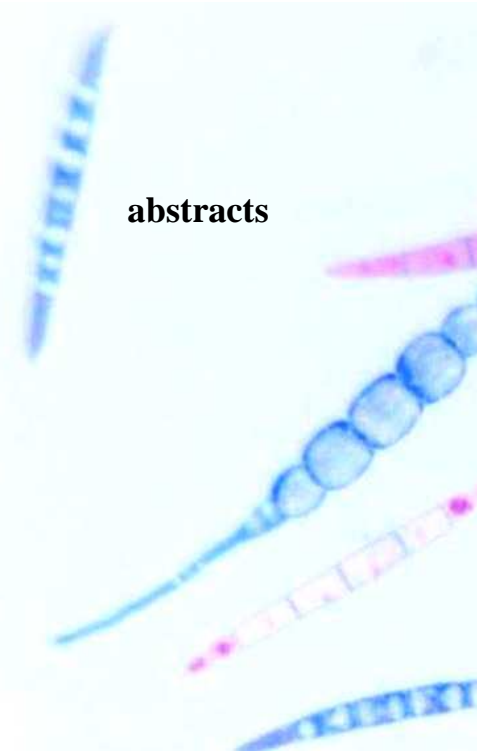


Fusarium Satellite Meeting 28-29 March 2010

abstracts

**Fusarium Satellite Meeting
Amsterdam, The Netherlands
28-29 March 2010**



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Program for Sunday 28 March

Location: Hotel Arena, 's Gravesandestraat 51

16.00-17.00 keynote presentation by Prof. Dr. Pedro Crous, Director of the Fungal Diversity Centre.

DNA barcoding in fungi: current realities and future prospects.

17.00-17.45 Fusigroup business meeting on needs and prospects for the Fusarium community and the election of the new steering committee members.

Evening - dinner cruise

Program for Monday 29 March

Oral presentations, 20 minutes each covering different themes

Location: Science Park 904, room C0.110 (ground floor)

Start 9.00

Genomics

Fusarium comparative genomics reveals lineage-specific chromosomes related to pathogenicity.

H. Corby Kistler, Li-Jun Ma, Martijn Rep and 62 co-authors of the Fusarium Genome Initiative [USA and other countries]

Exploring Lineage-specific chromosomes in *F. oxysporum* species complex

Li-Jun Ma, Shiguo Zhou, Liane R. Gale, Andrew Breakspear, Apratim Chakrabarti, Donald Gardiner, Wilfried Jonkers, Kemal Kazan, John Manners, Peter Dodds, David C. Schwartz, Jared White, Michael Koehrsen, Qiandong Zeng, James Galagan, Christina A. Cuomo Jeff Ellis, H. Corby Kistler [USA, Australia]

The Identification of a Virulence Factor-Enriched Micro-region in the Fusarium graminearum Genome.

Kim E Hammond-Kosack, John Antoniwi, Amy Freeman, Martin Urban, Sue Welham and Andrew Beacham [UK].

Secondary metabolites (toxins)

Variation in sequence and location of the fumonisin mycotoxin biosynthetic gene cluster in *Fusarium*

Robert H. Proctor, François Van Hove, Antonia Susca, Gaetano Stea, Mark Busman, Theo van der Lee, Cees Waalwijk, and Antonio Moretti [USA, Italy, Netherlands]

Coffee break 10.20-10.40

Novel pathways of regulation of deoxynivalenol production in *Fusarium graminearum*

Donald M. Gardiner, Kemal Kazan, Anca Rusu and John M. Manners [Australia]

Characterization of a novel regulatory gene involved in virulence in the phytopathogen *Fusarium graminearum*

Sean Walkowiak, Winnie Leung, Anne Johnston, Linda Harris and Gopal Subramaniam [Canada]

Epidemiology

Incidence of *Fusarium graminearum* and *Fusarium poae* from a 2-year wheat monitoring: factors promoting infection and mycotoxin contamination

Susanne Vogelgsang, Felix Wettstein and Hans-Rudolf Forrer [Switzerland]

***Fusarium* species, chemotypes and toxins in wheat from Luxembourg.**

Pasquali M., Giraud F., Cocco E., Hoffmann L., Bohn T. [Luxembourg]

Lunch 12.00-13.00

Taxonomy and identification

Taxonomy in *F. oxysporum* and detection of sp cubense tropical race 4 of the banana fusarium wilt pathogen.

