# Alternative splicing and genetic diversity of the white collar-1 (wc-1) gene in cereal Phaeosphaeria pathogens 

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#### Abstract

The white collar-1 (wc-1) gene encodes an important light-responsive protein (wc-1) that maintains circadian clocks and controls numerous light-dependent reactions including sporulation in ascomycete fungi. The structure and expression of the $w c-1$ gene in wheatbiotype Phaeosphaeria nodorum (PN-w) was studied. It was shown that the full-size ( $3,353 \mathrm{bp}$ in length) wc-1 gene in PN -w contained 4 introns in which introns 1 and 2 were flanked by GC-AG splice borders and were spliced constitutively. However, introns 3 and 4 of the $w c-1$ gene were alternatively spliced. As the result of alternative splicing (AS), six transcript variants were


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identified, encoding different lengths of deduced polypeptides (from 1,044 to $1,065 \mathrm{aa}$ ). Ratios of the wc-1 gene transcript variants in the RNA population were the same in the sporulated and non-sporulated PN -w isolate Sn37-1 and the sporulated PN-w isolate S-79-1, grown under light/dark conditions. The AS of the wc-1 gene may control various light-dependent reactions in $\mathrm{PN}-\mathrm{w}$, which leads to diverse morphological, physiological and pathological characters for pathogen infection and spread. Based on the nucleotide and deduced amino acid sequences, the wc-1 gene in cereal Phaeosphaeria pathogens was diverse. It appeared that the deduced

[^0]wc-1 polypeptide sequences of $P$. avenaria f . sp. avenaria (Paa), P. avenaria f. sp. triticea (Pat1 and Pat3) and barley biotype $P$. nodorum (PN-b) were more closely related than PN-w and Phaeosphaeeria sp. (P-rye) from Poland. Based on the wc-1 deduced polypeptide sequences, $P$. avenaria f. sp. triticea (Pat2) from foxtail barley (Hordeum jubatum L.) was evolutionary well separated from the other cereal Phaeosphaeria pathogens.

Keywords Alternative splicing • Phaeosphaeria . White collar-1 • Sporulation • Phylogenetic relationship

## Introduction

The white collar-1 ( $w c-1$ ) gene encodes an important light-responsive white collar-1 protein (wc-1) in ascomycete fungi that controls numerous light-dependent reactions and maintains circadian clocks in the organisms (Lee et al. 2003). The best-studied fungal systems in terms of the molecular mechanism of light responses are Aspergillus nidulans and Neurospora crassa. In response to light, the wc-1 protein acts as a transcription factor to induce gene expression for numerous physiological processes including carotenoid synthesis, photo-adaptation, and spore formation (Ballario et al. 1996). The wc-1 protein forms a heterodimeric photoreceptor white collar complex (WCC) with white collar-2 (wc-2) protein via PAS (Per-Arnt-Sim) domains (Crosthwaite et al. 1997). In response to a light pulse, WCC is phosphorylated and an additional wc-1 molecule is synthesized and joins the WCC, which acts as the positive component for frequency (frq) gene induction (Cheng et al. 2003). In Aspergillus nidulans, immediately after the induction of wc-1 and frq proteins by light, sequential expressions of the genes including bristle (brlA), abacus (abaA), wetwhite conidia (wetA) and stunted (stuA) establish a linear regulatory pathway required for the transition from vegetative hyphae to conidiophores and conidia (Prade and Timberlake 1993). In genetic and biochemical studies, this group of genes in concert with others has been proven to be the central regulatory pathway to control specific gene expression during asexual spore formation in A. nidulans (Fortwendel et al. 2004).

Numerous protein-coding genes of eukaryotes have introns that are removed constitutively from their precursor messenger RNA (pre-mRNA) by
splicing. Another mechanism that generates multiple transcription variants from a single gene and miscellaneous proteins is alternative splicing (AS). It was reported that AS affected the expression of nearly $20 \%$ and $65 \%$ of protein-coding genes in a flowering plant (Arabidopsis thaliana) and human (Homo sapiens), respectively (Kim et al. 2006).

Regulatory mechanisms of AS for specific gene expression and protein function diversity have recently been reviewed (House and Lynch 2008). AS was commonly reported in mammalian cells and their corresponding viruses. The multiple protein isoforms produced by AS may differ in structure, function and localization (Ogura et al. 2001). The polymorphic proteins encoded by spliced mRNA transcript variants of the human tumor suppressor gene (ING4) had different targeting signals, which governed their subcellular localizations and maintained their stability (Tsai et al. 2008). AS of a viral RNA transcript might also provide various proteins with specialized functions, which affected their interactions with other proteins/ factors and were important for certain stages of virus developmental cycle (Devireddy and Jones 1998). AS was also recently reported in a plant gene (JAZ.10) encoding a jasmonate ZIM-domain protein (JAZ). JAZ are reported to act as repressors of jasmonate signaling in plants (Chini et al. 2007). The JAZ. 10 gene contains 4 introns (accession no. NC_003076), and AS of the JAZ. 10 gene pre-mRNA in Arabidopsis thaliana produced three protein isoforms, a regular repressor (JAZ10.3, accession no. NP_974775, 185 aa) and two variants with either a longer C terminus (JAZ10.1, accession no. Q93ZM9, 197 aa) or a shorter one lacking the Jas domain (JAZ10.4, 167 aa) (Chung and Howe 2009). For regulation of plant defence reaction and normal growth development, both variant proteins (JAZ10.1 and JAZ10.4) were antagonistic to JAZ10.3 protein, and became highly resistant to jasmonateinduced JAZ degradation and attenuated jasmonate signaling (Chung and Howe 2009).

