformospora indica treatment

Roots of 3 days old seedlings of Falat were immersed into a suspension of *P. indica* spores

with 5×10^5 spore/ml for 12h. The colonized plants were then transplanted into the soil (perlite, clay and ceramic) and kept in growth chamber and irrigated weekly with Hoagland solution. To prove root colonization with *P. indica*, the roots of some plants were rinsed with tap water three weeks after inoculation and stained by vierheilig *et al.* (1998) method, then observed by light microscope. Colonized and control plants were inoculated with *Bgt* spores (50 conidia per mm²) and incubated in the same conditions.

Inoculating resistant and susceptible wheat cultivars with *Bgt*

Two-weeks old seedlings of resistance and susceptible cultivar were inoculated with Bgt spore (50 conidia per mm²) and incubated in a growth chamber.

RNA extraction and gene expression analysis

The first leaves of ten seedlings were harvested at 0, 6, 12, 24 and 48 hours post-infection (hpi) with Bet in treated and control plants and Tajan cv. Then, plant tissues (100 mg) were ground in liquid nitrogen. Total RNA was extracted by RNX-plus kit (Sinaclon, Cat, No: RN7713C). The extracted RNA was incubated with RNasefree DNase (Fermentas, Cat. No: EN0525). The first-strand cDNA of each sample was synthesized using Revert AidTM First Strand cDNA Synthesis kit (Fermentas, Germany) according to the manufacturer instructions. Quantitative Real-Time PCR (qRT-PCR) was performed in 15µl of SYBR Green Quantitative PCR Master Mix kit (Fermentas, Cat. No: K0221) in 96-well plates of C1000™ Thermal Cycler (BioRad). The synthesized cDNA was diluted 1:5 with nuclease- free water and 2 µl of the diluted cDNA (300 η g) was used as a template. The qRT-PCR program included 10 min at 95 °C, followed by 40 cycles of 95 °C for 15 s, and 60 °C for 30s. The expression was determined using delta-delta Ct method (Livak and Shmittgen 2001). For normalization, the Actin gene was applied as the housekeeping gene. Experiments were performed in 3 replicates and standard value and the results were analyzed by the student's T-test.

TaNPR1 gene isolation and bioinformatic assay

A pair of degenerated primers (*Tainpr1*) were designed on the basis of conserved nucleotide sequences of cereals NPR1 cDNA using Bio Edit 7.0.9.0 and Oligo Explorer V1.4 software (Table 1). PCR amplification program was as follows: 94 °C for 3min; 35 cycles of 94 °C for 30 s. 55 °C for 45 s. 72 °C for 1 min and 72 °C for 20 min. The PCR products of the predicted size (about 930 bp) were purified from the agarose gel and then sequenced by Bioneer Company. Then, on the basis of TaNPR1 gene, a pair of sense primer (TaNPR1) was designed for RT-PCR that amplified a fragment length of 157 bp (Table 1). Identification of purified nucleotide sequences was performed using BLAST program. Sequence alignment and comparison with sequences were conducted using ClustalW program from Bio Edit 7.0.9.0. The phylogenetic tree was generated from deduced nucleotide sequences for TaNPR1 and NPR1 homologues from other species using MEGA 5.0 software, maximum likelihood clustering method.

Results

Assessment of susceptible wheat colonized with *P. indica*

Microscopic studies on Falat cultivar's roots inoculated with *P. indica* showed the ability of this endomycorrhizal fungus in root colonization. Dense growth of *P. indica* hyphae were observed on the outer surface and in cortex sections of roots. Within the root cortex, fungal hyphae developed as a mass of spherical to pear shaped chlamydospores inside the root cells (Fig. 1). The microscopic studies indicated the appropriate establishment of *P. indica* in wheat roots.

In order to assay the role of *P. indica* and SA in the induction of resistance to *Bgt*, the rate of mildew colonies formed per unit area (2.5 cm²) on leaves of Falat cv. (control) were compared with those of Falat treated with SA and *P. indica*, 7 days post inoculation. The number of *Bgt* colonies was found to be 47%

and 42% less than control plants in *P. indica* and SA treated plants, respectively. Moreover, the size of established colonies on leaves was decreased in both treated plants compared to that in control plants.

Table 1 Nucleotide sequence of primers used for isolation and gene expression of *TaNPR1*.

Primer name	Forward primer (5'→3')	Reverse primer (5'→3')
Ta inpr1	GCA ACA AAT CTT	TCA TCR TCC ATG
	GCR TGA AAC TG	ATC TTG TC
Actin	GGA AAA GTG CAG	TAC AGT GTC TGG
	AGA GAC ACG	ATC GGT GGT
Ta NPR1	GCT TGT CAG GAT	GAA CAG TAT AAC
	GCT GCT	CTC TTG GGT TTC



Figure 1 Formation of *Piriformospora indica* chlamydospores inside colonized wheat root of Falat cultivar.