Cees Waalwijk, Kerry O'Donnell et al. [Netherlands, USA]

Pathogenesis

Characterization of fatty acid regulating transcription factors of *Fusarium graminearum*.

Giang Thi Thu Le, Long Nam Nguyen and Wilhelm Schäfer [Germany]

Infection cushions and mycotoxin induction of *Fusarium graminearum* on wheat florets

Marika J. Boenisch, Peter Ilgen and Wilhelm Schäfer [Germany]

Analysis of origin of tomato wilt pathogen: From a view of the history of tomato domestication and breeding

Keigo Inami, Masato Kawabe, Akiko Okabe, Tobin L. Peever, Motoichiro Kodama, Tohru Teraoka and Tsutomu Arie [Japan, USA]

Fungal virulence and host susceptibility genes in the *Fusarium oxysporum*-*Arabidopsis* interaction

John Manners, Louise Thatcher, Donald Gardiner and Kemal Kazan [Australia]

End 14.40

***Fusarium* comparative genomics reveals lineage-specific chromosomes related to pathogenicity.**

Li-Jun Ma¹, H. Corby Kistler², Martijn Rep³ and 62 co-authors of the *Fusarium* Genome Initiative. ¹The Broad Institute, Cambridge, MA, USA, ²USDA ARS, University of Minnesota, St. Paul, MN, 55108 USA, ³University of Amsterdam, The Netherlands, et al.

Fusarium species are important phytopathogenic and toxigenic fungi, having significant impact on agriculture. Distinctively, strains of *F. oxysporum* exhibit wide host range and are pathogenic to both plant and animal species, reflecting remarkable genetic adaptability. To understand the mechanism underlying such genetic plasticity and rapid pathogenic development, we compared the genomes of three economically important and phylogenetically related, yet phenotypically distinct phytopathogenic species, *F. graminearum*, *F. verticillioides* and *F. oxysporum* f. sp. *lycopersici*. Comparative analysis revealed diverse and co-ordinately transcribed secondary metabolite biosynthetic clusters in *F. graminearum* and *F. verticillioides* as well as greatly expanded lineage-specific (LS) genomic regions in *F. oxysporum* that include four entire chromosomes that account for more than one-quarter of the genome. LS regions are rich in transposons and genes involved in host-pathogen interactions, including known effectors, enzymes targeting plant substrates or processes, and genes involved in lipid signalling and gene silencing. We found evidence for the acquisition of the LS chromosomes through horizontal transfer, which may explain the polyphyletic origin of host specificity in *F. oxysporum* and the rapid emergence of new pathogenic lineages in distinct genetic backgrounds.

Exploring Lineage-specific chromosomes in *F. oxysporum* species complex.

Li-Jun Ma¹, Shiguo Zhou², Liane R. Gale³, Andrew Breakspear³, Apratim Chakrabarti⁴, Donald Gardiner⁵, Wilfried Jonkers³, Kemal Kazan⁵, John Manners⁵, Peter Dodds⁴, David C. Schwartz², Jared White¹, Michael Koehrsen¹, Qiandong Zeng¹, James Galagan¹, Christina A. Cuomo¹, Jeff Ellis⁴, H. Corby Kistler³

¹The Broad Institute, Cambridge, MA, USA, ²University of Wisconsin-Biotechnology Center, Madison, WI USA, ³USDA ARS, University of Minnesota, St. Paul, MN, 55108 USA, ⁴CSIRO Plant Industry, Black Mountain Laboratories, Black Mountain ACT 2601 Australia, ⁵CSIRO Plant Industry, Queensland Bioscience Precinct, St Lucia, Brisbane, Queensland, 4067 Australia