Wheat-biotype Phaeosphaeria nodorum (PN-w) is one of the important cereal diseases, causing stagonospora nodorum leaf blotch. Spore dissemination during the wheat growing season is an important tool for disease spread and severity. Sporulation in most PN-w isolates, such as $\mathrm{Sn} 37-1$, was reportedly activated by the near-ultraviolet light on the culture media (Cooke and Jones 1970). However, a PN-w S-79-1 isolate in B.M. Cunfer's culture collection was unique
in that it would produce pycnidiospores in the dark on the culture media, and its sporulation was not affected by blue, green and red light spectra (Personal communication). Since fungal sporulation is probably correlated with the activation of light-responsive genes, we determined to study the expression of the wc-1 gene in sporulated and non-sporulated PN-w cultures. We report here for the first time that the $w c-1$ gene in $\mathrm{PN}-\mathrm{w}$ is an AS gene. We also show that the wc-l gene transcription variants were formed proportionally in the population of both sporulated and un-sporulated cultures grown on the culture media under different light conditions. Also, based on the deduced wc-1 polypeptide sequences, phylogenetic relationships among cereal Phaeosphaeria pathogens were investigated.

## Materials and methods

Fungal isolates and DNA isolation
Cereal Phaeosphaeria species were either collected from Canada, Poland and the USA or purchased from the American Type Culture Collection (ATCC, Manassas, VA). Procedures for maintaining and growing fungal cultures were described previously. The genomic DNA (gDNA) was purified from 9 wheat-biotype $P$. nodorum (PN-w), 5 barley-biotype $P$. nodorum (PN-b), 9 P. avenaria f. sp. avenaria (Paa), 5 homothallic $P$. avenaria f. sp. triticea (P. a. f. sp. t., Pat1), 2 heterothallic $P$. a. f. sp. $t$. from foxtail barley (Pat2), 1 $P$. a. f. sp. $t$. from the state of Washington (Pat3) and 2 Phaeosphaeria sp. from Polish rye (P-rye) by CsCl gradient ultracentrifugation (Table 1).

Gene amplification and sequencing of genomic DNA
The full-length wc-1 gene sequence in $\mathrm{PN}-\mathrm{w}$ isolate SN15 was obtained by a protein BLAST search of the Stagonospora nodorum isolate SN15 database (http: www.broad.mit.edu) using the white collar-1 (wc-1) polypeptide sequence from Neurospora crassa (accession no Q01371, 1,167aa) as the query (Ballario et al. 1996). The 3,353-bp nucleotide sequence (accession no CH445345, nt235038-238390) encoding a hypothetical protein (SNOG_12044; accession no EAT80456, $1,079 \mathrm{aa}$ ) was chosen. Six primer sets ( $8 \mathrm{~A} / 5 \mathrm{~B}, 1 \mathrm{~A} / 1 \mathrm{~B}$, $2 \mathrm{~A} / 2 \mathrm{~B}, 3 \mathrm{~A} / 3 \mathrm{~B}, 4 \mathrm{~A} / 4 \mathrm{~B}$ and $5 \mathrm{~A} / 8 \mathrm{~B}$ ) were designed from the $3,669-\mathrm{bp}$ wc-1 gene nucleotide sequence of PN -
w isolate SN15, which included both $5^{\prime}$ and $3^{\prime}$ flanking-end partial sequences, to amplify the corresponding gene in other $\mathrm{PN}-\mathrm{w}$ isolates and six other Phaeosphaeria pathogens (Table 2). Based on partial sequencing data in PN-b, Paa and Pat1, specific primers were designed to produce the PCR fragments to cover the gaps and complete sequence analysis of the genes (Table 2). PCR amplification was performed in a $50-\mu \mathrm{l}$ reaction mixture containing 1 x colorless GoTaq $^{\circledR}$ flexi buffer ( pH 8.5 ), $1.5 \mathrm{mM} \mathrm{MgCl} 2,0.2 \mathrm{mM}$ dNTPs, $1.5 \mu \mathrm{M}$ of each primer, 80 ng of genomic DNA and 1.0 unit of GoTaq ${ }^{\circledR}$ DNA polymerase (Promega, Madison, WI). Reaction parameters were: denaturing at $94^{\circ} \mathrm{C}$ for 3 min , amplifying by 40 cycles of $94^{\circ} \mathrm{C}$ for $20 \mathrm{~s}, 55^{\circ} \mathrm{C}$ for 30 s and $72^{\circ} \mathrm{C}$ for 1 min , and then extending at $72^{\circ} \mathrm{C}$ for 10 min . Methods for isolation and direct sequencing of PCR products were reported previously (Ueng et al. 2003).

Gene expression in wheat-biotype $P$. nodorum isolates
To determine the wc-l gene expression in two PNw isolates, Sn37-1 and S-79-1, total RNA was isolated from these cultures grown in the dark and under continuous fluorescent lights at room temperature $\left(20 \pm 1^{\circ} \mathrm{C}\right)$ on sterile nitrocellulose membranes (BA-S 85, Schleicher \& Schuell Inc., Keene, NH), which were layered on V-8 juice agar ( $18 \%$ V8 juice, $0.2 \%$ calcium carbonate, and $2 \%$ agar). Extraction of total RNA mainly followed the protocols described previously (Wang et al. 2007). The mycelia were harvested and flash-frozen in liquid nitrogen. Total RNA was extracted from mortar/pestle pulverized mycelia using the RNeasy Plant Mini Kit according to the manufacturer's instructions (Qiagen Inc., Valencia, CA) and treated with RNase-Free DNase I enzyme (Roche Diagnostics Corporation, Indianapolis, IN). Lack of residual gDNA in total RNA was evidenced by not being able to amplify the histidinol dehydrogenase (Hdh1) gene fragment using the primer set 15A/12-1 (ATGCCGGCAGGACCCAGTGA/CTATCAAG CTACGCCAAGTCGC) (Wang et al. 2007). The firststrand (1x) cDNA was synthesized with random primer $\mathrm{p}(\mathrm{dN})_{6}$ and the First Strand cDNA Synthesis Kit (Roche Diagnostics Corporation, Indianapolis, IN). The thermal cycler settings were $25^{\circ} \mathrm{C}$ for 10 min , $42^{\circ} \mathrm{C}$ for $60 \mathrm{~min}, 99^{\circ} \mathrm{C}$ for 5 min , and $4^{\circ} \mathrm{C}$ for 5 min .

