Phylogenetic analyses and toxigenic profiles of *Fusarium equiseti* and *Fusarium acuminatum* isolated from cereals from Southern Europe

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ABSTRACT

Fusarium equiseti and Fusarium acuminatum are toxigenic species that contaminate cereal crops from diverse climatic regions. They are common in Spanish cereals. The information available on their phylogenetics and toxigenic profiles is, however, insufficient to assist risk evaluation. In this work, phylogenetic analyses were performed using partial sequences of the translation elongation factor gene (*EF*-1 α) of *E* equiseti and *E* acuminatum strains isolated from barley and wheat from Spain and other countries. The Northern and Southern European *F*. equiseti strains largely separated into two phylogenetically distinct clusters. This suggests the existence of two distinct populations within this species, explaining its presence in these regions of markedly different climate. Production of type A and B trichothecenes by the Spanish strains, examined in wheat cultures using a multitoxin analytical method, indicated that *E* equiseti could produce deoxynivalenol and nivalenol and other trichothecenes, at concentrations that might represent a significant risk of toxin contamination for Southern European cereals. *F. acuminatum* showed low intraspecific genetic variability and 58% of the strains could produce deoxynivalenol at low level. Neither species was found to produce T-2 or HT-2 toxins. The present results provide important phylogenetic and toxigenic information essential for the accurate prediction of toxigenic risk.

1. Introduction

Cereals are a dietary staple in most temperate regions. Unfortunately, they can become colonised by *Fusarium*, often resulting in severe crop disease, strongly reduced yields, and the accumulation of secondary metabolites toxic to humans and animals. Fusarium head blight (FHB) of small grain cereals is a disease complex that involves several *Fusarium* species causing largely indistinguishable symptoms. The species predominantly associated with FHB in Europe are *Fusarium graminearum*, *Fusarium avenaceum* and *Fusarium poae* (Nicholson et al., 2003; Somma et al., 2010; Xu et al., 2008). Less frequently isolated species are *Fusarium tricinctum*, *Fusarium sporotrichioides*, *Fusarium equiseti*, *Fusarium langsethiae* and *Fusarium culmorum* (Kosiak et al., 2003; Logrieco et al., 2003; Xu et al., 2005). Other species encountered sporadically include Fusarium acuminatum, Fusarium subglutinans, Fusarium solani, Fusarium oxysporum, Fusarium semitectum, Fusarium verticillioides and Fusarium proliferatum (Logrieco et al., 2003). Climate change scenarios predict increasing temperatures and variations in water availability could induce changes in the profile of FHB species on cereals. Since each species has a characteristic mycotoxin profile, the risk of mycotoxin contamination of cereals might also change (Miraglia et al., 2009).

E. equiseti is a cosmopolitan fungus distributed across regions with cool through to hot and arid climates (Leslie and Summerell, 2006). It behaves as a soil saprophyte associated with rotting fruit and other decaying plant material, and as a pathogen of a wide range of crops. This species is often detected in Norwegian cereals (Kosiak et al., 2005), but it is also common in Southern Europe (Logrieco et al., 2003), particularly in Spain (Jurado et al., 2006a; Marín, 2010; Soldevilla et al., 2005). Further, it belongs to the so-called *Fusarium incarnatum*—*F. equiseti* species complex, a genetically highly diverse group (O'Donnell et al., 2009) the members of which are associated with human disease.

F. acuminatum is widely distributed around the world, although mainly in temperate regions. It behaves as a soil saprophyte but is also found associated with the roots and crowns of plants (Leslie and Summerell, 2006; Pitt and Hocking, 2009). Its presence has recently been reported in Southern Europe, particularly Spain (Marín, 2010).

Both F. equiseti and F. acuminatum have been reported trichothecene producers (Adejumo et al., 2007; Kosiak et al., 2005; Logrieco et al., 1992). Trichothecenes are potent inhibitors of protein synthesis in eukaryotic cells (Brown et al., 2001), a consequence (mainly) of their interfering with peptidyl transferase activity. They cause different acute and severe diseases in humans and animals depending on the type of trichothecene ingested (Trenholm et al., 1989). F. equiseti produces trichothecenes such as T-2 toxin, 4-acetylnivalenol (FUS-X), deoxynivalenol (DON), nivalenol (NIV) and scirpentriol or its mono- and diacetyl derivatives (MAS and DAS) (Kosiak et al., 2005; Leslie and Summerell, 2006). It has been also reported to produce butenolide, beauvericin, equisetin (EQ), fusarochromanone (FUSCHR) and zearalenone (ZEA) (Leslie and Summerell, 2006). F. acuminatum has been reported to produce trace levels of trichothecene toxins such as diacetoxyscirpenol (DAS), monoacetoxyscirpenol (MAS), neosolaniol (NEO) and HT-2 toxin (HT-2) (Adejumo et al., 2007; Wing et al., 1993, 1994). Additionally, it produces enniatin B, steroids and moniliformin (Leslie and Summerell, 2006).