The Fusarium comparative genomes of *F. graminearum* (*Fg*), *F. verticillioides* (*Fv*) and *F. oxysporum* (*Fo*) revealed greatly expanded lineage-specific (LS) chromosomes in *Fo*. These mobile LS chromosomes contribute to fungal pathogenicity and host-specificity, providing an explanation for the polyphyletic origin of host specificity and the emergence of new pathogenic lineages in the *F. oxysporum* species complex (FOSC). Following this discovery, a comparative study focusing on the members of FOSC was developed to: 1) examine genome structural variation and confirm the presence of LS chromosomes among different isolates using optical mapping; 2) determine gene content variation among these selected isolates using next-generation sequencing (NGS); 3) identify all lineage-specific genes using targeted sequencing of the LS chromosomes and RNA sequencing via whole transcriptome approaches. One human isolate and 11 plant pathogenic isolates that represent eight *formae speciales* were included in the study. Preliminary results from the optical mapping confirm the existence of LS chromosomes in different isolates. Genomic data generated using NGS detects genome-wide patterns of mutation among isolates during their brief time of evolutionary divergence. RNA-seq data shows great promise in detecting novel genes encoded in the LS chromosomes and for determining gene expression profiles under different conditions.

The Identification of a Virulence Factor-Enriched Micro-region in the *Fusarium graminearum* Genome.

Kim E Hammond-Kosack, John Antoniwi, Amy Freeman, Martin Urban, Sue Welham and Andrew Beacham

Centre for Sustainable Pest and Disease Management, Department of Plant Pathology and Microbiology, Rothamsted Research, Harpenden, Herts, AL5 2JQ, UK.

Initial studies using a novel bioinformatics and statistical approach, identified a genomic micro-region in *Fusarium graminearum* that appeared to be enriched for homologues of verified pathogenicity genes in the other pathogenic species. Detailed analysis of this micro-region by a combination of bioinformatic and reverse genetics approaches has confirmed this micro-region has a role in *F. graminearum* pathogenicity and has led to the identification of a novel virulence determinant. This micro-region which is also found in other *Fusaria* genomes appears to be distinctly different from the virulence-associated biosynthetic and secreted protein clusters identified so far in other pathogenic fungi. Further investigation will reveal more about the properties of this small genomic region.

Rothamsted Research receives grant-aided support from the Biotechnology and Biological Sciences Research Council (BBSRC) of the UK. Andrew Beacham is supported by a BBSRC industrial CASE studentship awarded to Syngenta.

Variation in sequence and location of the fumonisin mycotoxin biosynthetic gene cluster in *Fusarium*.

Robert H. Proctor¹, François Van Hove², Antonia Susca³, Gaetano Stea³, Mark Busman¹, Theo van der Lee⁴, Cees Waalwijk⁴, and Antonio Moretti³

¹US Department of Agriculture, ARS, NCAUR, Peoria, Illinois, USA; ²Mycothèque de l'Université catholique de Louvain (MUCL), Louvain-la-Neuve, Belgium; ³National Research Council, ISPA, Bari, Italy; ⁴Plant Research International B.V., Wageningen, The Netherlands.

Several *Fusarium* species in the *Gibberella fujikuroi* species complex (GFSC) and rare strains of *F. oxysporum* can produce fumonisins, a family of mycotoxins associated with multiple health disorders in humans and animals. In *Fusarium*, the ability to produce fumonisins is governed by a 17-gene fumonisin biosynthetic gene (*FUM*) cluster. Here, we examined the cluster in *F. oxysporum* strain O-1890 and nine other species (e.g. *F. proliferatum* and *F. verticillioides*) selected to represent a wide range of the genetic diversity within the GFSC. Flanking-gene analysis revealed that the *FUM* cluster can be located in one of four genetic environments. Comparison of the genetic environments with a housekeeping gene-based species phylogeny revealed that *FUM* cluster location is correlated with the phylogenetic relationships of species; the cluster is in the same genetic environment in more closely related species and different environments in more distantly related species. Additional analyses revealed that sequence polymorphism in the *FUM* cluster is not correlated with phylogenetic relationships of some species. However, cluster polymorphism is associated with production of different classes of fumonisins in some species. As a result, closely related species can have markedly different *FUM* gene sequences and can produce different classes of fumonisins. The data indicate that the *FUM* cluster has moved within the *Fusarium* genome during evolution of the GFSC and further that sequence polymorphism was sometimes maintained during the movement such that clusters with markedly different sequences moved to the same genetic environment.