The rate of *NPR1* transcription in Falat colonized with *P. indica*

Inspection of gene expression pattern showed that in the absence of a pathogen challenge, expression of NPR1 gene was similar between P. indica colonized and non-colonized plants. While the NPR1 gene expression levels increased in both plants at early hours after infection by Bgt. NPR1 transcripts level elevated 1.5 fold more in 6 and 12 hpi in P. indica colonized plants compared to noncolonized plants. The maximum level of NPR1 gene expression was observed at 24 hpi in both non- and colonized plants (Fig. 2a). At this time, the expression rate of NPR1 in non- and colonized plants increased 5.1 and 8.9 folds, respectively. So that, it was revealed that NPR1 gene transcription levels in P. indica colonized plants was 1.7 fold higher than that of the non-colonized plants.

Rate of NPR1 transcriptome in Falat treated with SA

At 48 h after spraying plants with SA, the expression rate of NPR1 increased significantly in treated plants in comparison with control plants (Fig. 2b). In addition, at different hpi an increase in NPR1 gene expression level in both groups of plants was observed. However, in treated plants, NPR1 transcripts increased significantly by 1.7 fold over that of the control plants at 12 hai. The NPR1 expression reached its peak value in both groups at 24 hpi. Comparison of transcript levels at this time indicated that level of transcription in treated plants was about 2.1 fold higher than in control plants. Also, both groups of plants at 48hpi showed a reduced level of expression, but in treated plants, this rate was drawn lower than control plants.

The rate of *NPR1* transcriptomes in susceptible and resistant cv.

The gene expression increased 3.5 and 2.3 fold at 6 hpi, in cv. Tajan and Falat, respectively (Fig. 2c). Although, gene expression in Tajan increased about 1.5 fold more than Falat at 12 hpi. The *NPR1* expression at 24 hai in both cultivars demonstrated its maximum value. At this time, the rate of *TaNPR1* gene expression in Tajan was almost 1.7 fold higher than Falat. Also, Gene expression at 48 hpi in both groups decreased dramatically, but the expression level was higher in Tajan cultivar.

Isolation and sequence analysis of wheat *NPR1* gene

A partial-length cDNA of *TaNPR1* gene containing about 913 bp, which was registered in GenBank by accession No, KM017012 was used in this experiment. The *TaNPR1* is formed of 267 adenine (A), 184 cysteine (C), 212 guanine (G) and 250 thymine (T) nucleotides. This cDNA encoded a predicted polypeptide of 303 amino acids with an estimated molecular mass of 34.09 kDa and a theoretical pI of 7.8

and 100.7 aliphatic indexes. A BLAST search of *TaNPR1* in GenBank revealed that this gene from wheat shared the highest identity of 89% with wheat *TdNPR1* (JX424315), 88% with barely *HvNPR1* (AM050559) and 84% with Rice *OsNPR1* (DQ450947, AY923983).

Assaying the type and amount of amino acids in the TaNPR1 protein suggests that this protein contained all amino acids but tryptophan (Fig. 3). In the structure of this protein, leucine amino acid with over 40

molecules was the most abundant amine acid, while, tyrosine, cysteine, and glutamine with less than 7 molecules were the least abundant amino acids. Also, results of maximum likelihood phylogenetic tree indicated that TaNPR1 protein was grouped with the proteins encoded by Ta-NPR1, TdNPR1, HvNPR1 and OsNPR1. Furthermore, GhNPR1 and NtNPR1 were clustered together and AtNPR1, GmNPR1 and ZmNPR1 formed the third group (Fig. 4).

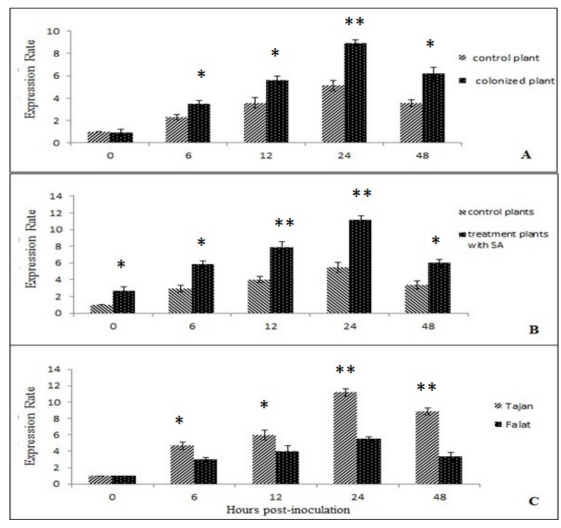


Figure 2 Expression levels of *NPR1* gene in *P. indica* colonized and control 'Falat' (A), treated with SA and control 'Falat' (B) and Resistant ('Tajan') and Susceptible ('Falat') plants challenged with *B. graminis*(C). * and ** = indicate significant difference at 0.05 and 0.01 levels, respectively.

