

The isoepoxydon dehydrogenase gene PCR profile is useful in fungal taxonomy

R Russell M Paterson

IBB-Institute for Biotechnology and Bioengineering, Centre for Biological Engineering, Braga, Portugal

Summary

This study evaluates the specificity of PCR isoepoxydon dehydrogenase (idh) primers on fungi associated with patulin production. The DNAs of 93 strains were extracted and analysed by PCR using primers of the *idh* gene of patulin biosynthesis. A single band at 620 bp was obtained on 17% of the analysed strains. Different molecular weight amplicons were observed in other strains. These were employed as binary characters for numerical analysis to obtain a dendrogram. Clusters were observed, which corresponded to morphological identifications in some cases. Amplicons at 400 and/or 500 bp were related to patulin non-detection for strains, whereas a 450 bp amplicon was associated with some Aspergillus and both of the Byssochlamys nivea strains tested. Hence, the idh primers are not specific for the gene and provide other amplicon products in other species. These results were useful providing (a) profiles of DNA to identify and classify fungi and (b) insights into patulin production. The DNA profiles in this study may be useful for determining patulin producing fungi. Obtaining multiple bands in culture-independent PCR of environmental samples by using the primers could indicate that more than one species is present.

Key words

Isoepoxydon dehydrogenase gene, Patulin, PCR, Numerical analysis, *Penicillium, Aspergillus, Byssochlamys*

Utilidad en taxonomía fúngica de los patrones obtenidos por PCR del gen de la isoepoxydon deshidrogenasa

Resumen

El interés del presente estudio fue investigar la especificidad de los cebadores para la PCR del gen de la isoepoxydon deshidrogenasa en hongos asociados a la producción de patulina. El DNA de 93 cepas fue extraído y analizado mediante PCR utilizando cebadores del gen idh implicado en la biosíntesis de la patulina. Se obtuvo una banda simple de 620 pb en un 17% de las cepas y bandas de peso molecular variable para el resto. Estos datos fueron utilizados como caracteres binarios en un análisis numérico. Se pudo comprobar que los diferentes clusters que aparecían en el correspondiente dendograma a veces se correspondían con determinados caracteres morfológicos de las cepas. Amplificados de de 400 y/o 500 pb se relacionaron con cepas no productoras de patulina y una banda de 450 pb se asoció con algunos Aspergillus y con las dos cepas de Byssochlamys nivea incluidas en el estudio. Ello demostraba que los cebadores no eran específicos para el gen y amplificaban otras regiones en algunas especies. Estos resultados demostraron ser útiles en para la identificación y clasificación de hongos y para un mejor conocimiento de la producción de patulina. Determinados patrones de DNA pueden ser útiles para detectar hongos potenciales productores de patulina. La obtención de múltiples bandas en una PCR de muestras ambientales no cultivadas puede indicar que más de una especie está presente.

Palabras clave

Gen de la isoepoxydon deshidrogenasa, Patulina, PCR, Análisis numérico, Penicillium, Aspergillus, Byssochlamys

Corresponding author: Dr. R. Russell M. Paterson

Dr. R. Hussell M. Paterson
IBB-Institute for Biotechnology and Bioengineering,
Centre for Biological Engineering
Campus de Gualtar
4710-057 Braga, Portugal
Phone: +351 253 604 400
Fax: +351 253 678 986
E-mail: russell.paterson@deb.uminho.pt

Aceptado para publicación el 18 de julio de 2007

©2007 Revista Iberoamericana de Micología Apdo. 699, E-48080 Bilbao (Spain) 1130-1406/01/10.00 €

Introduction

The significance of the polyketide patulin as a mycotoxin has increased greatly from the introduction of new European Union regulations that decrease the acceptable limits in fruit products and especially those used in baby foods. Evidence of this fact was recently reported in two patulin reviews [6,7]. Furthermore, the possibility for controlling the mycotoxin in food commodities has improved with greater understanding of the patulin metabolic pathway [17] and was further enhanced when the sequences of crucial genes were determined [4]. In particular, the sequencing of the isoepoxydon dehydrogenase gene (*idh*) which produces the enzyme to transform isoepoxydon to phyllostine, has been of greatest use.