Novel pathways of regulation of deoxynivalenol production in *Fusarium graminearum*.

Donald M. Gardiner, Kemal Kazan, Anca Rusu and John M. Manners

CSIRO Plant Industry, 306 Carmody Rd, Brisbane 4067, Queensland, Australia

Fusarium head blight of wheat, caused by *F. graminearum*, is one of the most important diseases of wheat not only because of yield losses but also the contamination of grain with trichothecene toxins such as deoxynivalenol (DON). An intriguing aspect of the pathogen's biology is that the production of DON occurs at much higher levels during the infection process than during axenic culture, even on plant-derived media such as autoclaved grain. Presumably, the fungus produces toxins in response to unknown signals of plant origin. We have used a reporter strain of *F. graminearum* carrying a *TRI*-gene promoter linked to the green fluorescent protein gene to identify compounds that induce high levels of DON production in culture. Through this system, we have identified a number of amines and polyamine compounds that induced the genes involved in the biosynthesis of DON to levels equivalent to those observed during infection, and resulted in high concentrations (>1500 ppm) of DON being produced in culture filtrate. Polyamines and other inducers increase in concentration in heads following inoculation suggesting that they may act as *in planta* DON inducers. The Affymetrix *Fusarium* GeneChip® was used to compare gene expression during culture under DON-inducing conditions, to that under non-inducing conditions. The polyamine inducer agmatine differentially regulated a large number of fungal genes, including both known and uncharacterised putative secondary metabolite biosynthetic gene clusters. *In silico* prediction of binding sites for the transcriptional regulator (TRI6) controlling *TRI* gene expression and gene expression analysis in a *TRI6* mutant of *F. graminearum* showed that three of the differentially regulated genes were under the control of TRI6. Gene knock-out mutations of two of these genes resulted in mutants with massively increased production of deoxynivalenol and, under our infection conditions, increased aggressiveness towards wheat. Our results identify a novel mechanism of negative regulation of DON production in *F. graminearum*.

Characterization of a novel regulatory gene involved in virulence in the phytopathogen *Fusarium graminearum*

Sean Walkowiak*, Winnie Leung, Anne Johnston, Linda Harris and Gopal Subramaniam

Agriculture and Agri-Food Canada | Agriculture et Agroalimentaire Canada, 960 Carling Avenue, Ottawa, Ontario, K1A 0C6

A study performed by Alexander et al. suggested *Tri15* may be negatively regulating some of the genes in the trichothecene biosynthetic pathway in *F. sporotrichioides*. In contrast, disruption of *Tri15* in *F. graminearum*, neither affected its ability to synthesize 15-ADON nor its pathogenicity. This study explores the role of *Tri15alt*, a homologue of *Tri15*. *Tri15alt* encodes for a protein that has three zinc fingers, two of which are highly homologous to the zinc fingers found in *Tri15*. Targeted disruption of *Tri15alt* in *F. graminearum* did not compromise the biosynthesis of 15-ADON. However, pathology studies performed on a susceptible variety of wheat (Roblin) revealed that *Tri15alt* disrupted strain is more virulent than the wildtype strain. We have performed microarray analyses on this mutant and results will be presented to identify genes involved in virulence.

Alexander N.J., S.P. McCormick, T.M. Larson and J.E. Jurgenson. 2004. Expression of *Tri15* in *Fusarium sporotrichioides*. *Curr Genet* 45: 157-162

Incidence of *Fusarium graminearum* and *Fusarium poae* from a 2-year wheat monitoring: factors promoting infection and mycotoxin contamination.