1 NKSCMKLLERCLEMV V QSNLDMITLEKV LP PDV V KQITDLRLSFGLASSEDMGFPNKHV R
61 RILRALDPDD V ELV RMLLKE GQTNLDDAFA LHYAVEHCDSKITTELLDIA LADV SLRNPR
121 GYTV LHIAAR RKDPKIIVSL LTKGARPSDF TFDGRKAVHISKRLTKHGDY FANTGEGKPY
181 PNDKLCIEIL EQAERRDSHF GKAYV SLALD GDV LRGRLLY LENRVALSRI LFPVE ARV AT
241 DIAQV DGTSE FALGHLV AIDLNGTPTKMKDEHLARMRALSKTV ELGKRY FPRCSNV LDKI
301 MDD

Figure 3 Sequence of amino acids in an ORF +3 protein of wheat TaNPR1.

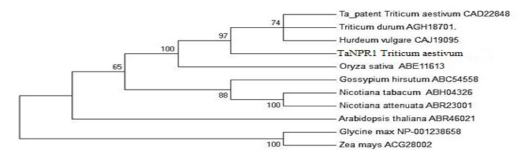


Figure 4 Maximum likelihood phylogenetic tree of NPR1 sequences from various plant species. Branches are labeled with plant species names followed by the protein data bank accession number. Bootstrap value is shown at every node (1000 replicates). Branches over 70 percent are confirmed.

Discussion

Powdery mildew would be one of the most damaging wheat diseases without the extensive use of conventional fungicides. Disease resistance strategies using the components of the JA and SA-signaling pathways have the potential to be effective and versatile. However, their success depends on managing between pathogen and plant defense mechanism. In this article, we have discussed the effect of Salicylic acid and *P. indica* which are classically described as inducers of disease resistance, in *NPR1* gene expression and reducers disease severity.

NPR1 is a novel protein with ankyrin repeats that is important in the induction of PR genes (such as PR1, PR2, and PR5) and eventually induced resistance to biotic stresses such as bacterial disease caused by Xanthomonas oryzae pv. oryzae and fungus Magnaporthe oryzae in rice (Takatsuji, 2014) and Fusarium

graminearum disease in wheat (Makandar et al., 2006) and abiotic stresses such as enhanced oxidative stress tolerance in transgenic tobacco (Srinivasan et al., 2009) and reduced tolerance to heat and salt in Mutant npr1 Arabidopsis (Larkindale et al., 2005). Partial-length of TaNPR1 gene isolated from Falat leaves contained a 913 bp. the BLAST search of TaNPR1 in GenBank revealed that this gene shared the highest identity with T. durum, H. vulgar and O. sativa. Also, results of maximum likelihood phylogenetic tree indicated a high level of conservation of TaNPR1 protein from different plant species, especially among the monocots such as wheat, barley, and Rice. To now many researchers have cloned homologues of NPR1 in several crop plants (Yuan et al., 2007; Zhao et al., 2009; Zhang et al., 2008).

Inspection of gene expression pattern showed a significant increase in *NPR1* gene expression in early hours after infection by *Bgt* in treated Falat and also in cv. Tajan compared

with control plant. Results of previous investigations indicated that P. indica and SA confer induced resistance (IR) in plants (Stein et al., 2008; Molitor et al., 2011). Meanwhile, based on the results, in absence of a pathogen (zero time), NPR1 gene were undistinguishable between non- and colonized plants by P. indica. So, the P. indica by skillful systemic effects caused priming plant to the rapid increase in defense gene upon infection occurred by Bgt. Results of previous investigations indicated that P. indica confers resistance reminiscent of ISR in plants (Molitor et al., 2011). Also, significantly increased NPR1 gene in plants treated with SA, indicated that SA has positive effects on NPR1 increasing, SAR induction, elevation of PR genes transcription and inducing disease resistance in plants. The contribution of SA and NPR1 on plant defense response were also reported (Yuan et al., 2007; Maier et al., 2011). Previous studies indicated that mutations in NPR1 and SA-responsive genes lead to suppress cell death and increase susceptibility to biotrophic pathogens (Rate et al., 1999; Fitzgerald et al., 2004).

The plant resistance mechanisms against pathogens, amount, and rate of defense genes expression play a central role. So, the SA and P. indica with higher and earlier induction of defense genes may cause an earlier induction of IR in treated plants during the penetration phase to prevent the growth of fungal hyphae in plant cells (Eichmann and Hückelhoven, 2008). So that, earlier and higher expression of NPR1 gene have been observed in treated plants at 6 to 12 hpi. This overexpression has occurred preceding the formation of haustorium at 12 to 24 hpi (Eichmann et al., 2006). These results were also reported by stein et al., (2008), Molitor et al. (2011) and Li et al., (2011) in P. indica colonized plants.