The identification of Fusarium species traditionally relies on the detection of morphological and physiological features. However, discrimination among similar species is often difficult. Neither do such methods detect intraspecific variability. Fortunately, phylogenetic analyses that make use of DNA sequence data have made significant contributions to our understanding of the systematics of Fusarium, providing species boundaries that are essential for establishing inter- and intraspecific relationships with respect to toxin profiles (Jurado et al., 2006a; Kristensen et al., 2005; Mirete et al., 2004; O'Donnell et al., 2009). Additionally, these techniques have provided the basis necessary for developing rapid, specific and accurate diagnostic methods based on PCR. These can be used to predict mycotoxin risk, providing the information necessary for early control strategies to be adopted (Jurado et al., 2005, 2006b; Knutsen et al., 2004; Konstantinova and Yli-Mattila, 2004). Several genomic sequences have been used to analyse intraspecific variability in Fusarium, including intron regions of histone coding genes, the β -tubulin gene (β *TUB*), the calmodulin gene (O'Donnell et al., 1998a; Steenkamp et al., 2002), and the translation elongation factor gene EF-1 α (O'Donnell et al., 1998b, 2000). EF-1 α gene has been used as a single-locus identification tool and is a suitable genetic marker for discriminating between Fusarium species (Geiser et al., 2004).

The literature contains little information regarding the phylogenetics of *F. acuminatum*. However, several reports suggest the existence of intraspecific diversity within *F. equiseti* (Kosiak et al., 2005; Kristensen et al., 2005; Jurado et al., 2006a). Kosiak et al. (2005) reported the existence of two groups with differences in morphological features and toxin production. However, the strains used in their study were basically all from Northern Europe. Later, a preliminary phylogenetic analysis conducted with *F. equiseti* strains from both Southern Europe (mostly isolated from Spanish cereals) and Northern Europe showed them to occupy one of two different clusters (Jurado et al., 2006a). The toxigenic profiles of the Spanish isolates were not analysed.

The aim of the present work was: (i) to examine, using partial *EF*- 1α gene sequences, the phylogenetics of *F* equiseti and *F* acuminatum strains isolated from cereal-producing regions in Spain, and (ii) to analyse the toxin profile of these Spanish strains in relation to their phylogenetics and those of Northern European strains.

2. Material and methods

2.1. Fusarium strains

Eight *F. equiseti* and 36 *F. acuminatum* strains were isolated from barley cultivated in two different regions of Spain (Castilla y Léon and Castilla-La Mancha) in 2006 (Table 1). Fungal cultures were maintained on potato dextrose agar medium (PDA) (Scharlau Chemie, Barcelona, Spain) at 4 °C and stored as spore suspensions in 15% glycerol in our collection at the Complutense University of Madrid (UCM).

2.2. DNA extraction and PCR amplification of a partial sequence of EF-1 α

Genomic DNA extractions of pure cultures of eight strains of F. equiseti and 36 strains of F. acuminatum were undertaken using three mycelium disks excised from 5 to 7 day-old PDA plate cultures, and making use of the DNeasy Plant Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer's instructions. A partial sequence of the $EF-1\alpha$ gene was obtained by PCR using the primers and protocol described elsewhere (O'Donnell et al., 1998b). PCR-amplified fragments were purified using the UltraCleanTM PCR Clean-UpTM kit (MoBio Laboratories Inc., USA) according to the manufacturer's instructions. Automated sequencing of both DNA strands was performed with the $EF-1\alpha$ pair of primers using a 3730 DNA Analyzer and the Big Dye[®] Terminator v 3.1 Cycle Sequencing Kit (Applied Biosystems, USA) at the UCM Genomic Unit. The sequences were corrected using Chromas v 1.43 software (Brisbane, Australia) and analysed and edited using Bioedit Sequence Alignment Editor v 7.0.9.0 software (Hall, 1999).

2.3. Phylogenetic analyses

Using PAUP v 4.0 b10 software (Swofford, 2003), individual maximum-parsimony (MP) phylogenetic analyses were performed for F. equiseti and F. acuminatum using the partial sequences of the *EF*-1 α gene obtained. Additional sequences previously obtained in our laboratory (Jurado et al., 2006a) were included, as were others retrieved from databases, along with sequences for species closely related to F. equiseti and F. acuminatum (Fusarium scirpi and F. avenaceum respectively). A total of 70 strains for F. equiseti and 63 for F. acuminatum were employed in the phylogenetic analyses (Table 1). An F. graminearum strain (AF212461) was used as an outgroup in both the F. equiseti or F. acuminatum MP analyses. Gaps were coded as missing data and were excluded from analyses. Unweighted parsimony analyses were performed on the individual data sets using the heuristic search option with 1000 random additional sequences with tree bisection-reconnection (TBR) branch swapping. Clade stability was assessed via 1000 bootstrap replications (Hillis and Bull, 1993). Additionally, phylogenetic analyses based on Neighbor Joining were performed using the Jukes-Cantor model (Jukes and Cantor, 1969).