Susanne Vogelgsang, Felix Wettstein and Hans-Rudolf Forrer

Agroscope Reckenholz-Taenikon Research Station ART, Reckenholzstrasse 191, 8046 Zurich, Switzerland

In a 2-year investigation, wheat samples and respective information on cultivation techniques were collected from Swiss growers. Wheat kernels were examined for incidence of *Fusarium* head blight (FHB) causing species and mycotoxin content (LC-MS/MS). From a total of 248 samples originating from 16 out of 26 cantons, three FBH species were dominant: *F. graminearum*, followed by *F. poae* and *F. avenaceum*. The average deoxynivalenol (DON) content was 940 ppb and thus barely below the European limit for unprocessed cereals (1250 ppb). With pre-crop maize and conservation tillage versus ploughing, an average DON content of 2670 ppb or 470 ppb, respectively, was obtained. We also measured the content of other trichothecenes and zearalenone (ZEA). Nivalenol (NIV) and ZEA contents in samples from the same two cropping systems showed a similar pattern as those of DON (NIV: average of 30 and 14 ppb for the two cropping systems; ZEA: 190 and 12 ppb). However, no correlation was found between *F. poae* incidence and the NIV content. Thus, we assume for *F. graminearum* the presence of NIV chemotypes in certain geographic areas. Current fungal incidence and toxin measurements from a third year of monitoring, chemotype investigations as well as in-depth analyses of the cultivation data should contribute to elucidate factors that influence the occurrence and toxin contamination by the most prevalent *Fusarium* species on wheat. The hypothesis of *F. graminearum* NIV chemotypes is in line with recent observations from other European wheat surveys. Hence, it would be worthwhile to discuss the establishment of a concerted initiative assembling data on fungal prevalence and toxins from various geographic areas in order to establish a European map on FHB chemotypes.

Fusarium species, chemotypes and toxins in wheat from Luxembourg.

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Seventeen wheat production sites in Luxembourg were chosen to represent the 3 different climatological conditions in Luxembourg. Grains sampled during the years 2007 and 2008 from all the 17 locations were analysed for the presence of Deoxynivalenol (DON), Nivalenol (NIV), T-2 and HT-2 and Zearalenone (ZON).

Seventy-five percent of the investigated fields were contaminated by DON (range 0-8111 µg/kg). Eight fields were also contaminated by NIV. Our study represents the first report of fusariotoxines in harvested grains in Luxembourg.

Species determination of *Fusarium* populations isolated from the same grains used for toxin analysis was carried out according to morphological criteria and confirmed by species-specific PCR. Major species found were *F. graminearum*, *F. poae*, *F. avenaceum* and *F. culmorum*.

In order to verify if chemotype may have an effect on toxin accumulation in grains, *F. graminearum* and *F. culmorum* were screened by using chemotype-specific primers.

NIV chemotype was the less frequent one and distributed non-homogeneously.

Investigating factors that may favour the presence of NIV chemotype in wheat grains, maize as preceding crop showed a significantly positive effect, suggesting its biological role as an ecological niche for the nivalenol chemotype.

Nivalenol presence in grains was correlated to the number of *F. culmorum* with NIV chemotype detected in grains (and not to *F. poae* nor to *F. graminearum* with NIV chemotype). Our finding suggests the potentiality of prediction of toxin content by analysing the chemotype of *Fusarium* population from the fields.

A molecular diagnostic for tropical race 4 of the banana.

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This study analysed genomic variation of the translation elongation factor 1 α (TEF-1) and the intergenic spacer region (IGS) of the nuclear ribosomal operon of *Fusarium oxysporum* f. sp. *cubense* (Foc) isolates, from different banana production areas, representing strains within the known races, comprising 20 vegetative compatibility groups (VCG). Based on two single nucleotide polymorphisms present in the IGS region, a PCR-based diagnostic tool was developed to specifically detect isolates from VCG 01213, also called tropical race 4 (TR4), which is currently a major concern in global banana production. Validation involved TR4 isolates, as well as Foc isolates from 19 other VCGs, other fungal plant pathogens and DNA samples from infected tissues of the Cavendish banana cultivar Grand Naine (AAA). Subsequently, a multiplex PCR was developed for fungal or plant samples that also discriminated *Musa acuminata* and *M. balbisiana* genotypes. It was concluded that this diagnostic procedure is currently the best option for the rapid and reliable detection and monitoring of TR4 to support eradication and quarantine strategies.

Characterization of fatty acid regulating transcription factors of *Fusarium graminearum*

Giang Thi Thu Le, Long Nam Nguyen and Wilhelm Schäfer

Molecular Phytopathology and Genetics, Biocenter Klein Flottbek, University of Hamburg, Germany.