Six primer sets for gDNA amplification were also used for PCR production with 1 x cDNA as templates. To

Table 1 Isolates of cereal Phaeosphaeria species used for analysis of the white collar-1 (wc-1) gene

| Species | Original host |  | Year | Geographic location |
| :--- | :--- | :--- | :--- | :--- |
|  |  |  |  | GenBank |
|  |  |  |  | accession |
| number |  |  |  |  |

Table 2 Primer sets used for genomic DNA and 1x cDNA amplifications and sequencings of the white collar-1 (wc-1) gene in cereal Phaeosphaeria pathogens

| Primer sequences ( $5^{\prime} \rightarrow 3^{\prime}$ ) | Nucleotide positions (nt) in the wc-l gene (SNOG_12044) of PN-w isolate SN15 | Species used ${ }^{\text {a }}$ |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | PN-w | P-rye | PN-b | Paa | Pat1 | Pat2 | Pat3 |
| $\mathbf{8 A} / \mathbf{5 B}{ }^{\text {b }}$ (CTCTGCAATGCTCCCAAACACAG/CTG CTCTCTGGCGACATTTC) | -115--93/293-274 | X | X | X | X | X | X | X |
| 1A/1B (ATGAATGGAAACGGCTATCC/ CTTGTGCGACTACCCAATCCAG) | 1-20/836-815 | X | X | X | X |  | X | X |
| VD1-1A/VD1-1B (ATGAATGGAAACGGCTATCC CTAC/CTACAGGTTGATGTGCTTTGAGAG) | 1-24/958-935 |  |  |  |  | X |  |  |
| $\mathbf{2 A}$ ² $\mathbf{2 B}$ (CTCGAGTCTCCTGTAACGCAAG/GTTG GTAGTTGACTCTGTACG) | 679-700/1652-1632 | X | X |  | X | X | X | X |
| VD1-2A/VD1-2BB (GTCTCTGGATTGGGTAGTC GCAC/GATTTCTGAGAATCGTTGAACAC) | 811-823/1698-1676 |  |  | X |  |  |  |  |
| 3A/3B (CATGAACTTGCTTACAATGATAC/GTGG TAAATGCCTGCAGCACC) | 1458-1480/2418-2398 | X | X | X |  |  | X | X |
| 3AA/3BA (CATGAACTTGCTGACAATGATAC/GT GGTAAAGGCCTGCAGCACC) | 1458-1480/2418-2398 |  |  |  | X |  |  |  |
| VD1-3AA/VD1-3BA (CAGGTGCTATGACAAACA AGAAC/GGGTCTGCCCAACAATGAAGGTGG) | 1550-1572/2477-2454 |  |  |  |  | X |  |  |
| 4A ${ }^{\mathrm{d}} / 4 \mathrm{~B}$ (GACAAGAATCGAAGGTCCAATTC/TCA TGAACCTGGCTCCGCCACTTC) | 2301-2323/3353-3330 | X | X | X | X |  | X | X |
| VD1-4A/4B (GCATTCTTGAACTTGCAAGATCTG/ TCATGAACCTGGCTCCGCCACTTC) | 2328-2351/3353-3330 |  |  |  |  | X |  |  |
| $\mathbf{5 A}{ }^{\mathrm{e}} \mathbf{8 8}$ (CAGCACCTACAGCAAGGTCG/CAGTCA CCTCTCCAAATGTAAC) | 3066-3085/3554-3533 | X | X | X | X | X | X | X |

${ }^{\text {a }}$ PN-w $=$ Wheat-biotype $P$. nodorum; P-rye $=$ Phaeosphaeria spp. from Poland; $\mathrm{PN}-\mathrm{b}=$ Barley-biotype $P$. nodorum; $\mathrm{Paa}=P$. avenaria f. sp. avenaria; Pat1, Pat2 and Pat3 $=P$. avenaria f. sp. triticea. ${ }^{\text {b }}$ Primer 5B substitution for Pat $2=5 \mathrm{BA}$ (CTG CTT TCT GGC GAC ATT TC); $\mathrm{PN}-\mathrm{b}=5 \mathrm{BB}(\mathrm{CTGCTCTCTGGAGACATTTC}) .{ }^{\mathrm{c}}$ Primer 2A substitution for Paa and Pat $1=2 \mathrm{AA}$ (CTCGAATCTCCTGT GACACAAG). ${ }^{\text {P Primer } 4 \mathrm{~A} \text { substitution for } \mathrm{Paa}=4 \mathrm{AA}(\mathrm{GACAAGAATCGAAAATCCAATTC}) .}{ }^{\mathrm{e}}$ Primer 5A substitution for Pat $1=$ 5AA (GTTAGCACCTACAGCAAGGTCGTG); Pat2 $=5 \mathrm{AC}(\mathrm{CAGCACTTACAGCAAGGTCG}) ;$ Pat $3=5 \mathrm{AB}$ (CAGCACCTACAG CAAGGCCG)
confirm the $3^{\prime}$ end sequence in 1 x cDNA, three primer sets, 10A/13B (GACAAGAGCAAGAAGAGTCCA TC/GTTGGATCGTGGTTCGTGAG, nt3117-3139/ nt3737-3718), 6A/4B (GTGTTTGACGAACTGAA GACTAC/TCATGAACCTGGCTCCGCCACTTC, nt2756-2778/nt3353-3330) and 7A/4B (GACCATCTG GTAACAGAGACTTG/TCATGAACCTGGCTCCGC CACTTC, nt2958-2980/nt3353-3330) were further used for amplification. Except for the PCR fragments amplified with primer sets $10 \mathrm{~A} / 13 \mathrm{~B}, 6 \mathrm{~A} / 4 \mathrm{~B}$ and $7 \mathrm{~A} / 4 \mathrm{~B}$, the amplified products were isolated and directly sequenced as reported previously (Ueng et al. 2003).

