Phylogenetics and Gene Structure Dynamics of Polygalacturonase Genes in *Aspergillus* and *Neurospora crassa*

Jin-Sung Hong¹²*, Ki-Hyun Ryu¹, Soon-Jae Kwon³, Jin-Won Kim⁴, Kwang-Soo Kim⁵ and Kyong-Cheul Park**

¹Department of Horticultural, Biotechnology and Landscape Architecture, Seoul Women’s University, Seoul 139-774, Korea
²Department of Applied Biology, College of Agriculture and Life Sciences, Kangwon National University, Chuncheon 200-701, Korea
³US Department of Agriculture-Agricultural Research Service, Western Regional Plant Introduction Station, 59 Johnson Hall, Washington State University, Pullman WA 99164, USA
⁴Department of Environment Horticulture, University of Seoul, Seoul 130-743, Korea
⁵Bioenergy Crop Research Center, National Institute of Crop Science, Rural Development Administration, Muan 534-833, Korea

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Polygalacturonase (PG) gene is a typical gene family present in eukaryotes. Forty-nine PGs were mined from the genomes of *Neurospora crassa* and five *Aspergillus* species. The PGs were classified into 3 clades such as clade 1 for rhamno-PGs, clade 2 for exo-PGs and clade 3 for exo- and endo-PGs, which were further grouped into 13 sub-clades based on the polypeptide sequence similarity. In gene structure analysis, a total of 124 introns were present in 44 genes and five genes lacked introns to give an average of 2.5 introns per gene. Intron phase distribution was 64.5% for phase 0, 21.8% for phase 1, and 13.7% for phase 2, respectively. The introns varied in their sequences and their lengths ranged from 20 bp to 424 bp with an average of 65.9 bp, which is approximately half the size of introns in other fungal genes. There were 29 homologous intron blocks and 26 of those were sub-clade specific. Intron losses were counted in 18 introns in which no obvious phase preference for intron loss was observed. Eighteen introns were placed at novel positions, which is considerably higher than those of plant PGs. In an evolutionary sense both intron loss and gain must have taken place for shaping the current PGs in these fungi. Together with the small intron size, low conservation of homologous intron blocks and higher number of novel introns, PGs of fungal species seem to have recently undergone highly dynamic evolution.

**Keywords**: *Aspergillus*, gene structure, intron loss/gain, intron phase, *Neurospora*, polygalacturonase

*Corresponding author.
Phone) +82-2-970-5613, FAX) +82-2-970-5610
E-mail) jshong@swu.ac.kr and kyongcheul.park@kangwon.ac.kr

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There are two contradictory schools of thought on the origin of introns, “intron-early” (Doolittle, 1978; Gilbert, 1978) versus “intron-late” (Palmer and Logsdon, 1991; Sverdlov et al., 2000). A synthetic theory of merging both ‘intron early’ and ‘intron late’ theories was proposed to explain the new as well as ancient introns in the current genes (de Souza, 2003; Roy, 2003). Due to the functional redundancy, extra copies of gene families of some paralogs might have undergone purifying selection while others were degenerated by accumulation of evolutionarily neutral or loss-of-function mutations (Prince and Pickett, 2002). Therefore, comparison of gene structures of orthologs and paralogs in gene families among related species might give insights into the intron evolution in eukaryotic genes.

Aspergillus is a filamentous fungal genus with over 185 species (Jones, 2007; Timberlake and Marshall, 1989). It contains important species for genetic model, human health and industry. A. nidulans and A. niger have been studied for eukaryotic cellular physiology and molecular biology by a well-characterized sexual cycle and genetic systems (Coppen et al., 1997; Pontecorvo et al., 1953). A. fumigatus is a serious life-threatening human pathogen (Denning, 1998), and A. oryzae is a beneficial food-industry fungus in the production of sake and soy sauce (Abe et al., 2006). With the importance of human health and industrial impact, genome sequencing was done in three species, A. nidulans, A. fumigatus, and A. oryzae (Galagan et al., 2005a; Machida et al., 2006). These analyses provide good venues for the evolutionary insights of the fungi by comparative genome analysis (Galagan et al., 2005a; 2005b; Payne et al., 2006; Wortman et al., 2006). Neurospora crassa is also a filamentous fungus and has been used for genetic model species (Reviews in Hynes, 2003, references therein). The completed whole genome sequences of N. crassa are now providing valuable resources in comparative genomics in fungal genetics (Galagan et al., 2003; 2005b).