Paterson et al. [8] were the first to employed idh primers to screen fungi specifically in the context of patulin as a mycotoxin: the fungi that were analysed were all penicillia. In addition, primers were employed seminally to analyse environmental samples as a cultural independent PCR (CIP). The analyses were extended to many other species within penicillia using culture dependant PCR (CDP), indicating that strains could be classified as to whether they were positive for idh and patulin detection [10]. Varga et al. [18] indicated the presence of idh and patulin production within Aspergillus section Clavati and included an evolutionary perspective on fungi with, or without, the trait. Paterson [12] analysed species from Penicillium, Aspergillus, and Byssochlamys that were associated with patulin production: B. nivea was positive for idh and patulin detection. In addition, positive idh results were obtained for Paecilomyces lilacinus and a strain of B. fulva [11]. Paecilomyces is the anamorphic form of Byssochlamys. Finally, an applied scheme for the classification of penicillia was developed using idh and patulin production as characters [13] to compliment other schemes e.g. Samson and Frisvad [16].

DNA sequencing demonstrated that the *idh* of *P. griseofulvum* was different from *P. expansum* [2] and that maximum parsimony trees based on rDNA and *idh* sequences were congruent [3] in the cases of these two important terverticillate penicillia. Unrelated *B. nivea* had a high degree of homology (88%) with the two penicillia: The *B. nivea* strains were identical independent of geographical region of isolation [1]. Puel et al. [15] determined that *B. fulva* could not produce patulin because of the absence of the 6-methylsalicylic acid synthase gene and *idh*. However, one other strain was *idh* positive in Paterson [11] and the situation requires clarification. White et al. [19] determined that *idh* was up-regulated under patulin permissive culture conditions.

During the investigations involving the use of *idh* primers on fungi it was noticed by the present author that other amplicons, apart from the expected product, appeared on gels at 620 bp. An assessment is made in the current report of the complete profiles for the classification of various fungi associated with patulin production using *idh* primers as such novel characters may be useful for the classification and identification of these organisms.

Material and methods

Ninety-three strains were obtained from the CABI Bioscience, UK culture collection and were identified by expert taxonomists as associated with that organisation. Methods of growth were as in Paterson et al. [10]. Isolates were grown on slopes of potato dextrose agar or 2% (w/v) malt agar. The isolates were harvested and extracted. The

PCR analysis used has been described thoroughly [10] where images of representative gels are available. The band patterns were recorded to the nearest 50bp and subjected to numerical analysis using the statistical programme, Statistical Package for the Social Sciences version 14.0 and a dendrogram was obtained (Figure 1). However, the *idh* product was recorded as 620bp, which is the typical size. Control taxa were a *Ganoderma* sp. (IMI 357185), a basidiomycete that may cause disease in oil palm; and *Aspergillus flavus* (IMI 380661), which produces aflatoxins that are polyketides, as is patulin. Neither of these fungi is known to produce patulin.

The PCR mixture consisted of 200 mM of dNTPs (Pharmacia, Herts), 1.25 units of Tth polymerase, Tth buffer (both from HT Biotechnology, Cambridge, UK), 0.8

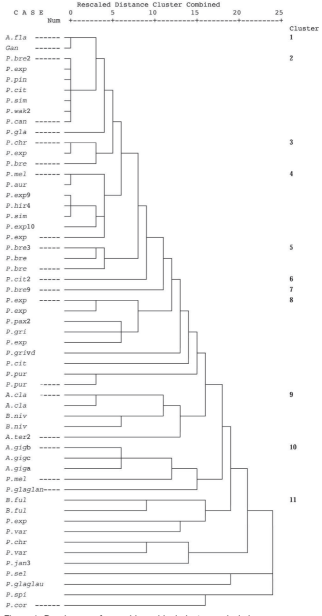


Figure 1. Dendrogram from a hierarchical cluster analysis by average linkage (between groups) of the bands resulting from the isoepoxydon dehydrogenase primers applied to total DNA from the fungal strains. The numbers after the abbreviated species names are the number of strains of the same species that were analysed. The broken horizontal lines indicate the limits of each cluster. The full names of the species can be determined from table 2.