Analysis of the deduced amino acid sequence
Based on the deduced polypeptide sequences of the $w c-1$ gene, phylogenetic relationships among 8 cereal Phaeosphaeria pathogens were analyzed using Mega

Version 4.0 (http://www.megasoftware.net/index. $\mathrm{html})$. The blue-light regulator 1 protein sequence of Bipolaris oryzae (BAF35570) was used as the outgroup in analysis. The amino acid sequences were aligned with complete deletion option, which eliminated the positions containing gaps. The bootstrap consensus tree was inferred from 10,000 replicates with the neighbor-joining method to represent the evolutionary history (Saitou and Nei 1987).

## Results

Expression of the wc-1 gene in wheat-biotype P. nodorum (PN-w)

Five exons and four introns (nt127-177, nt15771625, nt3014-3068 and nt3223-3273) were deter-
mined in the full-length ( $3,353 \mathrm{bp}$ ) wc-1 gene of PNw isolate Sn37-1 (Fig. 1). In PN-w isolates, it was unique that introns 1 and 2 were flanked by a GC-AG splice border, instead of GT-AG (Table 3). Similar splice borders were found in the first two introns of the $w c-1$ gene in P-rye, which is closely related to PN-w. In the other five cereal Phaeosphaeria pathogens, only intron 1 of the $w c-1$ gene contained the GC-AG splice border (Table 3).

AS was found for introns 3 and 4 of the $w c-1$ gene in PN -w (Fig. 2). With either the 6A/4B or the 7A/4B primer set, there were two smaller PCR products amplified from the 1 x cDNA than the product from the gDNA (Fig. 1). When the 10A/13B primer set was used, two PCR products were amplified when the 1x cDNA was used as template. However, size of one of these PCR products was the same as the amplified product from the gDNA. The PCR products amplified with primer set 7A/ 4B, which included both introns 3 and 4, were purified
with the PCR advanced ${ }^{\mathrm{TM}}$ PCR clean up mini prep system (Viogene BioTek, Taipei, Taiwan), cloned with the pGEM ${ }^{\circledR}$-T Easy vector system (Promega, Madison, WI) and sequenced. The size of spliced nucleotides in intron 3 was either 55 or 64 bp . For intron 4, the region was either retained in the RNA transcript or spliced. The size of spliced nucleotides in intron 4 was either 51 or 54 bp . Through AS at the post-transcriptional level, six $w c-1$ gene transcription variants (1-6) were identified in $\mathrm{PN}-\mathrm{w}$ (Fig. 2A).

Duncan's multiple range test was used to study the significance of difference among transcript variants and the isolates (SAS Version 9.1, 2003).Variance analysis showed that a significant difference of the clone numbers was observed in the randomly cloned population among six transcript variants, but not among two PN-w isolates, Sn37-1 and S-79-1, grown under different light conditions (Table 5). The analysis showed that the means of the number of clones were not


Fig. 1 Amplification of the white collar-1 (wc-1) gene in wheat biotype Phaeosphaeria nodorum isolate Sn37-1. Five translated exons of the wcl gene are shown as black boxes, and the noncoding region at the $3^{\prime}$ end as an open box. Four introns are indicated as the thin lines. The alternative splicing of the introns 3 and 4 are marked with asterisks "*". The wcl gene was amplified with 6 primer sets $(1 \mathrm{~A} / 1 \mathrm{~B}, 2 \mathrm{~A} / 2 \mathrm{~B}, 3 \mathrm{~A} / 3 \mathrm{~B}, 6 \mathrm{~A} / 4 \mathrm{~B}$,

7A/4B and 10A/13B). The numbers below the grey boxes are the fragment sizes in base pairs (bp) produced from genomic DNA (gDNA); the numbers in parentheses the fragment sizes from 1 x cDNA prepared from the fungal culture grown under fluorescent light. $\mathrm{a}=$ Bands produced from genomic DNA; $\mathrm{b}=$ bands produced from cDNA. $\mathrm{M}=$ Molecular markers of 50 and 100 bp ladders

Table 3 Nucleotide sequences of $5^{\prime}$ and $3^{\prime}$ end exon-intron boundaries in 4 introns of the white collar-1 (wc-1) genes in cereal Phaeosphaeria pathogens ${ }^{\text {a }}$

| Species ${ }^{\text {b }}$ | Intron 1 | Intron 2 | Intron 3 | Intron 4 |
| :---: | :---: | :---: | :---: | :---: |
|  | 5' end 3' end | 5' end 3' end | 5' end 3' end | 5' end 3' end |
| PN-w | CAGIgcaagt cagIA | CAGIgcaagt taglA | CAGlgtatgt cagıICACCTACAG $\\| \underline{\mathbf{C}^{\text {c }}}$ | CGAIgtaaga aagı $\underline{C A G} \\| \underline{\mathbf{C}^{\text {c }}}$ |
| P-rye | CAGIgcaagt cagIA | CAGIgcaagt taglA | CAG\|gtatgt cag|CACCTACAG $\\| \underline{\text { C }}$ | CGAlgtaaga aagICAG $\\| \underline{\text { C }}$ |
| PN-b | CAGIgcaagt cagıA | CTGIgttagt tag A | CAG\|gtatgt cag|CACCTACAG $\\| \underline{\text { C }}$ | CAAIgtaaga aagıICAG $\\| \underline{\text { C }}$ |
| Paa | CAGIgcaagt cagıA | CTGlgtaagt tagı A | CAG\|gtatgt cag|CACCTACAG $\\|$ C | CGAigtaaga aagICAG $\\|$ C |
| Pat1 | CAGIgcaagt cagıA | CGGlgtaagt tag ${ }^{\text {A }}$ | CAG\|gtacgt tag|CACCTACAG $\\|$ C | CAAIgtaaga aagıLAG $\\|$ C |
| Pat2 | CAGIgcaagt cagıA | CAGıgtgagt tagıA | CAGIgtgcgt cagıCACCTACAG $\\|$ C | CGAlgtaaga aagıCAG $\\|$ C |
| Pat3 | CAGIgcaagt cagIA | CTGIgttagt tag $\mid$ A | CAG\|gtatgt cag|CACCTACAG\| C | CAAlgtaaga aagı $\underline{\text { CAG }} \\| \underline{\text { C }}$ |