Nucleotide diversities estimated as the average number of differences per site between two homologous sequences (π) were calculated using DnaSp v 4.50.3 software (Rozas et al., 2008), employing Eq. (10.5) of Nei (1987). Molecular diversities were calculated in each species for both the total sequence dataset and for the sequence dataset of each cluster. Gaps present in the alignment were excluded from analysis.

To determine the proportion of the total genetic variance attributable to inter-population differences, Wright's Fst statistic was determined via analysis of molecular variance (AMOVA) using the Arlequin v 3.01 software (Excoffier et al., 2005). The Wright's Fst statistic for the different groups was estimated using DNAsp v

Table 1 *Fusarium* strains used in this study, indicating host, origin and accession number of *Fusarium* strains used in this study, indicating host, origin and accession number, were given to different strains grouped in the same cluster. Strains in bold were obtained in the present work. The same accession number was given to identical *EF*-1 α partial number.

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lsolate name	Strain	Host	Origin	Accession number	SCI3 SCI3 SCI4
F. equiseti		-		interest	
EQUI FOID	H35A.042 C1SA 060	Barley Barley	Spain	JF490508" IF496560	F. gramme
EOI13	C1SA 063	Barley	Snain	IE496570	
EOU4	C3RA.065	Barlev	Snain	JF496571 ^j	T amining
EOU5	C1SA.073	Barley	Spain	IF496568-1 ^j	J. ucunimu
EOU6	D24SZ.090	Barlev	Spain	IF496572 ^j	
EQU7	C1SA.102	Barley	Spain	[F496573 ^j	
EQUS	C3SH.103	Barley	Spain	JF496574 ^j	ACT 14
EQU9	H2-2-5B (Type II)	Durum wheat	Spain	JF496575 ^{a,j}	ACUS
EQU10	L1-2-2 (Type II)	Durum wheat	Spain	JF496575-1 ^{a.j}	ACU6
EQU11	L3-1-2J (Type II)	Durum wheat	Spain	JF496576 ^{aJ}	ACU7
EQU12	U6-1-1 (Type II)	Durum wheat	Spain	JF496577 ^{aJ}	ACU8
EQU13	VI01066 (Type II)	Soil	Malta	M543571 [°]	ACU9
EQU14	VI0106/ (Type I)	Beet B	Denmark	Aj543558°	ACU10
EQU15	VI01068 (Type I)	Barley	Sweden	AJ5435577 ATF 47F C1b	ACU11
EQUI0	VI01009 (Type I) VI01070 (Tyme I)	Onion	Denmark	AJ545201 AI543567 ^b	ACU12
EOU18	VI01071 (Type I)	Wheat	Denmark	AI543563 ^b	ACU13
EOU19	VI01072 (Tvpe I)	Barlev	Denmark	AI543559 ^b	ACU14
EQU20	VI01079 (Type I)	Wheat	Norway	AJ543564 ^b	ACITI6
EQU21	VI01084 (Type I)	Oats	Norway	AJ543565 ^b	ACU17
EQU22	VI01087 (Type II)	Wheat	Norway	AJ543570 ^b	ACU18
EQU23	VI01093 (Type I)	Barley	Norway	AJ543566 ^b	ACU19
EQU24	VI01095 (Type I)	Barley	Norway	AJ543560 ^b	ACU20
EQU25	VI01096 (Type I)	Barley	Norway	AJ543567°	ACU21
EQU26	VI01104 (Type I)	Wheat Octo	Norway	AJ543568° ATE 42E 60b	ACU22
EQU2/	11 7P 2	Uals Soil	Canada Canada	AJ242209- D/0842055	ACU23
FOI131	34/211	Fenarto orace	Snain	D0854854 ^c	ACU24
EOU31-1	45/1.2.1	Esparto grass	Snain	D0854855 ^c	
EQU33	DAOM194187	Wheat	Canada	D0842084	ACI 127
EQU35	DA0M232362	Barley	Canada	DQ842096	ACU28
EQU36	DA0M236361	Wheat	Canada	DQ842099	AC1129
EQU38	G4_2_QC_ND_	Soybean	Canada	DQ842101	ACU30
	3_2_1_2	:			ACU30-1
EQU43	11_ZP_1	Ginseng soil	Canada 2	DQ842054	ACU30-2
EQU44	16_ZP_Z	Wheat	Canada	DQ842058	ACU30-3
EQU40	22_22_2 م77 م	WPIIC	Canada	DQ842001	ACU30-4
EQU48 E0118-1	2_21_2 5_70_1	Straw	Canada	008420777	ACU30-5
E01148-2	6 ZP 2	Straw	Canada	D0860274	ACU30-6
EOU48-3	9 ZP 2	Straw	Canada	D0842080	ACU30-7
E0U48-4	19 ZP 2	Straw	Canada	D0842059	ACU30-8
EQU48-5	27_ZP_2	Straw	Canada	DQ842063	ACU30-10
EQU49	7_ZP_1	Ginseng root	Canada	DQ842078	ACU30-11
EQU49-1	45_ZP_2	Ginseng root	Canada	DQ842076	
EQU50	60	Sugar beet	France	FJ939674 ^d	ACU30-12
EQU52	113	Sugar beet	US Sector	FJ9396784	ACU30-13
EOU53-1	90 115	Sugar beet	Sweden	FI939679 ^d	ACU30-14
EQU58	157	Sugar beet	Germany	FJ939684 ^d	ACU30-15 ACU30-16
EQU60	149	Sugar beet	Germany	FJ939680 ^d	
EQU60-1	150	Sugar beet	Germany	FJ939681 ^d	
EQU60-2	151	Sugar beet	Germany	FJ939682 ^d	ACU30-17
EQU60-3	159	Sugar beet	Germany	FJ939685 ^u	ACU31
EQU60-4	150 174	Sugar beet Sugar beet	uermany Italv	r1939686 ^d F1939686 ^d	ACU31-1
EQU62	DAOM194188	Wheat	Canada	DQ842085	ACU31-2 ACU31-3
EQU64	DA0M215463	Corn	Canada	DQ842094	ACU31-4
EQU65	DAOM232364	Wheat	Canada	DQ842098	ACU31-5
EQU66 FOLI68	NRRI 20697	kice Beet	rtaly Chile	GO5055942° GO505594 ^f	ACU31-6
EQU69	NRRL26419	Soil	Germany	GQ505599 ^f	ACU31-7 ACU32
EQU70	NRRL36136	I	I	GQ505644 ^f	ACU33
EQU71	NRRL36321 NRRL36321	Soil Pototo neel	Netherlands	GQ505647 ¹	ACU34
ЕŲU / 2 F01173	NKKL30400 NRRI 13636	Potato peei Doo	Denmark	GQ202326 ⁻ CAFAF63 ^f	ACU35
FUU J	NUNLAUOU	DUS	c n	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	ACU36