The expression of *NPR1* gene in Falat was delayed and significantly lower than that in Tajan after infection. So, it seemed that delaying expression of *NPR1* gene provides an opportunity for the fungus to spread on Falat. We obtained a high potential of plant defense mechanisms through *NPR1* gene in cv. Tajan.

Thus, if *NPR1* gene expression is increased at the right time, then it would induce resistance to pathogens that coincides with results by Sugano *et al.* (2010) and Feng *et al.* (2011).

In addition, the evaluation of gene expression indicated that, at 48 hpi, the transcription level of *NPR1* gene dampened, indicating an effective suppression of defense-associated genes upon haustorium establishment (Spanu *et al.*, 2010; Molitor *et al.*, 2011). However, reduction of expression level in non-treated plants was higher than that of the treated plants, So, the treated plants with increased expression levels try to prevent the growth of fungal secondary hyphae.

Previous studies represented that unlike SAR. ISR requires only cytosolic but not nuclear localization of NPR1 (Stein et al., 2008; Molitor et al., 2011). According to the significant increase in NPR1 gene expression in treated plants we believe that this gene if located in the nucleus and/ or cytosol, plays a positive role in the induction of resistance responses during biotic stress. The role of NPR1 gene in resistance to Fusarium graminearum in wheat (Makandar et al., 2006), and resistance to Powdery mildew in Arabidopsis (Stein et al., 2008; Molitor et al., 2011) has been proven. As a result, wheat NPR1 gene can be used as the desired gene for induced resistance to the pathogen, that high response of this gene to inducers and its high gene expression in Tajan cv. after infection is in agreement with this hypothesis. Thus, using NPR1 gene in genetic engineering for increasing the potential of plant resistance to pathogens can be considered. As well as, high expression of NPR1 gene in treated Falat indicated that SA-dependent pathway and P. indica play important roles in IR against Bgt. Therefore, application of these two inducers is proposed for inducing resistance processes.

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بررسى الگوى بيان ژن NPR1 گندم تحت بيماري سفيدك سطحي

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چکیده: یکی از استراتژیهای مدیریت بیماری براساس استفاده از القاکنندههای مقاومت می باشد. در مطالعه اخير، بهمنظور بررسي اثر سالسيليک اسيد (القاكننده شيميايي) و قارچ Piriformospora indica (القاكننده بيولوژيكي) در القاي مقاومت به بيماري سفيدك سطحي گنده .(Blumeria graminnis f. sp. ماري سفيدك سطحي tritici) بيان ژن (Non-expresser of pathogenesis-related genes1 (NPRI) با استفاده از تكنيك qPCR مورد بررسی قرار گرفت. در ابتدا دو رقم فلات و تجن بهترتیب بهعنوان حساس ترین و مقاوم ترین ژنوتیپ به سفیدک سطحی انتخاب شدند. سپس بهمنظور بررسی الگوی بیان ژن، فلات همزیست شده با P. indica به همراه گیاهان کنترل در معرض آلودگی با سفیدک سطحی قرار گرفتند. سپس در آزمایش جداگانهای فلات تیمار شده با اسید سالسیلیک بعد از ۴۸ ساعت بـهمـراه گیاهـان کنتـرل بـا عامل بیماری مایه کوبی شد. علاوه براین میزان بیان ژن NPRI در رقم فلات نیز در مقایسه با رقم تجن مور د بررسی قرار گرفت. سپس نمونهبرداری در بازههای زمانی ۰، ۶، ۱۲، ۲۴ و ۴۸ ساعت پس از آلودگی در تمامی تیمارها انجام گرفت. مقایسه الگوی بیان ژن نشان داد که پس از اعمال آلودگی سطح بیان ژن NPRI در فلات القا شده و کنترل و همچنین رقم تجن افزایش یافت. بیشترین سطح بیان ژن در ۲۴ ساعت پس از آلودگی مشاهده گردید. اما سطح بیان ژن در این بازه زمانی در گیاهان القا شده و رقم تجن بهطور معنی داری بیش تر از کنترل بود. نتایج این بررسی بیانگر این است که ژن NPR1 می تواند در استراتژی مقاومت دخیل باشد. لذا ژن NPRI را میتوان به عنوان ژن مطلوب در مهندسی ژنتیک برای افزایش پتانسیل القای مقاومت به پاتوژن درنظر گرفت. علاوه براین، پاسخ بالای NPRI در گیاهان تيمار شده نشان دهنده اين است كه دو القاكننده SA و P. indica نقش مهمي را بـراي القـاي مقاومـت ايفا مينمايند.

واژگان کلیدی: گندم، القای مقاومت، Piriformospora indica سالسیلیک اسید، ژن NPRI