${ }^{\text {a }}$ Lowercase letters indicate the intron nucleotides spliced, and uppercase letters were the adjacent nucleotides in the exons. "l" represented the splice cut points of the introns, in which the surrounding nucleotide positions were always -1 and +1 . Exonic nucleotide positions in $5^{\prime}$ donor sites were negative numbers in order ( -$3,-2$ and -1 ), and those in $3^{\prime}$ acceptor sites were positive ( +1 ). The $\underline{C A G} \| \underline{C}$ in introns 3 and 4 were the alternative $3^{\prime}$ end acceptor splicing sites. ${ }^{\text {b }}$ Cereal Phaeosphaeria pathogens included wheat-biotype $P$. nodorum ( $\mathrm{PN}-\mathrm{w}$ ), Phaeosphaeria sp. from rye (P-rye), barley-biotype $P$. nodorum ( PN -b), $P$. avenaria f. sp. avenaria (Paa), and $P$. avenaria f. sp. triticea (Pat1, Pat2 and Pat3). ${ }^{c}$ Shaded nucleotides in introns 3 and 4 had low frequencies in the corresponding nucleotide positions in the 5' and 3' ends exon-intron boundaries of the mammalian genes.
statistically different among transcript variants 1, 2, 3, 5 and 6 , but variants 1,2 and 3 were significantly more common than transcript variant 4 based on the Duncan test at $5 \%$ significance level (Table 5).

Six lengths of deduced protein isoforms (from 1,044 to $1,065 \mathrm{aa}$ ) could be obtained through the regulation of AS in the wc-1 gene of the PNw pathogen (accession no GU322809-GU322813 and GQ149725) (Fig. 2B). Based on the polypeptide sequence deduced from the protein isoform 1 (accession no GU322809), deletions and substitutions of amino acids occurred between aa971 and 1042 .

Structure of the deduced wc-1 polypeptide in wheat-biotype $P$. nodorum ( $\mathrm{PN}-\mathrm{w}$ )

SMART sequence analysis (http://smart.embl-heidelberg. de) revealed that the deduced wc-1 protein isoform 3 in PN-w contained 3 Per-Arnt-Sim (PAS) motifs (PAS $1=$ aa370-499, PAS A $=$ aa565-678 and PAS B $=$ aa679 750 ) and a single putative GATA-type zinc finger (Znf) domain (aa933-985) (Fig. 3). When compared to the PAS motif in the wc-2 protein of PN-w isolate SN15 (accession no. EAT78433), the identity of the PAS 1, A and B motifs of wc-1 protein in PN-w was $19 \%, 33 \%$ and $18 \%$, respectively. The PAS 1 also contained a conserved 8 amino acids light-oxygen-voltage (LOV) domain (GRNCRFLQ, aa415-422), which included a reactive cysteine (C) that formed a covalent bond with the $\mathrm{C}(4 \mathrm{a})$ carbon of a flavin on photoexcitation (Ballario et al. 1998). The Znf domain, which recognizes and binds to DNA consensus sequences (A/T)GATA(A/G) of other genes for transcriptional activation, was well
conserved and belonged to zinc finger type IVb ( $\mathrm{C}-x_{2}-$ $\mathrm{C}-\mathrm{x}_{18}-\mathrm{C}-x_{2}-\mathrm{C}$ ) (Teakle and Gilmartin 1998). Like the wc-1 protein in Neurospora crassa, a putative nuclear targeting sequence (KKKRK) upstream (aa923-927) and 2 potentially phosphorylated serine ( S ) residues (aa993 and 995) downstream of the Znf domain were identified (Fig. 3).

Diversity of the $w c-1$ genes and their deduced polypeptides in cereal Phaeosphaeria pathogens

The wc-1 gene coding sequences amplified from cereal Phaeosphaeria pathogens differed in nucleotide length, from 3,334 bp in Paa to $3,361 \mathrm{bp}$ in Pat2 (Table 4). Due to the intron size discrepancy, the protein isoform 3 deduced from transcription variant 3 were from 1,043 to 1,050 aa (Table 4). There were 2-57 amino acid substitutions in wc-1 protein isoform 3 of the examined 6 Phaeosphaeria pathogens as compared with PN-w isolate Sn37-1 (Table 4). Based on the wc-1 protein isoform 1 sequence in $\mathrm{PN}-w$, most amino acid divergences were present in the N terminus (aa1-330), between PAS B and Znf domains (aa800-900) and the carboxy terminus (after aa990) among cereal Phaeosphaeria pathogens (Fig. 3).

Phylogenetic relationships based on the deduced wc-1 protein isoform 3 sequence showed that Pat1 isolates were more closely related to Paa and other Phaeosphaeria pathogens than PN-w and P-rye (Fig. 4). All Phaeosphaeria pathogens appeared to form a single clade, with the exception of Pat2, which is from foxtail barley (Hordeum jubatum L.).