F. graminearum is a major pathogen of cereals worldwide. Recently, we identified secreted lipases as general virulence factors. To study the regulation of lipase genes we investigate the role of so called cutinase transcription factors. The cutinase transcription factor protein family is extensively present and conserved among filamentous fungi. We identified several putative cutinase transcription factor genes in *F. graminearum* and characterized them by gene disruption. Disruption of *Far1* (*fatty acid regulator1*), a homolog of *Aspergillus FarA* gene, indicates that it is important for long chain fatty acid utilization. Disruption of *Far2*, a homolog of the *Aspergillus FarB* gene, demonstrates that *Far2* is required for very short chain fatty acid assimilation by the fungus. *Lr1* (*lipase regulator1*), which belongs to the *Far1* clade, leads to reduced total extracellular lipolytic activity and transcriptional repression of several lipase genes in culture. These results suggest that *Lr1* mediates expression of genes involved in fatty acid hydrolysis. In summary, our results show that transcription factors of the plant pathogen *F. graminearum* are involved in regulation of genes important for fatty acid assimilation and lipid hydrolysis.

Infection cushions and mycotoxin induction of *Fusarium graminearum* on wheat florets.

Marike J. Boenisch, Peter Ilgen and Wilhelm Schäfer

Molecular Phytopathology and Genetics, Biocenter Klein Flottbek, University of Hamburg, Germany.

The mycotoxin producing pathogen *Fusarium graminearum* is the causal agent of Fusarium head blight (FHB) of small grain cereals on fields worldwide. Although *F. graminearum* is one of the best investigated phytopathogens, detailed information about fungal development on host surfaces and the penetration strategy of the pathogen is limited. We established a bioassay that allows a comprehensive investigation of the inoculated host surfaces. Detection of mycelium was facilitated by constitutive expression of a *dsRed* reporter gene, thereby allowing bioimaging with white light and fluorescence stereomicroscopy, as well as confocal laser microscopy. Additionally, a *GFP* coupled *TRI5*-promotor allows monitoring of the mycotoxin desoxynivalenol production during infection. Combining bioimaging with scanning electron microscopy we identified penetration structures and alterations of the host surface on a three-dimensional level. For the first time we demonstrate the formation of infection cushions during *F. graminearum* infection on host tissues. We discovered that the infection cushions are attended by an intensive subcuticular growth stage of the pathogen and exhibit a high mycotoxin induction. Surprisingly, a *TRI5*-k.o. mutant exhibits the same infection strategy and efficacy. We conclude that mycotoxin production is specifically induced in infection structures but not necessary for penetration.

Loss-of-function of the avirulence gene, *SIX4*, by transposon-insertion in tomato wilt pathogen *Fusarium oxysporum* f. sp. *Lycopersici*.

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²Kochi Agricultural Research Center, Kochi, Japan

Fusarium oxysporum f. sp. *lycopersici* (*FOL*) is the soilborne pathogen of tomato wilt. In the pathogen, three races 1, 2, and 3 have been determined based on the specific pathogenicity to tomato varieties. The compatible or incompatible relationships between races and varieties can be explained by the interactions between the avirulence genes carried by *FOL* and resistance genes carried by tomato varieties according to gene-for-gene theory (Flor, 1956). For example, race 1 carrying *AVR1* is avirulent to tomato cultivars with a resistance gene *I*, and races 2 and 3 carrying no *AVR1* is virulent to the tomato cultivars with *I*. Houterman et al. (2008) reported *SIX4* corresponding to *AVR1* in *FOL* race 1.

In 2008 a strain of *FOL* (KoChi-1), overcoming *I*-mediated resistance, emerged in Japan. Although KoChi-1 is not race 1, PCR revealed that KoChi-1 carried *SIX4*. Sequence analysis showed that *SIX4* ORF in KoChi-1 was truncated by a transposon (759 bp). The inserted transposon is non-autonomous and belongs to *hAT* family (Hua-Van et al., 2000). According to the Genome Databases of Broad Institute, 72 copies of the identical transposon exist in *F. oxysporum*. Integration of an intact *SIX4* derived from a race 1 isolate into KoChi-1 genome complemented avirulence to a tomato cultivar possessing *I*. This is the first report of an avirulence gene truncated by transposon-insertion in *F. oxysporum*.