Polygalacturonase (EC3.2.1.15) is a pectin-digesting enzyme containing a glycoside hydrolase family 28 domain. The number of PGs mined was 8 in A. nidulans, 6 in A. terreus, 7 in A. niger, 9 in A. fumigatus, 15 in A. oryzae, 2 in A. flavus, and 2 in N. crassa, respectively. The actual number of PGs in each species may be higher than these numbers because amino acid sequences with higher E-value than > 10^4 against query PG or with truncated glycohydrolase 28 domain less than 70% in coverage were not selected. Also, PGs without either mRNA or genomic DNA sequences were excluded from gene structure analyses. The higher number of PGs may be related to the largest genome size of A. oryzae, among the five aspergilli, as shown in other gene families.
such as cytochrome p450 enzymes and nonribosomal peptide synthases, which were 151 and 24 copies in *A. oryzae* and 14 and 65 in *A. fumigatus*, respectively (Machida, Asai et al., 2005; Payne, Nierman et al., 2006). By whole genome comparison between *A. oryzae*, *A. fumigatus*, and *A. nidulans*, sequence acquisition was found to increase the genome size in *A. oryzae* (Galagan et al., 2005a).

Polygalacturonase containing the glycohydrolase family 28 domain is one of the largest glycohydrolases encoded by a gene family in eukaryotes (Markovic and Janecek, 2001). The number of PG copies in *Aspergillus* is considerably lower than those in Arabidopsis (67 copies) and rice (48 copies) (Yokoyama and Nishitani, 2004; Kim et al., 2006).

Gene families were derived via gene duplication and subsequent regional or segmental duplication, which would ultimately be scattered throughout the genome by genome rearrangement (Lynch and Conery, 2000). If the duplication occurred prior to divergence eukaryotic phylogenetic lineage, as shown in the Cytochrome P450 gene family in four filamentous Ascomycetes fungal species, *Fusarium graminearum*, *Magnaporthe grisea*, *A. nidulans*, and *N. crassa* (Deng et al., 2007), the genes could spread into lineage-specific expansion. In our analysis of *Aspergillus* and *N. crassa*, the PGs showed some clade or sub-clade specific gene structures that support the notion of lineage-specific expansion after the genus diverged. In contrast to with the diploid-prominent organisms such as *Arabidopsis* and rice, the haploid stage is prominent during the life cycle of fungi. Therefore, the duplicated genes in fungi might have subjected different selection pressure from those of the diploid-prominent species. After undergoing duplication, one of the pair may either degenerate to a pseudogene or acquire a novel function (neo-functionalization) (Prince and Pickett, 2002). Expression of the fungal PGs in the current analysis is obvious since mRNA derived cDNA sequences are available in most of them. Functional redundancy of PGs was noted in plants such as corn (Allen and Lonsdale, 1992) and *Arabidopsis* (Hadfield and Bennett, 1998) because cell wall modification by polygalacturonases is critical in development of plants. However, pectin is not a major component making up cell wall of fungi. Hyper expression of PGs may be related with virulence of the phytopathogenic fungi by softening the plant cell wall to permit penetration by fungal hyphae during the process of infection (Markovic and Janecek, 2001). Except for *A. flavus*, the other four aspergilli in the current analysis are not known to be phytopathogenic.

**Phylogenetic analysis.** The 49 PGs that were analyzed in this study were separated into three clades, which were also divided into 11 sub-clades, based on the amino acid similarities (Fig. 1). While PGs in clade I were rhamno-PGs, those in clade II were exo-PGs. In clade III, PGs were endo-PGs except for the three PGs of sub-clade III-I. PGs in the PGs in sub-clade III-I, which were placed as the most outgroup in clade III, were exo-PGs. Hadfield and Bennett (1998) classified plant PGs into three clades of A, B, and C where clade C was composed exclusively of exo-PGs. In the analysis of diverse origin of PGs from plant, fungus, insect, and bacteria (Markovic and Janecek, 2001), fungal PGs formed two separate clusters, one exclusively with exo-PGs and the other with endo-PGs and rhamno-PGs, in which the latter were closely clustered with insect PGs. The difference of our classification from others a further analysis.