Fig. 2 Alternative splicings in introns 3 and 4 of the white collar-1 (wc-1) gene and their deduced polypeptide sequences in wheat-biotype Phaeosphaeria nodorum isolate Sn37-1. With the primer set 7A/4B (GACCATCTGGTAACAGAGACTTG/ TCATGAACCTGGCTCCGCCACTTC) and 1 x cDNA as the template, partial nucleotide sequences (nt2958-3353, see " $\downarrow$ ") of the amplified products represented 6 transcript variants (A); their deduced polypeptide isoforms (B) (from 1 to 6 ) are

## Discussion

The automated annotations of fungal genome sequences sometimes incorrectly identify intron boundaries in the deduced genes. Two introns, at nt1577-1625 and nt3014-3068, were hypothesized in the wc-1 gene of PN-w isolate SN15 by the Broad Institute (SNOG_12044). When the wc-l genomic sequence was analyzed by Softberry (http://linux1.softberry. com/berry.phtml) with Aspergillus as the organism parameter, the same two introns but with a 9 -bp longer splicing sequence in the second intron (nt3014-3077) were reported. Therefore, the deduced WC-1 polypeptide ( 1,076 aa) analyzed by Softberry was three amino acids fewer as compared to the analysis from the Broad
presented. In A, the coding sequences are dark boxed, and the spliced introns are indicated by thin lines with their nucleotide positions (nt) following the genomic sequence of the $w c-1$ gene (GQ149725). The GenBank accession numbers are in parentheses. In B, the splicing positions of 6 transcript variants are indicated in their polypeptide isoform sequences as "|". Amino acids involved in deletions (....) and substitutions (boxed) are in bold

Institute. These two introns predicted in the wc-1 gene by both annotations corresponded to introns 2 and 3 when the RNA transcripts were determined by sequencing in this study.

Two AS patterns were found in the wc-l gene of PN-w (Fig. 2). One was the retention and excision of the intron from the pre-mRNA, and the other was the alternative $3^{\prime}$ splice sites that shortened the downstream exon (Black 2003). Since the intron sizes in the $w c-1$ gene were relatively small (ca. 50 bp ) as compared to the exons, regulation of AS was suggested to be an intron definition (Romfo et al. 2000).

Consensus nucleotide sequences positioned at intron/exon borders were essential for precise pre-

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| 76 | MSPESRRLSMLNFGDPGDAGDFQFDMHAAGMNDMMRNNPSFPRASGDMSQHNRNSANNMGLNTQFQNQNQNPPYS | 150 |
| 151 | TMAAPGSAYASPLHPSAPLDLDMSPYPNGMNMSMDLDDSLNMMSGDMNMFSNNQFNTPMLESPVTQEFIGPMPAP | 225 |
| 226 | NQDNSMSAIPQGHFKRPSLSNTPETRSGVSGLGSRTSSQDQNSAPSQSRPQSEQRSSSKNNLPTQMSLSSLKAHQ | 300 |
|  | PAS 1 |  |
| 301 | PVALDPAQDLPEEKMNQLKDYRTAWKPPAGGFPSTMHSNPHQKTQFKDAYSSTGFDMLGVLMRVATRPD [PEIDIG | 375 |
| 376 | SVDLSCAFVVCDAEMDDIPIVYCSENFERLTGYTRHMILGRNCRFLQSPDGKVEAGIKRKYVDDDSVLYLKNMIN | 450 |
|  | (2) |  |
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| 526 | VFNDSQKPQSEQGQTISKDEVSNVLATYTGSGDSEITRR [IWDKVLLENTDDVVHVLSLKGLFLYLSPSSSRILEY | 600 |
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|  | PAS B |  |
| 676 | PVY] [TLSKTIVRESGGVGDNELWTKMSTSGMFLFVSSNVRQLLDKQPEDLVGTSIQALMRQESKVQFGRILELARS] | 750 |
| 751 | GRKGEVKHEMINKRGQVLQAFTTIYPGDATEGQKPTFIVGQTRLLKYSRNSANHRPSMYTNKERLGGGSTHSIPT | 825 |
| 826 | QNTLNSPYPGSVQDHTGSNTPITSSTDGRFVTTETNATTHAGHNGLQLGHQDQSLASDDNVFDELKTTRSTSWQY | 900 |</table-markdown></div> <br> 901 ELRQMEKRNRYLAEEVQTLLAAKKKRKRRKGA[GQMQKDCANCHTRTTPEWRRGPSGNRDLCNSCGLRWAKQ|HLQ 974 

(4)