 Table 1 (continued)

Isolate name	Strain	Host	Origin	Accession number
F. scirpi				
SCI1	NRRL36478	Pasture soil	Australia	GQ505654 ^f
SCI2	NRRL29134	Pasture soil	Australia	GQ505605 ^t
SC13	NRRL26922	Soil	France	GQ505601 ^f
SCI4	NRRL13402	Pine soil	Australia	GQ505592 ^f
T contraction of the				
r. grammeun GRA1	un NRRL29169	Wheat	SU	AF212461 ^g
F actiminatio	8			
ACU1	C1RA.007	Barley	Spain	JF496578 ^j
ACU2	C1RA.038	Barley	Spain	JF496578-1
ACU3	C1RA.045	Barley	Spain 2	JF496579
ACU4	C1RA.066	Barley	Spain Spain	JF496580 TE406E00 1
ACUS	C1BA 070	Barley	uieus Snain	JF450030-1 IE496578-7
ACU7	CISA.047	Barley	Spain	JF496580-2
ACU8	C2SA.024	Barley	Spain	F496578-3 ^j
ACU9	C2SA.035	Barley	Spain	JF496578-4
ACU10	C3RA.022	Barley	Spain	JF496580-3 ^j
ACU11	C3RA.032	Barley	Spain 2 ·	JF496581 ^J
ACU12 ACU13	C3RA.U37 C3RA.075	Barley Barley	Spain Snain	JF496581-1 IE496582
ACU14	C3SA.013	Barley	Spain	JF496580-4 ^j
ACU15	C3SA.048	Barley	Spain	JF496578-5
ACU16	C3SA.049	Barley	Spain 5	JF496583
ACU17 ACU18	HISA.014 HISA.017	Barley Barley	Spain Snain	JF4965781-0 ⁷ IF496581-0 ¹
ACU19	H15A.052	Barley	Spain	IF496578-7
ACU20	H1SA.053	Barley	Spain	JF496584
ACU21	H2RA.033	Barley	Spain	JF496585 ^j
ACU22	H2RA.054	Barley	Spain	JF496585-1
ACU23	H2SA.019 U2CA.020	Barley	Spain Spain	JF496585-2 ⁰ IE406585-2 ⁰
ACU25	H2SA.043	Barley	Snain	IF496580-5
ACU26	H2SA.044	Barley	Spain	JF496585-4
ACU27	H2SA.077	Barley	Spain	JF496580-6
ACU28	H2SA.059 U2DA.021	Barley	Spain Spain	JF496580-7 TE406570 of
ACU30	H3RA.039	Barley	Spain	JF4302/8-8
ACU30-1	DA0M170306	Rice	India	ĎQ855948
ACU30-2	DAOM194173	Ι	Canada	EF521137
ACU30-3	DAOM194174	-	Canada	EF521138 D0842086
ACU30-5	DAOM215457	Corn	Canada	D0842091
ACU30-6	DA0M230348	I	Canada	EF521139
ACU30-7	DAOM238674	I	Canada C	EF521134
ACU30-8 ACU30-9	DAOM238676	1 1	Canada	EF521130 FF521135
ACU30-10	DAOM238678	I	Canada	EF521133
ACU30-11	F30	Astragalus racemosus	I	EF531698
ACU30-12	R-2165	Lavender	France	FI154733 ^h
ACU30-13	R-6678	Hibiscus	Egypt	FJ154735 ^h
ACU30-14	R-7408 Niddi 54719	Morning glory	NSA	FJ154737 ⁿ UMAG0216
ACU30-15 ACU30-16	NRRL 34218 NRRL 36147	– Human	1 1	GQ505420 ^{f, i}
		bronchial secretion		
ACU30-17	NRRL 45994	Cloaca	NSA	G0505432 ^e
ACU31	H3RA.040	Barley	Spain	JF496585-5
ACU31-1 ACU31-2	E5 NRRL 54210		1 1	പ്പാട്ട2925 HM068308
ACU31-3	NRRL 54211	I	I	HM068309
ACU31-4	NRRL 54212 NEBL 54214	I	I	HM068310
ACU31-5	NRRL 54215			HM068313
ACU31-7	NRRL 54217 11204 055	- Defect	- Constraint	HM068314 E406578 0
ACU33	H3RA.078	Barley	Spain	JF496578-10
ACU34 ACU35	H3SA.018 H3SA.050	Barley Barley	Spain Spain	JF496578-11 ^J JF496578-12
ACU36	H3SA.051	Barley	Spain	JF496578-13
			(continue	d on next page)