 Table 1. List of strains employed and PCR products from the isoepoxydon dehydrogenase gene primers. The six-figure number is the IMI culture collection identification code.

Fungus				PC	R produc	ts (base pairs)	idh is 62	0	
Aspergillus	Species/strain								
Subgen. Aspergillus	•								
Sect. Flavi	A. flavus 380661						800		
Sect. Terrei	A. terreus 135817, 016043	250	450		620				1100
Subgen. <i>Fumigati</i>									
Sect. Clavati	A. clavatus 015949v	300	450		620		800		1200
Goot. Glavan	232883	300	450		620		800		1100, 1200
	A. gigantus 343711	350		500	620	700, 750	850		1100
	016154	350		500	620	750	850		
	358435	350		500	620		850		1100
Byssochlamys nivea	361545		450		620		800	900	1050
	058423		450		620		800		
B. fulva	058422		400		620	750	850		1050, 1100
	040021	200	400	500	620	750			1100, 1050
Ganoderma sp.	357185						800		
Paecilomyces variotii	321342		400	500	620				1050, 1300
	204127	200	400			700	800		1050, 1300
Penicillium									
Subgen. Aspergilloides	P. glabrum 380331						-	-	1000
	P. spinulosum 380642	350			620	700		950	1050, 1200,
Cularan Discriticiii	D pipophilum 000050	No berel							1500, 2000
Subgen. Biverticillium	P. pinophilum 380659 P. purpurogenum 380952	No bands		500		700			1100, 1400
	380948			500		100			1100, 1400
	P. waksmanii 381974, 381990		400	500	650	750			1200
Subgen. Furcatum	P. canescens 380320	No bands							1200
Subgen. Furcatum	P. citrinum 380961, 380969	140 bands							1250, 1400
	380323			500			800		2000
	380342	No bands							
	P. corylophilum 380322	350			620				1500, 1550, 200
	P. janczewskii 381948,			550			800	900	1050, 1300
	380968, 380633								
	P. melinii 304279	350	400		620	700	850		
	040216ii		400		620	700			
	P. paxilli 380639, 381272 P. selandiae 304284		400 400	500	620	750 700		950	1000, 1400
	P. simplicissimum 380333		400	300	620	700		930	1000, 1400
	380971	No bands			020				
Subgen. Penicillium	P. aurantiogriseum 265302				620	700			
	P. brevicompactum								
	381970, 380944, 380346,				620				1150, 1050
	380347,380348, 380329, 380341, 380656, 380321								
	380334				620				1500
	380330				620				
	380352							900	1100
	380349, 380643, 380648							900	
	380353		400					900	
	380634, 380645	No bands							
	P. chrysogenum 380958								1500
	380332					700			1050, 1300
	P. expansum 381952, 381969, 381268,				620				
	380345, 380350, 380665,								
	380657, 380662, 380653								
	299046				620	750			
	380324, 380326, 380327, 380325, 380344, 380337,				620				1200
	380336, 380335, 381265,								
	380970	400		F00	000	750			
	232297	400		500	620	750 750			4000
	380960	400			620	750 750			1200
	380654 381266				620	750			1200 1500
	380636	No bands							1500
	028619	200	400	500	620				1200
	P. griseofulvum 075832ii	300	400	550	620	750			1200
	P. griseofulvum				5_5				
	var. <i>dipodomyicola</i> 296935	350			620	750		900	
	P. glandicoladicola							-	
	var. glandicoladicola 321513	300		500	620		850		1150
	P. glandicoladicola var. glaucovenetum 321511	200, 300	400	500	620	700		900	
	P. hirsutum 381967,	200, 000	700	500	620	100		500	
	381951, 382063, 382064				020				