975 QGRVSPRTSSA]ASDKSKKSPSPRHHMTLPNSVPETGQSAAIPSAMAPAT|ARVEAAVSQRFIPPPKIEEVAEPGS 1048

Fig. 3 Deduced polypeptide isoform 3 sequence of the white collar-1 (wc-1) gene in wheat-biotype Phaeosphaeria nodorum isolate $\operatorname{Sn} 37-1$. The numbers $1-1,048$ on both side columns represent amino acid sequence. The polypeptide isoform 3 sequence was deduced from transcript variant 3. PAS motifs (PAS $1=\mathrm{aa} 370-499$, PAS A $=\mathrm{a} 565-678$ and PAS B $=\mathrm{a} a 679-$ 750) and 1 GATA-type zinc finger (Znf) domain (aa933-985) are in bold and bracketed. The conserved amino acids in PAS 1
mRNA splicing. On the other hand, AS was a consequence of $5^{\prime} / 3^{\prime}$ splice sites with weak binding potential for spliceosome components. In GT-AGtype introns, they were $(\mathrm{C} / \mathrm{A})$ AGIGT $(\mathrm{a} / \mathrm{g})$ agt and $(\mathrm{t} / \mathrm{c})_{\mathrm{n}} \mathrm{n}$ $\left({ }^{c} / \mathrm{t}\right)$ AGIG, where " I " indicated the cut positions, at $5^{\prime}$ donor and $3^{\prime}$ acceptor sites, respectively (Mount 1982). In a study of known gene splice sites from five eukaryote species (Homo sapiens, Mus musculus, Drosophila melanogaster, Caenorhabditis elegans and Arabidopsis thaliana) and ascomycetes, more than $98 \%$ of them belonged to the canonical GT-AGtype, and only $0.74-1.0 \%$ of introns had non-
and Znf domains in Phaeosphaeria and other ascomycetes are dark boxed. The 8 amino acids (aa415-422) in the gray area of the PAS 1 domain are the conserved residues of the light-oxygen-voltage (LOV) domain. A putative nuclear targeting sequence (KKKRK) is double underlined (aa923-927), and two potential phosphorylation sites at serine (S) (aa993 and aa995) are single underlined
canonical GC-AG splicing sites (Kupfer et al. 2004). Non-canonical GC-AG-type introns could be spliced constitutively if nucleotides in other positions around the $5^{\prime}$ and $3^{\prime}$ exon-intron borders were as well conserved as those in high-strength GT-AG-type introns (Aebi et al. 1987). It was shown that a GT-AG-type intron with a point mutation at the +2 position of the 5 ' donor site from " t " $\rightarrow$ " c " would retain some 5 'splice signal and produce less of the normal mRNA (Iida 1990). It appeared that the presence of consensus nucleotide sequences CAGIgcaagt, except for the "c" at the +2 position, in the 5 ' donor site of intron 1 and

Table 4 Structure of the white collar-1 (wc-1) gene in cereal Phaeosphaeria species ${ }^{\text {a }}$

| Species/ <br> Isolates | Gene size $(b p)^{\mathrm{b}}$ | \# of nucleotide substitutions ${ }^{\text {c }}$ | Transcription variant 3 size (bp) ${ }^{\text {b }}$ | Deduced protein isoform 3 size (aa) ${ }^{\text {b }}$ | \# of amino acid substitutions ${ }^{\text {c }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Phaeosphaeria nodorum (wheat-biotype) (PN-w) |  |  |  |  |  |
| Sn37-1 | 3353 | - | 3147 | 1048 | - |
| Phaeosphaeria sp. (from Poland) (P-rye) |  |  |  |  |  |
| Sn23-1 | 3353 | 13 | 3147 | 1048 | 2 |
| Phaeosphaeria nodorum (barley-biotype) (PN-b) |  |  |  |  |  |
| S-80-603 | 3338 | 278 | 3132 | 1043 | 37 |
| Phaeosphaeria avenaria f. sp. avenaria (Paa) |  |  |  |  |  |
| ATCC12277 | 3334 | 276 | 3132 | 1043 | 37 |
| Phaeosphaeria avenaria f. sp. triticea (Pat1) |  |  |  |  |  |
| Sat24-1 | 3342 | 297 | 3135 | 1044 | 48 |
| Phaeosphaeria avenaria f. sp. triticea (Pat2) |  |  |  |  |  |
| ATCC26370 | 3361 | 344 | 3153 | 1050 | 57 |
| Phaeosphaeria avenaria f. sp. triticea (Pat3) |  |  |  |  |  |
| S-81-W10 | 3340 | 272 | 3132 | 1043 | 42 |

${ }^{\text {a }}$ Based on the hypothetical protein gene SNOG_12044 in the Stagonospora nodorum data base of the Broad Institute (www.broad. mit.edu/annotation/genome). ${ }^{\text {b }} \mathrm{bp}=$ base pairs; aa $=$ amino acids. ${ }^{\text {c }}$ Substitutions as compared with PN-w isolate Sn37-1
part of intron 2 in the wc-1 gene of cereal Phaeosphaeria pathogens would ensure their constitutive splicing in pre-mRNA (Table 3).

The presence of weakly conserved nucleotides surrounding the intron splice sites would allow AS to occur. A nucleotide " a ", instead of " t " found at the +6 position of the 5 ' donor site "gtaagt" of intron 4 in the $w c-1$ gene might not affect splicing efficiency since a mutation from " t " $\rightarrow$ " c " at this position in a human gene reported almost normal splicing (Table 3, Iida 1990). However, a low frequency of nucleotide "A" at the -1 position of the $5^{\prime}$ donor site in intron 4
of the $w c-1$ gene as compared to mammalian genes might make the splicing activity vulnerable (Table 3). Mutation at the -1 position of the $5^{\prime}$ donor site from "G" $\rightarrow$ "C" affected the splicing of human $\beta$-globin pre-mRNA (Vidaud et al. 1989). It was possible that relaxation of the $5^{\prime}$ splice site recognition gave ca. $50 \%$ of intron 4 retention and excision in the wc-1 gene of Phaeosphaeria pathogens (Table 3).

In comparison to the conservation in nucleotides surrounding the $3^{\prime}$ acceptor site in mammalian genes (Lopez 1998), a low frequency of "C" at the +1 position of the $3^{\prime}$ acceptor site in introns 3 and 4 of the


Fig. 4 Phylogenetic relationships based on the deduced polypeptide isoform 3 sequences of the white collar-1 (wc-1) gene in cereal Phaeosphaeria pathogens. The polypeptide sequences deduced from the $w c-1$ transcript variant 3 of Phaeosphaeria pathogens were aligned and analyzed. The
blue-light regulator 1 protein sequence of Bipolaris oryzae (BAF35570) was used as the out-group in analysis. GenBank accession numbers for nucleotide sequences in parentheses were used for Phaeosphaeria pathogens. Bootstrap values with 10,000 replications of the internal branches are indicated
$w c-1$ gene in cereal Phaeosphaeria pathogens might furnish weak splicing activities and produce multiple isoforms of mRNA transcripts (Fig. 2). It was reported that "(c/t)ag" consensus nucleotides were commonly found at -3 to -1 positions of the $3^{\prime}$ splice site in fungal genes (Kupfer et al. 2004). The presence of a low frequency of "a" at the -3 position of the 3 ' acceptor site in intron 4 would affect the splicing activity of the wc-1 gene in Phaeosphaeria pathogens.