Table 1 (continued)

Isolate name	Strain	Host	Origin	Accession number
ACU50 ACU52	R-6934 R-9382	Soil Chrysanthemum	Australia China	FJ154736 ^h FI154738 ^h
ACU59	NRRL 54216	-	-	HM068314
F. avenaceum AVE1	VI01057	Wheat	Norway	AJ543518 ^b
 ^a Jurado et al ^b Kristensen e ^c Maciá-Vicer ^d Nitschke et ^e Amatulli et 	., 2006a. et al., 2005. nte et al., 2008. al., 2009. al., 2010. tal. 2000.			

^r O'Donnell et al., 2009. ^g O'Donnell et al., 2000.

^h Nalim et al., 2009.

ⁱ O'Donnell et al. 2010

o Donnen et al., 2010.

^j Isolates tested for trichothecene production.

4.50.3 software (Rozas et al., 2008) using Eq. (3) of Hudson et al. (1992) to take into account the genetic distances between the groups in both species.

To show more clear dendrograms for each species, the MP analyses were repeated eliminating some of the isolates that occurred in the same cluster in the first analyses, and which either had identical sequences or had differences of up to five singletons (non-parsimonious informative sites). These isolates were given the same name followed by different numbers. In both species, the dendrograms obtained in both the first and second MP analyses showed identical topologies. The same was also recorded when using the Jukes-Cantor model.

2.4. Growth conditions for toxin detection

Twelve strains of *F. equiseti*, including four isolated in a previous study (Jurado et al., 2006a) and 12 strains of *F. acuminatum* isolated in the present work were tested for trichothecene production (Table 1). Fifty gram samples of autoclaved wheat kept at a moisture content of 45% for one night were inoculated with a small quantity of each strain. Cultures were incubated at 25 °C under fluorescent light (12 h photoperiod) for four weeks, then dried at 48 °C for 24 h and ground to a fine powder. Control (non-inoculated) wheat was treated in the same way.

2.5. Chemical analysis

A multitoxin analytical method, combining high-performance liquid chromatography (HPLC), atmospheric pressure chemical ionisation (APCI) and triple quadrupole tandem mass spectrometry (LC-MS/MS) under the selected reaction monitoring (SRM) mode, was used to detect the following mycotoxins: NIV, DON and its derivatives, FUS-X, NEO, HT-2, T-2 and DAS. Standards of these toxins were purchased from Sigma–Aldrich (Milan, Italy) and stored at 4 °C in the dark. The details of the procedure have been previously described (Somma et al., 2010). The limits of detection for NIV, DON, FUS-X, NEO, HT-2, T-2, DAS were 0.0033, 0.0005, 0.0015, 0.0033, 0.0033, 0.001, 0.0025 and 0.001 μ g/g respectively.