mM of primer IDH1, and IDH2, and template DNA. The IDH gene was amplified at 94 °C for 1 min, 52 °C for 1 min, 72 °C for 1 min for 30 cycles with a Hybaid, Omn-E thermal cycler. The initial denaturing step was 94 °C for 3 min and the programme was completed with an extension step of 72 °C for 5 min. The primers (GIBO BRL, Paisley) were: IDH1 sequence: 5k-CAATGTGTCGTACT GTGCCC-3k, and IDH2 sequence: 5k-ACCTTCAGTCGCTGTTCCTC-3k. A list of the 93 strains used is provided (Table 1) together with the DNA products observed.

Results

A single amplicon at 620 bp was obtained by 17% of the 93 strains tested wich were mainly strains of P. simplicissimum, P. expansum, P. brevicompactum and P. hirsutum (Table 1). The remainder strains showed 2 to 7 amplicons, with the exception of P. spinulosum, which showed 8 bands. The different banding patterns are shown in table 1 some of which are specific for species as revealed in the dendrogram (Figure 1). The strains representing each cluster are provided in table 2. From "top" to "bottom" of the dendrogram, the first cluster is formed by the *A. flavus* and the Ganoderma sp strains, (outgroup). The second cluster was made of strains, which bands were not apparent (Table 1). A single P. glabrum strain was loosely linked to this cluster. A cluster of three unrelated subgenus penicillia formed the fourth cluster. This was followed by a cluster containing most of the P. expansum strains and which were all idh positive. Approximately 50% of these strains produced also a band above 1000 bp. However, this expansum cluster included the P. hirsutum strains together with one P. aurantiogriseum. P. melinii and P. simplicissimum from subgenus Furcatum were also included in this cluster. A cluster of five idh negative P. brevicompactum strains was followed by a cluster of two P. citrinum. However, these were followed by most of the P. brevicompactum strains, which possessed the idh band at 620 bp and other higher molecular weight products at 1050 bp and 1150 bp. Those strains without the 620 amplicon showed a high molecular weight band at approximately 900 bp. A mixed collection of predominately single strains clustered below these strains, although the P. paxilli strains were identical and the P. purpurogenum were similar to each other in terms of the banding patterns. The next group was formed by A. clavatus, A. terreus and B. nivea strains, all of which were idh positive. However, A. giganteus clustered separately and were linked loosely to two penicillia. Strains with complex patterns containing an average of 5 bands were separate from the other strains and formed numerous "single strain clusters" towards the lower portion of the dendrogram. However, two strains of B. fulva clustered here and were idh positive: Three strains of P. janczewskii were *idh* negative and formed a loose group with two other idh negative strains of Paecilomyces variotii and P. chrysogenum. The other strains in this group were idh positive, the predominant bands being at 400 bp and 500 bp.

Discussion

It is apparent that the *idh* primers were not specific for the expected 620 bp product. The bands produced did not allow for complete separation of taxa into individual clusters in all circumstances. Nevertheless, the banding patterns provided are useful taxonomic characters in addition to determining the presence of *idh*. A group of predominately *P. expansum strains* was observed, all of which

Table 2. Strains included in each cluster. The six-figure number is the IMI culture collection identification code.