Fungal virulence and host susceptibility genes in the *Fusarium oxysporum*-*Arabidopsis* interaction.

John Manners, Louise Thatcher, Donald Gardiner and Kemal Kazan

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The interaction of the root infecting fungal pathogen *Fusarium oxysporum* with *Arabidopsis* is a highly tractable system for a molecular analysis of fungal virulence and host susceptibility and immunity. We have completed a rigorous analysis of 6868 T-DNA insertion mutants of *Arabidopsis* Col-0 ecotype, selected lines with altered disease phenotype ($P < 0.01$) and retested these to identify mutants with significant & reproducible increased resistance or susceptibility. Second allele insertions are currently being tested to provide certainty on specific gene functions. These studies have identified ~100 novel genes with previously unidentified roles in immunity and susceptibility to this pathogen. To complement this we have identified a small range of fungal mutants with altered pathogenicity and virulence. One of these includes mutants in the *SIX4* gene which is required for full virulence. Experiments are underway to attempt to match putative functions in the host that are necessary for susceptibility with functions in the pathogen required for virulence. Initial experiments are focusing on the role of host jasmonate signalling in susceptibility and how the pathogen may intervene in this.

Title: Role of a mucin-like membrane protein in signalling and pathogenicity of *Fusarium oxysporum*.

Elena Perez-Nadales and Antonio Di Pietro

The soilborne fungus *Fusarium oxysporum* causes vascular wilt in a wide range of plant species by penetrating roots, invading the cortex and colonizing the vascular tissue. Fmk1, a mitogen activated protein kinase (MAPK) orthologous to *S. cerevisiae* Fus3 and Kss1, is essential for plant infection. The signalling components upstream of the Fmk1 cascade are currently unknown. In yeast, the membrane mucin Msb2 functions at the head of the filamentous growth MAPK cascade. We identified a gene from *F. oxysporum* whose predicted product has sequence homology with yeast Msb2 and shows a similar domain structure, including an N-terminal signal sequence, a predicted serine-threonine rich mucin region, a transmembrane domain and a short cytoplasmic tail. Western analysis using an HA-tagged Msb2 version showed that *F. oxysporum* Msb2 is an integral membrane protein which is expressed during vegetative growth and tomato root infection. Deletion mutants lacking *msb2* showed reduced phosphorylation levels of Fmk1, suggesting that Msb2 may function upstream of this MAPK. In contrast to $\Delta fmk1$ strains, $\Delta msb2$ single and $\Delta fmk1/\Delta msb2$ double mutants exhibited enhanced sensitivity to the cell wall-targeting compounds Congo Red and Calcofluor White, suggesting that Msb2 also signals in an Fmk1-independent pathway functioning in the cell wall stress response. The $\Delta msb2$ strains showed delayed invasive growth across cellophane membranes and significantly reduced virulence on tomato plants. Our results suggest that Msb2 is a mucin-like membrane protein that contributes to invasive growth and virulence of *F. oxysporum* by signalling partly via the Fmk1 MAPK cascade.

Transcriptional analysis of the response to extracellular pH changes in *Fusarium graminearum* *Pac1* mutants and effect on trichothecenes B accumulation

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Fusarium graminearum infects wheat and maize and produces type B trichothecenes. These mycotoxins cause serious problems when consumed via contaminated cereals. *Tri* genes, located in the “*Tri* cluster”, are responsible for the biosynthesis of trichothecenes B. *In vitro*, *Tri* genes of *F. graminearum* strain CBS 185.32 are expressed at day 3 with the toxin starting to accumulate one day later. Strikingly, the induction of *Tri* genes expression always seems concomitant with a sharp pH drop in the media. Acidic pH seems a determinant factor for induction, as neither the toxin nor the *Tri* genes are detectable at neutral pH. The pH regulation of gene expression in fungi is mediated by the *Pac1* transcription factor involved in various secondary metabolites regulation. An *FgΔPac1* deletion mutant and a strain expressing a constitutively active form (*FgPac1^C*) were constructed in *F. graminearum*. Expression of this constitutive *Pac1^C* factor strongly reduces expression of *Tri* genes and toxin accumulation at acidic pH. Unexpectedly, deletion of *Pac1* does not induce toxin production at neutral pH. However, it causes an earlier *TRI5* induction and toxin accumulation at acidic pH. In order to determine the interference with other *Tri* genes regulatory mechanisms, exploring general transcriptional response to pH variation for mutants and wild-type strains were also performed using microarrays. Preliminary results will be presented.