3. Results

3.1. Phylogenetic analyses

The amplification of *EF*-1 α produced a sequence of 616 bp and 612 bp for *F. equiseti* and *F. acuminatum* respectively. In *F. equiseti*,

the total number of nucleotides analysed, excluding indels, was 584. Of these, 95 nt were polymorphic sites and 60 were parsimony-informative sites. In the case of *F. acuminatum*, excluding indels, the total number of nucleotides analysed was 589. Of these, 11 nt were polymorphic sites and 9 nt were parsimony-informative sites. Nucleotide diversities per site (π) were 0.02846 \pm 0.00197 (standard deviation) and 0.00433 \pm 0.00073 for the *F. equiseti* and *F. acuminatum EF*-1 α sequences respectively.

Results from the first (data not shown) and second MP phylogenetic analyses (Figs. 1 and 2) generated the same EF-1 α genealogies for F. equiseti and for F. acuminatum. Figs. 1 and 2 show the bootstrap 50% majority consensus trees based on MP analysis plus the consistency (CI), retention (RI) and rescaled consistency (CR) indices. For F. equiseti, both types of phylogenetic analysis (MP and Jukes-Cantor model [data not shown for the latter]) revealed three distinct clusters of isolates corresponding to F. equiseti types I and II and F. scirpi. The genetic distance, in terms of Fst estimated using DNAsp software, between F. equiseti type I and type II was 0.87166, between F. equiseti type II and F. scirpi it was 0.56589, and between F. equiseti type I and F. scirpi it was 0.63569. The Fst fixation index value, determined by AMOVA for the three groups as a whole was 0.90646. The F. equiseti type I and type II groups were homogeneous and showed low intra-group variability. The nucleotide diversity per site within these groups was $\pi = 0.00359 \pm 0.00104$ and $\pi = 0.00607 \pm 0.00110$ respectively. Variability was higher among the strains of the *F*, scirpi group ($\pi = 0.03026 \pm 0.00343$). The results did not support any influence of the host. However, the Northern European and Southern European F. equiseti strains largely separated into the type I and type II phylogenetic clusters (Fig. 1), with 16 of the 17 Spanish strains in the type II cluster and all 20 Northern European strains in the type I cluster. Strains from other locations (Table 1) fell into either the type I, type II or F. scirpi clusters.

The phylogenetic analyses revealed three defined groups for *F. acuminatum* (A, B and C) (Fig. 2) plus a set of sequences that did not belong to any of the above clusters. The Fst fixation index obtained for this species was 0.84570. The intraspecific variability for the *EF*-1 α sequences was very low, with π values between groups A, B and C less than 0.002 (data not shown).

3.2. Toxin production

To characterize the chemical profile of F. equiseti and F. acuminatum toxin production, 12 isolates of each species were examined (Table 2). None of the 12 isolates of F. equiseti tested produced 3/15Ac-DON, and none of the 12 F. acuminatum isolates produced NIV, 3/15Ac-DON, DON derivatives, NEO or DAS. T-2 and HT-2 were absent in all cultures of both species. In F. equiseti, NIV, DON, DON derivatives, FUS-X, NEO and DAS were produced by 3 (25%), 10 (83.33%), 4 (33.33%), 5 (41.67%), 3 (25%) and 3 (25%) isolates respectively (Table 3). Production levels were very variable, ranging from 227 to 7005 µg/kg for NIV, from 46 to 1035 µg/kg for DON, from 367.5 to 10,150 µg/kg for DON derivatives, from 32.5 to 16,750 µg/kg for FUS-X, from 38.1 to 1665 µg/kg for NEO, and from 9.65 to 197 µg/kg for DAS. In F. acuminatum the production of DON and FUS-X was detected in 7 (58.33%) and 1 (8.33%) isolate respectively. The production level was constant, ranging from 45.1 to 61.1 µg/kg for DON, and from 32.5 to 16,750 µg/kg for FUS-X (Table 3). Three isolates of F. equiseti produced both type A (NEO and DAS) and type B (NIV, DON, 3/15Ac-DON, DON derivatives and FUS-X) trichothecenes, whereas seven produced only type B trichothecenes. Only two isolates produced no type of trichothecene. In F. acuminatum, seven isolates produced only DON, and only one of these produced FUS-X. Five isolates did not produce any of the trichothecenes analysed (Table 2).



Fig. 1. Bootstrap 50% majority-rule consensus tree based on MP analysis of F. equiseti isolates. CI: Consistency index, RI: Retention index, CR: Rescaled consistency index.



Fig. 2. Bootstrap 50% majority-rule consensus tree based on MP analysis of F. acuminatum strains. CI: Consistency index, RI: Retention index, CR: Rescaled consistency index.