None of the sub-clades contained solely paralogous genes from a single species, which implies that PGs were diverged before species divergence within *Aspergillus* took place. Also, the separation of two PGs from *N. crassa* into separate clades implies that the PG divergence into current clades represents predated divergence of *Aspergillus* and *Neurospora* which occurred approximately 300 M (Galagan et al., 2005b; Padovan et al., 2005). Clade I contained seven PGs of *A. oryzae*, *A. terreus*, and *A. fumigatus*, which grouped into two sub-clades where bootstrap value in the node of the two sub-clades was 100, but the bootstrap values in the deepest branch in the sub-clade I-I were below 50. Nineteen PGs of *A. oryzae*, *A. terreus*, *A. fumigatus* and *A. nidulans* were in the clade II which was subdivided into four sub-clades. The bootstrap values in the nodes of sub-clades were high. However, they were lower than 50 in the nodes tying with the sub-clades. One PG from *N. crassa* (Ncra957508) was tied with four PGs of *Aspergillus* with 100 bootstrap value in the sub-clade II-II. In a previous study, five PGs of *A. oryzae* in this clade (Aory66570 in the sub-clade II-II, Aory54924, A ory63240, Aory56683, Aory61240 in sub-clade II-IV) were grouped with plant PGs (Park et al., 2008). However, they ruled out the possibility for fungal PGs being ancestral to the plant PGs since there were no PGs with intermediate gene structure between plant and fungus found in their study. The clade III was the largest one with 23 PGs from *A. oryzae*, *A. terreus*, *A. fumigatus*, *A. nidulans*, *A. niger*, *A. flavus* and *N. crassa*. There were five sub-clades of the *Aspergillus* PGs and the PG from *N. crassa* was not included in the *Aspergillus* sub-clades of clade 3.

*Aspergillus* species have evolved and diverged over 200 million years (Galagan et al., 2005a; 2005b). Whole genome duplication and subsequent gene loss predated the *Aspergillus* speciation in eukaryotic evolution (Achaz et al., 2001; Kellis et al., 2004). The distribution of PGs from each *Aspergillus* species in the sub-clades supports concept of duplication of PG predating a *Aspergillus* speciation. The
Fig. 1. Phylogenetic dendrogram and gene structures of PGs of *Aspergillus*, *N. crassa*, and *E. coli*. In the phylogenetic tree, the numbers at the nodes are bootstrap values. Lower than 50 bootstrap values were not shown. In gene structure, exons are filled bars and introns are lines. The lengths of the filled bars are relative sizes of the exons, but the lines for introns are not depicting the intron sizes for illustration. The numbers over the gene structure are the number in nucleotides of exons and introns. The colors showing the numbers in intron size are blue for phase 0, red for phase 1, and green for phase 2, respectively. The clusters in right column are the sub-clades; clades are indicated by Roman numerals.
absence of PGs from *A. niger* in clades I and II might be due to the underestimation of the PG sequences in the partially finished genome project. Among the seven pairs of the PGs in the deepest branch with 100% bootstrap values, one was a pair of paralogous sets of *A. niger* (Anig12554 and Anig42809 in sub-clade III-III) and two were orthologous PGs from *A. oryzae* and *A. flavus* (Aory58591 and Afla05020 in sub-clade III-IV, Afla05015 and Aory58737 in sub-clade III-V) which taken together support the close species relationship of *A. oryzae* and *A. flavus* as also shown by a comparison of large genome sequences (Payne et al., 2006).

**Gene structure analysis.** Five of the total 49 PGs analyzed did not carry introns. The number of introns per gene varied from one to seven among the rest of the PGs. The average number of introns per gene was 2.5, which was similar to the number of exons in whole genomes of *A. nidulans* (3.6), *A. fumigatus* (2.8), and *A. oryzae* (2.9) (Galagan et al., 2005a). The average number of introns was 3.7 in clade I, 2.9 in clade II, and 1.8 in clade III, respectively (Table 1). The lengths of introns ranged from 20 to 424 bp with an average of 65.9 bp. Most of the introns were between 31 to 90 bp at a peak of 51 to 60 bp (Fig. 2). Parsch (2003) also noted a narrow range of intron length distribution in 15 orthologous genes of *Drosophila*. Based on these observations, natural selection seems to have played a role in maintaining intron size in contrast to the selective neutrality of the nucleotide sequence variation in introns in various organisms. The average length of the introns of all genes of *A. fumigatus* was 112 bp which is similar to the introns of another filamentous fungus *Phanerochaete chrysosporium* (117 bp) (Martinez et al., 2004). The average length of the introns of *Arabidopsis* PGs was 100 bp (Torki et al., 2000) and minimal intron length in genes of *Drosophila* was 61 ± 10 bp (Yu et al., 2002). The reason for the short introns in PGs of *Aspergillus* is not clear at this moment although a negative correlation was reported between intron length and gene expression level in *Caenohabditis elegans* and *Homo sapiens* (Castillo-Davis et al., 2002). Another interesting observation was that two or more introns of the same length within a gene were clustered in 8 PGs (Fig. 1). For example, Aory57693 (I-II) had three introns of 66 bp with different intron phases. Afum742685 (II-III) had four introns of 51 bp with one of phase 1 and other 3 in phase 2. The sequences of these introns were highly variable except of those sequences for proper splicing (Fig. 3). An interesting speculation is a concerted evolution for intron size in the

**Table 1.** Intron phase distribution and average number of introns in each clade of PG genes in *Aspergillus* and *Neurospora* species analyzed

| Clade | Phase | Total | Average number of introns |
|-------|-------|-------|----------------------------|
| I     | 0     | 14    | 26                         | 3.7 |
| II    | 1     | 36    | 56                         | 2.9 |
| III   | 2     | 30    | 42                         | 1.8 |
| Total |       | 80    | 124                        | 2.5 |

Definitions of intron phase: phase-0, introns between codons; phase-1, introns between the first and the second nucleotides of a codon; phase-2, introns between the second and the third nucleotides of a codon.

