A new method for fungal genetics: flow cytometry of microencapsulated filamentous microcolonies.

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INTRODUCTION AND OBJECTIVE

Genetic analysis of non-filamentous microorganisms is facilitated by the isolation of consistent, well-defined colonies on solid media and the handling of individual cells by flow cytometry. In contrast, some filamentous fungi are hard to be analyzed using these procedures; in particular by flow cytometry. The combination of single spores microencapsulation and large particle flow cytometry is a possible alternative analysis of filamentous for the fungi. Microencapsulation allows the early detection of fungal growth by monitoring the development of hyphae from encapsulated individual spores. Mycelium proliferation inside the microcapsules can be detected using COPASTM large particle flow cytometry. Here we show the successful application of the Flow Focusing® technology to the filamentous microencapsulation of fungi in monodisperse alginate microspheres, using Aspergillus and Trychoderma as model systems. Using a Cellena® Flow Focusing microencapsulator, we managed to produce monodisperse microparticles containing individual spores and to develop microcolonies of these fungi upon germination in the appropriate conditions. Proliferation inside the particles was monitored by microscopy and large particle flow cytometry without requiring fluorescent labeling. Sterility was preserved during the microencapsulation procedure, preventing undesired contaminations. Conditional mutants were utilized to demonstrate the feasibility of the method. This procedure allows for the handling, screening and analysis of clonal colonies in liquid culture. Examples of applications are provided.

MATERIALS AND METHODS





Figure 1 : (left) Cellena® portable micro encapsulator (right) Spherical sized monodisperse alginate microcapsules



Figure 2 : Microencapsulation technology





Figure 3 : COPAS® Large Particle Flow Cytometer and sorting principle.

These instruments re now available to automate the process of sorting large (20-1,500 micron) particles in a continuously flowing stream at a rate of 10-50 objects/second. Using object size (TOF), optical density (EXT), and intensity of fluorescent markers as sorting criteria, selected objects in this size range, can be dispensed in multi-well plates for further analysis. A gentle pneumatic sorting mechanism located below the flow cell avoids harming or changing sensitive objects, thereby making the instrument suitable for live biological materials or sensitive chemistries.

Multiple fluorescence excitation and emission wavelengths are available. Today, automation, increased sensitivity, and quantitative measurements enable larger / faster screens of model organisms, cell clusters and bead-based assays.

RESULTS AND DISCUSSION

Monitoring the proliferation of *Trichoderma reseei* spores by optical microscopy and flow cytometry

In this test the spores were encapsulated in $400\mu m$, 1,66% alginate capsules . After encapsulation the beads were incubated in shaking flasks and samples were recovered after 2h, 4h, 6h, 8h, 10h, 12h and 14h of incubation. Aliquots were analyzed by COPAS SELECT flow cytometry allowing the measurement of different optical parameters: size (TOF), optical density (EXT), green self-fluorescence and red self-fluorescence signals.



Figure 4 : Pictures of encapsulated spores during germination at different times points.

The germination of spores is associated with an increase in density as measured by the COPAS system. These measurements are represented in the next graphs showing the increase of EXT over time (left) and the EXT distribution within the bead (right).





Figure 5 : Graphs showing the increase in EXT (scattered light) and the distribution of EXT during time of incubation



Figure 6 : Optimisation of encapsulation by increasing alginate the percentage

CONCLUSION

Encapsulation of filamentous fungi can be used for genetic screenings using large particle flow cytometry of single spores. Encapsulation allows to screen for growth / no growth and/or fluorescence. Examples will be given during the poster presentation

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