Toxin production by the *F. equiseti* and *F. acuminatum* strains isolated from Spanish cereals. NIV: nivalenol; DON, deoxynivalenol; FUS-X, 4-acetylnivalenol; NEO, neosolaniol; DAS, diacetoxyscirpenol; DON derivatives; n.d., not detected; 1, type A (NEO and DAS) and type B (NIV, DON, 3/15Ac-DON, DON derivatives and FUS-X) trichothecene producer; 2, type A trichothecene producer; 3, type B trichothecene producer; 4 non-trichothecene producer.

Isolate Name	Toxin pro	duction (µg/kg)					Toxin profile
	Type B tri	chothecene				Type A trich	othecene	
	NIV	DON	3/15Ac-DON	DON derivatives	FUS-X	NEO	DAS	
F. equiseti								
EQU1 (type II)	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	4
EQU2 (type II)	n.d.	46	n.d.	n.d.	n.d.	n.d.	n.d.	3
EQU3(type II)	n.d.	62.4	n.d.	n.d.	n.d.	n.d.	n.d.	3
EQU4 (type II)	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	4
EQU5 (type II)	119.5	67.01	n.d.	118.5	121	n.d.	n.d.	3
EQU6 (type II)	n.d.	61.5	n.d.	n.d.	n.d.	n.d.	n.d.	3
EQU7 (type II)	7005	1035	n.d.	10150	16750	1665	197	1
EQU8 (type 1)	n.d.	52.1	n.d.	n.d.	n.d.	n.d.	n.d.	3
EQU9 (type II)	n.d.	116.5	n.d.	578.5	2320	60.65	15.9	1
EQU10 (type II)	227	62.3	n.d.	367.5	1115	38.1	9.65	1
EQU11(type II)	n.d.	56.55	n.d.	n.d.	n.d.	n.d.	n.d.	3
EQU12 (type II)	n.d.	60.35	n.d.	n.d.	32.5	n.d.	n.d.	3
F. acuminatum								
ACU1	n.d.	58.65	n.d.	n.d.	n.d.	n.d.	n.d.	3
ACU8	n.d.	48.4	n.d.	n.d.	n.d.	n.d.	n.d.	3
ACU10	n.d.	60.11	n.d.	n.d.	46.7	n.d.	n.d.	3
ACU11	n.d.	61.1	n.d.	n.d.	n.d.	n.d.	n.d.	3
ACU14	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	4
ACU17	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	4
ACU18	n.d.	58.2	n.d.	n.d.	n.d.	n.d.	n.d.	3
ACU21	n.d.	45.1	n.d.	n.d.	n.d.	n.d.	n.d.	3
ACU23	n.d.	58.35	n.d.	n.d.	n.d.	n.d.	n.d.	3
ACU24	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	4
ACU29	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	4
ACU34	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	4

4. Discussion

Accurate predictions of mycotoxigenic risk basically rely on the correct identification of the fungal species in agrofood products and the determination of the toxigenic profiles of strains from different origins that might adequately represent the species. Conventional methods for identifying fungal species may overlook diversity at intraspecific level, i.e., the existence of cryptic populations. Phylogenetic analyses, however, provide a useful tool for revealing such populations, they can efficiently assist in the identification of fungal strains, and permit toxigenic profiles to be associated with particular populations or species. In the present work this approach was used to investigate a sample of isolates from two *Fusarium* species,

Table 3

Percentage of *F. equiseti* and *F. acuminatum* strains producing any of the toxins tested in this study, and the minimum and maximum quantities produced. NIV: nivalenol; DON, deoxynivalenol; FUS-X, 4-acetylnivalenol; NEO, neosolaniol; DAS, diacetoxyscirpenol; DON derivatives; n.d., not detected.

		Toxin producti (µg/kg)	on
Toxin	Producer isolates (%)	Minimum	Maximum
F. equiseti			
NIV	25	227	7005
DON	83.33	46	1035
DON derivatives	33.33	367.5	10150
FUS-X	41.67	32.5	16750
NEO	25	38.1	1665
DAS	25	9.65	197
F. acuminatum			
DON	58.33	75.1	61.1
FUS-X	8.33	46.7	46.7

F. equiseti and *F. acuminatum*, which often occur in cereals grown in Spain, and which have not been characterized to date. Their phylogenetic analysis also included a representative sample of isolates of different origin in order to situate them within a wider geographical context, particularly that of Europe.