![Fig. 2. Distribution of intron size in PGs of Aspergillus. The numbers in X and Y axis are the number of nucleotides and number of introns, respectively.](image)

![Fig. 3. Multiple nucleotide sequence alignment of the same length of Aory57693 and Afum742685, respectively. The 5' (GT) and 3' (AG) ends of the introns are highlighted with red and the conserved sequences for lauriate structure formation are highlighted with blue where the adenine was red.](image)
spliceosomal introns, which requires further analyses with more data sets.

Among the 124 introns, 18 introns were present in novel positions in only one PG and the rest were arranged in 29 blocks of homologous intron sets. In sub-clade III-II, the introns of the first (50 bp) of Anid6656, the second (50 bp) of Aory55286, and the second (58 bp) of Anig72931, which were differentiated only by 1 amino acid, were counted as a homologous intron set since the slightly different positions might have been derived from intron “sliding” by clustering algorithms (Stoltzfus et al., 1997). Although the homologous introns corresponded in their positions, their nucleotide sequences and length were somewhat variable. None of the intron blocks was common in all clades. Twenty-six of the 29 homologous intron blocks were sub-clade specific.

There were higher numbers of introns retaining their positions in clade III compared to those in clade I and II. Intron position conservation in Aspergillus seemed to be less stringent compared to the plant PGs reported by Park et al. (2008), where numerous introns corresponded in their positions between clades. In the analysis of 446 introns from 108 PGs of plants, they showed 19 homologous intron blocks among which only two homologous intron blocks were sub-clade specific and novel introns were as rare as 3 out of the 446 introns. Therefore, 18 novel introns of the 123 introns in the current study are significantly higher than those of plant PGs. The novel introns should have been derived from recent insertion of intronic sequences. Roy (2004) proposed transposon insertion for the origin of recent intron novelty. However, none of the introns in the PGs of Aspergillus had significant homology with known transposon sequences in BLAST analysis. Intron phase distributions among the 18 novel introns were 8 for phase 0, 7 for phase 1 and 3 for phase 2, respectively, to show no phase preference for the insertion site.

Phase distribution of introns of the 124 PGs was 80 for phase 0 (64.5%), 27 for phase 1 (21.8%), and 17 for phase 2 (13.7%), respectively, which was similar to those of plant PGs (65.5% for phase 0, 19.7% for phase 1, and 14.8% for phase 2) (Park et al., 2008), but dissimilar to those of other eukaryotic genes (50% for phase 0, 30% for phase 1, and 20% for 2) (Fedorov et al., 2002). PGs in clade 1 showed a significantly higher number of phase 1 introns compared to those of clade 2 and 3. Since loss of phase 0 introns does not occur in all clades, the absence of introns in the PGs of Aspergillus was consistent with the results of previous studies (Park et al., 2008).
not disrupt the protein coding frame, Fedorov et al. (2002) argued that intron loss might have occurred only in phase 0. In our analysis, intron loss was assumed to have occurred if one PG did not have an intron while more than two other PGs had introns at the corresponding position in a sub-clade (Fig. 4). There were 18 intron losses recorded in this study which were disproportionately distributed over the intron phases; 15 for phase 0, 2 for phase 1, and 1 for phase 2, respectively. Figure 4 is showing the intron gain and loss of sub-clade II-III where only one intron gain (phase 0) is evident SPN/WHN in Anid9045 and the other 7 introns (4 of phase 0, 2 of phase 1, 1 of phase 2) were lost in one or two orthologous genes. Both intron loss and intron gain might have occurred in leading up to the present PGs in Aspergillus and N. crassa, which is congruent with the synthetic theory of merging both ‘intron early’ and ‘intron late’ theories (de Souza, 2003; Roy, 2003).

Overall evolutionary dynamics of gene structures in PGs of Aspergillus and N. crassa with their evidences of small introns, less conservation of intron positions, and higher number of novel introns seemed to be different from plant PGs. The reason for shorter introns of PGs than other fungal genes is not clear at this moment. Although the physiological and developmental roles of PGs are obscure in fungi, their expression is abundant because the cDNA sequences are available in most of the PGs. Elucidation of the cellular and physiological functions of the PGs in fungi will help to understand the role of PGs in phytopathogenicity.

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