Characterization of the serine-/threonine protein kinase *gad8* in the phytopathogenic fungus *Fusarium graminearum*.

Katja Schäfer, Jörg Bormann and Wilhelm Schäfer

Molecular Phytopathology and Genetics, Biocenter Klein Flottbek, University of Hamburg, Germany

Fusarium graminearum is the causal agent of Fusarium head blight and a highly destructive disease of all cereals.

A knock-out of the serine-threonine protein kinase *gad8* in this fungus leads to a severe inhibition of growth *in vitro* as well as of virulence *in planta*. To determine the role of this gene in the regulatory pathways involved in fungal development, a complementation study in yeast was accomplished.

The aimed homologue of *gad8* in *S. cerevisiae*, is coding for the AGC-type protein kinase. Ypk1 and is part of a signalling module which activates a phosphorylation cascade. This pathway is stimulated by sphingolipid base phytosphingosine, a metabolic product of sphingolipids which are upregulated by several stimuli and serve as second messenger in signal transduction pathways and controls a wide range of cellular processes including growth, cell wall integrity, stress resistance, endocytosis and aging. Ypk1 is a high copy suppressor gene that allows growth when the synthesis of sphingolipids is inhibited.

We could show that the Δ pk1 strain in *S. cerevisiae* is less stress tolerant than the wt in regard to temperature and toxic agents. Additional tests with the complemented yeast as well as studies with the *gad8* knock-out in *F. graminearum* will be presented.

The tetraspanin FgPls1 is involved in fitness and pathogenicity of *Fusarium graminearum*

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Tetraspanins are a group of membrane proteins with four transmembrane domains common among different species like mammals, fish, insects, and fungi. Their capacity is complex and involved in a broad range of physiological processes where they function as “molecular facilitator”, interacting with proteins from different families like integrins, proteoglycans, growth factors and growth factor receptors as well as members from the Ig superfamily. In fungi, three different families of tetraspanins have been characterized so far: Pls1, which is found in both ascomycetes and basidiomycetes, Tsp2, which is unique to basidiomycetes, and Tsp3, which is exclusively found in ascomycetes. Pls1 knock out mutants in three appressorium producing plant pathogenic fungi, *Magnaporthe grisea*, *Botrytis cinerea*, and *Colletotrichum lindemuthianum* were non pathogenic on their respective host plants which proves that Pls1 is a pathogenicity factor in these fungi. In this study, we identified a tetraspanin Pls1 like protein, named FgPls1 in the wheat scab fungus *F.graminearum*. Results show that FgPls1 is important for vegetative growth and the production of macroconidia as well as pathogenicity of *F. graminearum*.

RNA interference in *Fusarium graminearum* using intron containing hairpin vectors.

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RNA interference is a posttranscriptional gene silencing mechanism. In multinucleate filamentous fungi, RNAi is an efficient way to silence multiple gene copies at the same time. We used transgenic strains of *Fusarium graminearum* expressing the GFP gene from *Aequorea victoria* and the DsRed gene from *Discosoma sp.*, respectively, under control of the *gpdA* promoter. To maintain a permanent down regulation of the target genes, we used an intron containing hairpin vector carrying fragments of the DsRed/GFP gene. To monitor the effects of silencing, we investigated the down regulation on the transcriptional level through RT-PCR and quantitative real-time PCR, and on translational level through western blotting, fluorescence microscopy, and measurement of the fluorescence level in 96 well plates. Most of the transformants showed a down regulation compared to their respective wild type, albeit to various degrees. Our experiments confirmed the results concerning siRNA based silencing by hairpin constructs in *Fusarium graminearum*. We report here that 400bp sense and antisense fragments are sufficient to maintain silencing. The applied method will be an efficient means to down regulate lethal genes and create new insights in the genome of *Fusarium graminearum*.