Regulation of AS in Phaeosphaeria pathogens is unknown. Discrete repeated sequence elements in numerous genes in eukaryotes were reported to act as exonic splicing enhancers, which positively affect the accuracy and efficiency of splicing of upstream introns (Webb et al. 2005). In a search of tandem repeats (http://tandem.bu.edu), two $14-\mathrm{bp}$ continuous consensus repeats, TGGGACACACGACT and TGGGCACACGAGCT, were found at nt 1163 downstream of the stop codon (TGA) of the wc-1 gene premRNA (accession no CH445345, nt239553-239580). These repeats are purine rich, and their potential splicing mechanism needs further study.

In Neurospora crassa, expression of the wc-1 gene was regulated by AS at the $5^{\prime}$ end, which produced transcriptional isoforms in various environments. Under different light conditions, two separate transcription initiation sites located upstream of the wc-l gene-coding sequence and the one within the gene open reading frame were detected (Káldi et al. 2006). Therefore, two similar wc-1 polypeptides and one truncated molecule were produced in N. crassa. Generation of 6 AS
transcription variants from the wc-1 gene in PNw might not result from near-UV light damage, since isolate S-79-1 produced the same transcription variants in the dark (Table 5, Muñoz et al. 2009).

Functions of gene activation by the $6 \mathrm{wc}-1$ protein isoforms in PN-w were not clear and needed further study. One possibility was that all $6 \mathrm{wc}-1$ protein isoforms might confer only one function, which was to activate the downstream frq gene. However, the significance of 25 amino acid residues carboxyterminal to the Znf domain of the AREA protein, which mediated nitrogen metabolite repression in Aspergillus nidulans, in gene expressions was reported (Stankovich et al. 1993). Mutations in this region would result in enhancing, de-repressing and impairing various gene expressions (Stankovich et al. 1993). The amino acid substitutions in particular motifs of the carboxy-terminal of transcription factor Pdr3p in yeast (Saccharomyces cerevisiae) also was reported to affect gene activations and induce multidrug resistance (Nourani et al. 1997). Deletions, retentions and substitutions of amino acid residues occurred between aa971 and aa1042 in the carboxy terminal of the wc-1 polypeptide in PN-w and might have diverse gene-activation functions (Fig. 2B). The abundance of six different AS transcripts during sporulation in planta might give a fascinating insight into gene regulation during asexual sporulation on the natural host rather than artificially in vitro conditions.

It appeared that the $w c-1$ gene did not lead to direct activation of the gene(s) associated with asexual

Table 5 Variance analysis of six transcription variants of the white collar-1 (wc-1) gene in sporulated (S) and non-sporulated (NS) cultures of wheat-biotype Phaeosphaeria nodorum (PN-w) isolates

| Transcription variants | Introns spliced | Fragment sizes ${ }^{\text {a }}$ | Isolates |  |  |  | Total | Average ${ }^{\text {b }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Sn37-1 |  | S-79-1 |  |  |  |
|  |  |  | Dark (NS) | Light (S) | Dark (S) | Light (S) |  |  |
| 1 | 3 | 341 | 7 | 13 | 6 | 14 | 40 | 10.0 a |
| 2 | 3 | 332 | 6 | 13 | 10 | 5 | 34 | 08.5 a |
| 3 | 3 and 4 | 290 | 3 | 6 | 17 | 12 | 38 | 09.5 a |
| 4 | 3 and 4 | 287 | 1 | 1 | 2 | 4 | 8 | 02.0 bc |
| 5 | 3 and 4 | 281 | 8 | 7 | 8 | 6 | 29 | 07.3 ab |
| 6 | 3 and 4 | 278 | 5 | 4 | 4 | 4 | 17 | 04.3 ab |
| No. of clones |  |  | 30 | 44 | 47 | 45 | 166 |  |

[^1]sporulation in PN-w (Table 5). Inactivation of the mannitol-1-phosphate dehydrogenase gene (Mpd1) in $\mathrm{PN}-\mathrm{w}$ is reported to prevent the pathogen from sporulating on infected detached leaves (Solomon et al. 2005). Mannitol-1-phosphate dehydrogenase is one of the important enzymes for mannitol synthesis in fungi, and mannitol sugar is reported to be a major carbon and energy source for sporulation and spore germination. However, addition of mannitol to culture medium did not restore sporulation in naturally derived non-sporulated $\mathrm{PN}-\mathrm{w}$ culture suggesting that the expression of the $M p d 1$ and other sporulation genes was regulated by other upstream activator gene(s).

Based on phylogenetic analyses with nucleotide and deduced amino acid sequences of numerous genes, it appeared that genes in closely related cereal Phaeosphaeria pathogens had a common ancestor and evolved independently. In the glyceraldehyde-3phosphate dehydrogenase (gpd), $\beta$-glucosidase (bgll) and $w c-1$ genes, Pat1 was grouped together with Paa, Pat3 and PN-b as a single clade, while the RNA polymerase II (RPB2) gene of Pat1 was closely related to PN-w (Reszka et al. 2005). In the $\beta$ tubulin (tubA) and trifunctional histidine biosynthesis (his) genes, Pat1 was phylogenetically separated from all these Phaeosphaeria pathogens (Malkus et al. 2005; Wang et al. 2007). Of the studied $w c-1$ and other gene nucleotides and their deduced polypeptide sequences, the Pat2 appeared to be the most diversified, and evolved separately from all other cereal Phaeosphaeria species.

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[^1]:    ${ }^{\text {a }}$ Fragment sizes in base pairs (bp) were based on the full length PCR products amplified with primer set $7 \mathrm{~A} / 4 \mathrm{~B}$ and 1 x cDNA as templates. ${ }^{\text {b }}$ Based on Duncan test at $5 \%$ significant level

