Aspergillus strain typing in the genomics era

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Abstract: Multiple reasons may justify a need for strain typing purposes, but the most common reason is to delineate the epidemiological relationships between isolates. The availability of whole genome sequences has greatly influenced our ability to develop highly targeted and efficient strain typing methods for these purposes. Some strain typing methods may serve dual goals: not only can they be used to discriminate between multiple isolates of a certain species, they can also aid in the recognition, identification, description and validation process of a fungal species.

Key words: AFLP, coding tandem repeats, high resolution typing, identification pathogenic aspergilli, MLST.

INTRODUCTION

Strain typing can fulfill many needs both in clinical settings and otherwise. Among the many potential applications for strain typing are outbreak analysis and environmental monitoring, patient monitoring and treatment follow-up, local and global epidemiology, database construction, strain identification (e.g. with production organisms) and many more. Apart from these applications at the subspecies level, molecular methods are also increasingly used at the genus level for the definition and recognition of fungal species.

Over the years, many different molecular methods have been developed for Aspergillus strain typing. Because of its clinical significance, these methods were primarily directed at A. fumigatus. The most promising techniques are either PCR based, such as analysis of microsatellite length polymorphisms (MLP)/short tandem repeats (STR) (Bart-Delabesse et al. 1998; de Valk et al. 2005) and amplified fragment length polymorphism (AFLP) analysis (Warris et al. 2003; de Valk et al. 2007), or based on non-coding repetitive sequences (such as the Mut1 element) in combination with restriction fragment length polymorphisms (RFLP) (Girardin et al. 1993). Use of these and other methods has been reviewed by Varga (2006). Three recent additions to this diverse list are multilocus sequence typing (MLST) (Bain et al. 2007), coding tandem repeats (Balajee et al. 2007; Levdansky et al. 2007) and retrotransposon insertion-site context (RISC) typing (de Ruiter et al. 2007). Depending on the exact reason for strain typing and on the technical resources in a particular setting, the choice for either of these methods could be appropriate.

Classically, without the availability of genomic sequence information, the process of developing a new strain typing method often involved many laborious selection and optimisation experiments. At present, in the genomics era, the availability of whole genome sequences has had a great impact on our options to develop novel and state-of-the-art fingerprinting methods. We now can develop new fingerprinting methods using highly targeted approaches with much higher à priori chances of being successful than before. Naturally, as more genomic sequence information is becoming available, these chances will continue to increase.

Here, we will present a number of applications for several of these genotyping methods and discuss the impact of the availability of genomic sequence data on the applications of these methods.

High resolution exact strain typing using short tandem repeats

Microsatellites or STR’s are ubiquitously present in the genomes of many fungi including Aspergillus spp. Microsatellites, as tools for the identification of and discrimination between individual organisms, already have a relatively long history in human forensic applications where they currently comprise the global “gold standard” in the identification process of individuals. The use of STR’s offers a number of technical advantages over many other fingerprinting techniques including: ease of amplification, multiplex options, extremely high discriminatory power, an exact unambiguous (numerical) and highly portable and exchangeable typing result, ability to detect mixed samples, construction of databases, etc. Because of these advantages, there is a growing interest in the use of STR based methods for strain typing in the microbial field as well.

Bart-Delabesse et al. (1998) reported the first application of microsatellites for A. fumigatus. These markers were obtained by screening genomic DNA libraries of A. fumigatus for suitable, microsatellite containing sequences, a process that proved quite laborious in the pre-genomics era. A panel of 4 dinucleotide repeats was selected that performed well in comparative genotyping experiments (Lasker 2002). Recently, based on genomic sequence data that has become available, de Valk et al. (2005) reported a novel set of 9 tandem repeats for typing A. fumigatus isolates, the so-called STRA1 assay (STR’s of A. fumigatus). In contrast to the previously developed typing scheme, this panel also contained tri- and tetranucleotide repeat markers and, in addition, all loci contained a single uninterrupted repeat element. By using multicolor multiplex approaches with these novel markers, large numbers of isolates can be analyzed in a short period of time. Because of the larger number of loci, the STRA1 assay yielded a superior discriminatory power for typing A. fumigatus isolates. In Fig. 1, a graphical representation of
the diversity with the \textit{A. fumigatus} population is shown. The minimal spanning tree represents 99 presumably unrelated \textit{A. fumigatus} isolates. Almost all isolates could be discriminated from each other. The ones that could not be discriminated by the STR\textsubscript{Af} assay also proved to be indistinguishable using other molecular methods such as AFLP analysis. Furthermore, the STR\textsubscript{Af} assay proved to be an extremely robust typing assay. It has been shown that deliberate and significant changes to the experimental protocol did not lead to wrong typing results with this assay (de Valk \textit{et al.} 2007a).