The phylogenetic results for F. equiseti reveal the existence of wide genetic variability and two different clusters, type I and type II, that predominantly group the Northern and Southern European F. equiseti strains respectively. These results agree with those of a previous study by our group (Jurado et al., 2006a). Further, four out of five strains from Northern Europe described by Kosiak et al. (2005) fell into the type I cluster, while the other fell into the F. scirpi cluster. The existence of two distinct populations within this species might explain its presence in these two regions of markedly different climate. The genetic isolation of these two populations (which would depend greatly on the amount of sexual reproduction practised) may have occurred and genetic differences providing better adaptation to ecophysiological factors may have arisen in both. Although the life cycle of F. equiseti includes the perfect stage Gibberella intrincans (Leslie and Summerell, 2006), perithecia of G. intrincans have only been observed in laboratory experiments. Therefore, the predominant mode of reproduction of F. equiseti is considered to be asexual. If no sexual events occur between the two proposed populations, the different features and toxigenic profiles of each will persist. Additional studies would be useful for characterizing these two populations, including an examination of the ecophysiological characteristics of individuals from both populations and their relationship with their climatologically different origins.

Several studies indicate that *F. equiseti* strains can produce a wide array of toxins (Adejumo et al., 2007; Kosiak et al., 2005). In the present study, the toxigenic profile of the type II Spanish strains showed differences with respect to previously reported strains from Northern Europe (Kosiak et al., 2005), which fell into the type I cluster. In neither set of strains, however, was the production of toxins T-2 nor HT-2 detected. Kosiak et al. (2005) reported F. equiseti strains to produce higher quantities of type A trichothecenes, no detectable levels of DON nor DON derivatives, but significant amounts of NIV and FUS-X. In the present study, the Spanish F. equiseti strains did produce DON, DON derivatives and the highly toxic NIV at higher levels. Over 80% of the strains produced at least DON and 25% produced NIV as well. Further, the type II population showed diversity among individuals regarding the set of toxins produced and the relative quantities manufactured. In any event, the low level of non-toxigenic strains (less than 20%), the importance of some of the toxins produced, and their wide occurrence in cereals, highlight the potential contribution of F. equiseti to the toxin risk associated with the consumption of Spanish cereals, as well as the need to design early detection and control strategies for this species. The PCR-based F. equiseti detection protocol (Jurado et al., 2005, 2006b) may be useful in this respect.

The topology of the dendrogram obtained, which included the closely related species F. scirpi as a reference, suggests that further studies are needed for clear species distinctions to be made. Indeed, the present results, and those of a recent study on the clinically important F. incarnatum-F. equiseti species complex involving the use of multilocus DNA sequence data (O'Donnell et al., 2009), show the strong variability of F. equiseti and F. scirpi, O'Donnell et al. (2009) concluded that the F. incarnatum-F. equiseti complex contained 28 phylogenetically distinct species in which both F. equiseti and F. scirpi were represented in two distinct clusters. These F. equiseti and F. scirpi strains were also included in the present phylogenetic study, and they clustered within F. equiseti type I and F. scirpi respectively. Unfortunately, it would seem that no isolate of F. equiseti type II was included in the study performed by O'Donnell et al. (2009). Further studies should examine a fully representative sample of the diversity of F. equiseti in order to obtain a reliably robust phylogeny.

Diversity was also found for F. acuminatum, with different clusters detected, although no relationship with host or geographic origin could be established (all these groups contained F. acuminatum strains isolated from Spanish barley). However, analyses of the EF-1 α genomic sequence revealed less intraspecific variability than that recorded for F. equiseti, with a low number of parsimony-informative sites and little nucleotide diversity despite the different origins of the strains analysed. Thus, these results are suggestive of a quite homogenous population. This species is clearly phylogenetically distinct from other related species with similar morphological features within the F. avenaceum/F. acuminatum/F. tricinctum species complex (Harrow et al., 2010; Leslie and Summerell, 2006). Harrow et al. (2010) suggest that F. acuminatum may not be as abundant as F. avenaceum, probably as a consequence of biogeographic limitation, narrow host preferences and/or competitive disadvantages restricting its presence in different environments, and indicate that this might be related to the scant variability it shows. As mentioned above, a number of authors have reported different toxigenic profiles for F. acuminatum. However, to our knowledge, the production of DON by F. acuminatum has not been previously reported. In the present study seven out of 12 strains analysed were able to produce DON, although at a very low level. This suggests that this species should not be considered a DON non-producing species, although additional studies are needed to confirm this. The potential toxin risk of DON contamination should not, therefore, be dismissed. A study including both phylogenetic and toxin analyses are essential for accurate predictions of toxin risk since this could detect intraspecific variability that otherwise might be overlooked, and associate toxigenic profiles to groups/lineages/species.

In summary, the present results suggest the existence of two phylogenetically distinct populations of *F. equiseti* apparently associated with two different geographical/climatic regions: Northern and Southern Europe (type I and type II respectively). Toxin production by the type II population indicates that its members could pose a risk of cereal contamination in Southern Europe, particularly involving DON and NIV. In the case of *F. acuminatum*, although the potential DON risk is probably small, it should not be disregarded.

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