Clusters	Fungi				
1.	A. flavus 380661, Ganoderma sp. 357185.				
2.	P. pinophilum 380659, P. waksmanii 381974, 381990. P. citrinum 380342, P. simplicissimum 380971, P. canescens 380320, P. brevicompactum 380634, 380645, P. expansum 380636.				
Single strain	P. glabrum 380331.				
3.	P. brevicompactum 380334, P. chrysogenum 380958, P. expansum 381266.				
4.	P. expansum 299046, 381952, 381969, 381268, 380345, 380350, 380665, 380657, 380662, 380653, 380324, 380326, 380327, 380325, 380344, 380337, 380336, 380335, 381265, 380970. P. hirsutum 381967, 381951, 382063, 382064. P. aurantiogriseum 265302. P. melinii 040216ii, P. simplicissimum 380333.				
5.	P. brevicompactum 380352, 380349, 380643, 380648, 380353.				
6.	P. citrinum 380961, 380969.				
7.	P. brevicompactum 381970, 380944, 380346, 380347 380348, 380329, 380341, 380656, 380321.				
8.	P. expansum 380960, 380654.				
	P. paxilli 380639, 381272.				
	P. griseofulvum 075832ii.				
	P. expansum 232297				
	P. griseofulvum var. dipodomyicola 296935				
	P. citrinum 380323				
	P. purpurogenum 380952, 380948.				
9.	A. clavatus 015949v, 232883.				
	B. nivea 361545, 058423.				
	A. terreus 135817, 016043.				
10.	A. giganteus 343711, 016154, 358435.				
	P. melinii 304279				
	P. glandicoladicola v. glandicoladicola 321513				
11.	B. fulva 058422, 040021.				
	P. expansum 028619				
	Pa. variotii 321342				
	P. chrysogenum 380958				
	Pa. variotii 204127				
	P. janczewskii 381948, 380968, 380633				
	P. selandiae 304284				
	P. glandicoladicola var. glaucovenetum 321511				
	P. spinulosum 380642				
	P. corylophilum 380322				

produced detectable amounts of patulin and were *idh* positive. Some non-*P. expansum* species were observed in the same cluster. However, these tended to be negative for patulin production as reported in [8,10,12]. Differences within *P. expansum* require further investigation as the fungus is an important pathogen of economic food plants and a known mycotoxin producer.

The *P. brevicompactum* strains consist of at least two separate groups, one of which contains strains that were positive for *idh*. These may form the other members of the *Olsonii* series, as the morphological methods employed to identify the fungi would not necessarily separate the different species of the series (see [14]). The *Aspergillus* and *B. nivea* were separate from *Penicillium*, as would be anticipated from conventional taxonomic analysis. Related to this, the *A. clavatus*, *A. terreus* and *B. nivea* all provided a unique amplicon at 450 bp compared to the other species. More work is necessary to determine if this is a

genuine difference. The *B. nivea* were also separate from *B. fulva*, which is interesting as these species are difficult to differentiate by conventional methods. It is notable that the *B. fulva* strains were positive for *idh*, contradicting the hypothesis of Puel et al. [15] that *B. fulva* are not capable of producing patulin because the species does not possess *idh*, amongst other genes. It is necessary to consider the possibility of inhibition of PCR in the circumstances where CDP is employed [5,9]. The other banding patterns obtained provide additional characters for identification of taxa.

In conclusion, PCR using *idh* primers provides more information than a simple, single band at 620 bp. This needs to be appreciated when undertaking CIP analysis where many bands can be obtained [8] and, for example, multiple bands may not be indicative that numerous

species could be present in environmental samples. The data provided additional taxonomic characters that may be useful in classification and identification of species, in addition to indicating whether strains possess the *idh* gene.

RRM Paterson is funded by grant SFRH/BPD/34879/2007 from Fundação para a Ciência e a Tecnologia, Portugal. Dr Zofia Kozakiewicz is acknowledged gratefully for the morphological identifications which she undertook.