The key element in the use of microsatellites is to translate the electrophoretic mobility of the obtained fragment (reflected as the size of the fragment in bp and obtained on a high resolution electrophoretic platform) to the corresponding number of repeats. Unfortunately, this mobility is dependent on many factors such as the presence/absence of denaturing compounds, the sieving matrix that is used, the exact base composition and sequence of the fragment, run temperature, presence of different fluorescent labels and even something that may appear only trivial such as the sizing marker (de Valk \textit{et al.} 2007a; Tu \textit{et al.} 1998; Vainer \textit{et al.} 1997). In order to transfer a microsatellite based assay to a different electrophoresis platform, a careful calibration of the new platform has to be established. Similar to the situation in human forensics, a series of allelic ladder was constructed that contain reference fragments with established repeat numbers. By running these allelic ladders, every platform can be calibrated to yield exchangeable typing data with any given set of isolates (de Valk \textit{et al.} submitted for publication). Thus, the STR\textsubscript{Af} assay has all the key ingredients to be successfully used for global standardisation of \textit{A. fumigatus} typing.

\textbf{Simultaneous identification and strain typing}

In recent years, there has been a growing interest in the use of more accessible techniques such as MLST approaches for fungal identification purposes and strain typing. This approach that is exclusively based on sequencing data has the advantage of the development of accurate databases totally reliable for taxonomy. However, whereas MLST performs well at the genus and species level, in the case of \textit{Aspergillus} (and in contrast to other species like \textit{Candida}) the discriminatory power at the subspecies level turns out to be disappointing (Bain \textit{et al.} 2007).

AFLP analysis is a highly discriminatory method at the intraspecies level. In AFLP analyses, fragments are amplified from random locations throughout an organisms’ genome in a highly reproducible manner (Vos \textit{et al.} 1995). The discriminatory power of AFLP analysis equals that of the STR panels (de Valk \textit{et al.} 2007b) and \textit{Afut1} RFLP analysis. However, like with any other fingerprinting method based on DNA banding patterns, its long-term stability and reproducibility may be quite challenging. Development of AFLP fingerprinting requires no prior sequence information. However, depending on the genome composition (GC-content and distribution, presence of multicopy elements), certain combinations

\begin{figure}
\centering
\includegraphics[width=\textwidth]{Fig1.pdf}
\caption{Minimal spanning tree of 99 presumably unrelated \textit{A. fumigatus} isolates based on microsatellite data. The tree was generated using the multi-state categorical similarity coefficient. Each circle represents a unique genotype. The size of the circle corresponds to the number of isolates with the same genotype. Genotypes connected through a shaded background differ by a maximum of 2 out of 9 markers. The numbers correspond to the number of different markers between the genotypes. No less than 96 different genotypes were discriminated. Data are taken from de Valk \textit{et al.} (2005).}
\end{figure}
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Fig. 2. Example of AFLP analysis showing the discriminatory power as well as the ability to discriminate between isolates belonging to different species. The figure shows 16 fingerprints from A. fumigatus cultured from 16 IA patients (1 isolate per patient). Based on the differential presence or absence of one or more bands, all isolates can be discriminated from each other. One isolate with a clearly different fingerprint turned out to be N. fischeri. The dendrogram was calculated by UPGMA clustering using the Pearson correlation coefficient. The scale bar indicates the percentage similarity.

of restriction enzymes and selective residues could prove to be more suitable than others. At present, based on available genomic sequence data, one can predict in silico which fragments will be obtained with any known genome (Bikandi et al. 2004).

Whereas AFLP has originally been presented as a tool for strain typing purposes, it is also very well suited to simultaneously resolve isolates belonging to different species from each other. This is relevant in the case where the identification of a microorganism may be uncertain such as the species from the morphologically very similar section Fumigati. If not properly identified, use of such typing data could easily lead to false conclusions. In a way, AFLP can be considered the perfect PCR alternative to DNA-DNA reassociation studies. Classical DNA-DNA reassociation studies rely on sequence similarities. If two species share a certain amount of sequence information, it is to be expected that they will also share a certain amount of similarity in banding patterns from an AFLP fingerprint. In fact, this has already been demonstrated for a variety of bacterial and fungal species. According to our own observations, AFLP fingerprints of isolates of the same species are usually > 60 % similar whereas fingerprints for isolates representing different species are usually < 40 % similar (Fig. 2). We also used AFLP analysis for confirmation of the identity of 67 isolates representing 26 species in Aspergillus section Fumigati. These isolates have previously been identified using a variety of other methods. Although the majority of isolates were correctly identified, this exercise clearly showed that: i) some isolates were misidentified, ii) that some recognised species are comprised of multiple clearly discernible subgroups and iii) that several isolates that are currently recognised as different species should actually be grouped into a single species. Thus, AFLP can very well complement the use of MLST and other methods in the recognition and validation process of fungal species.

Strain typing based on coding tandem repeats

Coding tandem repeats are adjacent in-frame coding DNA sequences of 2 to 200 nucleotides in length that are directly repeated; these repeated units may be completely identical or partially degenerate (Li et al. 2004). The number of these coding-repeat copies often varies among different isolates leading to expansion or contraction of amino-acid blocks. Coding repeats have been observed in a number of prokaryotic and eukaryotic genomes where they play an important role in generating variability in cell-surface immunogenic antigens and adhesins, thereby evading the immune system or enhancing pathogenicity (Gravekamp et al. 1998; Jordan et al. 2003; Verstrepen et al. 2005). The inter-strain variability in the number of coding sequences can also serve as an extremely robust and rapid typing technique. Sequence analysis of a single, highly-variable gene, Protein A (spa) or clumping factor (cflb) has been successfully applied to strain differentiation amongst Staphylococcus aureus isolates which generally exhibit low variability and poorly discernible population genetic structure (Shopsin et al. 1999). Recently, an analysis of the genome of A. fumigatus identified as many as 292 genes with internal repeats. Fourteen of 30 selected genes showed size variation of their repeat–containing regions among 11 clinical A. fumigatus isolates. One of these, the cell wall protein Afu3g08990 is involved in conidial germination and adhesion (Levdansky et al. 2007). Importantly, the repeat containing region of Afu3g08990 or CSP (cell-surface protein) was shown to vary significantly between A. fumigatus isolates from various origin (Levdansky et al. 2007; Balajee et al. 2007) (Fig. 3). By simply sequencing the Afu3g08990
repeat region in the various isolates and performing a phylogenetic analysis using the maximum parsimony method, it was possible to successfully “sub-type” fifty-five epidemiologically linked A. fumigatus isolates from six nosocomial outbreaks of invasive aspergillosis. The results were concordant with another discriminatory genotyping technique, the Afut1 RFLP typing method. However, while Afut1 typing is labor and time intensive, needs specialised equipment and is not high throughput, Afu3g08990 / CSP-typing requires only the ability to perform PCR and have access to an automated sequencer. Also, interpretation of the sequence information does not require sophisticated algorithms nor dedicated software and thus can be seamlessly integrated into any clinical microbiology laboratory.

It is worthy to note that in the A. fumigatus genome there is a substantial enrichment of putative cell-surface and/or secreted proteins that contain internal repeats. While 2.9 % of the ~9 900 genes in the A. fumigatus genome code for protein repeats, at least 12.5 % of all putative cell-wall encoding genes do so, a greater than 4-fold increase. This suggests that as found in a number of other fungal genomes, repeats in A. fumigatus may play an important role in generating variability in cell-surface immunogenic antigens and adhesins, thereby evading natural predators in its natural environment and the immune system in its inadvertant host.

CONCLUSIONS

Many different reasons may exist to explain the wish for being able to discriminate between different isolates of a given species. In the light of increasing reports of misidentified isolates, there is also clearly a need for more accurate and accessible methods to identify a fungal isolate to the species level. Ideally, these two parameters are achieved in one typing/identification method. Based on the availability of genomic sequence data, highly targeted approaches allow strain typing methods to be directed at (a) highly specific part(s) of a specific fungal genome. Several of these typing methods can be used for typing purposes and may simultaneous be used for identification confirmation of fungal isolates: e.g. any clinical A. fumigatus isolate lacking the characteristic amplification products using a highly species specific typing method such as the STRAF assay or any isolate lacking the typical banding patterns obtained with the Afut1 RFLP method is most likely not a true A. fumigatus. The true identity of such an isolate yet remains to be established using other methods. In contrast, more universally applicable typing methods that are not hampered by the species barrier such as MLST and AFLP analysis are not only suitable for strain typing and identification purposes but they could additionally serve as parameter in the description and validation process of fungal species and in the delineation of the relationships between them.

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![Fig. 3. The TR region in A. fumigatus gene Afu3g08990 containing a leader sequence and GPI anchor shows size variability. The TR region of Afu3g08990 was amplified by PCR in 11 clinical A. fumigatus isolates, including the genome-sequenced reference strain Af293. This figure shows size variation of the TR region between the strains.](image-url)
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