References

- 1. Dombrink-Kurtzman MA, Engberg AE. Byssochlamys nivea with patulin-producing capability has an isoepoxydon dehydrogenase gene (idh) with sequence homology to Penicillium expansum and P. griseofulvum. Mycol Res 2006; 110: 1111-1118.
- Dombrink-Kurtzman MA. The isoepoxydon dehydrogenase gene of the patulin metabolic pathway differs for Penicillium griseofulvum and Penicillium expansum. Anton van Leeuw 2006; 89: 1-8.
- Dombrink-Kurtzman MA. The sequence of the isoepoxydon dehydrogenase gene of the patulin biosynthetic pathway in Penicillium species. Anton van Leeuw 2006; 91: 179-189.
- Fedeshko RW. Polyketide enzymes and genes in *Penicillium urticae*. PhD Dissertation, University of Calgary, Alberta, Canada 1992.
- Hoorfar J, Cook N, Malorny B, de Medici D, Abdulmawjood A, Fach P. Diagnostic PCR: making internal amplification control mandatory. J Appl Microbiol 2004; 96: 221-222.
- 6. Jackson L, Dombrink-Kurtzman MA. Patulin. In: Sapers GM, Gorny JR, Yousef AE (Eds.) Microbiology of Fruits and Vegetables. Boca Raton, FL, CRC Press, 2006: 281-311.
- 7. Moake MM, Padilla-Zakour OI, Worobo RW. Comprehensive Review of Patulin Control Methods in Foods. Comp Rev Food Sci Food Safety 2005; 1: 8-21.
- Paterson RRM, Archer S, Kozakiewicz Z, Lea A, Locke T, O'Grady E. A gene probe for the patulin metabolic pathway with potential use in novel disease control. Biocon Sci Technol 2000; 10: 509-512.

- Paterson RRM. Internal amplification controls have not been employed in diagnostic fungal PCR hence potential false negative results. J Appl Microbiol 2007; 102: 1-10.
- 10. Paterson RRM, Kozakiewicz Z, Locke T, Brayford D, Jones SCB. Novel use of the isoepoxydon dehydrogenase gene probe of the patulin metabolic pathway and chromatography to test penicillia isolated from apple production systems for the potential to contaminate apple juice with patulin. Food Microbiol 2003; 20: 359-364.
- Paterson RRM. The isoepoxydon dehydrogenase gene of patulin biosynthesis in cultures and secondary metabolites as candidate PCR inhibitors. Mycol Res 2004; 108: 1431-1437.
- 12. Paterson RRM The isoepoxydon dehydrogenase gene of the patulin biosynthetic pathway, patulin detection, and the utility of species names in patulin-producing fungi. In: Lima N, Smith D (Eds.) Biological Resource Centres and the use of Microbes, European Culture Collections Organisation XXII. Braga, Micoteca da Universidade do Minho, 2003; 259-266.
- Paterson RRM, Venâncio A, Lima, N. A novel identification system based on 318 penicillia strains using the isoepoxydon dehydrogenase gene and patulin production. Rev Iberoam Micol 2006; 23: 155-159.
- Peterson SW. Multilocus DNA sequence analysis shows that Penicillium biourgeianum is a distinct species closely related to P. brevicompactum and P. olsonii. Mycol Res 2004; 108: 434-440.

- 15. Puel O, Tadrist S, Delaforge M, Oswald I, Lebrihi, A. The inability of Byssochlamys fulva to produce patulin is related to absence of 6-methylsalicylic acid synthase and isoepoxydon dehydrogenase genes. Inter J Food Microbiol 2006; 115: 131-139.
- Samson RA, Frisvad JC, Penicillium subgenus Penicillium: New taxonomic schemes, mycotoxins and other extrolites. Stud Mycol 2004; 49: 1-258.
- Sekiguchi J, Shimamoto T, Yamada Y, Gaucher GM. Patulin biosynthesis: enzymatic and nonenzymatic transformations of the mycotoxin (E)-ascladiol. Appl Environ Microbiol 1983; 45: 1939-1942.
- Varga J, Rigó K, Molnár J, Tóth B, Szencz A, Téren J, Kozakiewicz Z. Mycotoxin production and evolutionary relationships among species of Aspergillus section Clavati, Anton van Leeuw 2003; 83: 191-200.
- White S, O'Callaghan J, Dobson ADW. Cloning and molecular characterization of Penicillium expansum genes upregulated under conditions permissive for patulin biosynthesis. FEMS Microbiol Lett 2006; 255: 17-26.