Cell type specific transcriptional profiles of the dimorphic pathogen *Penicillium marneffei* reflect distinct reproductive, morphological and environmental demands.

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Figure S1  Confirmation of differential gene expression. The expression of a subset of differentially expressed genes identified in the microarray analysis was confirmed by RT PCR on RNA isolated from vegetative hyphae at 25° (A), asexual developing cultures at 25° (B) and from yeast cells at 37° (C). Genes shown are as follows: Hyphal specific; PMAA_076130 meiosis induction protein kinase (Ime2), PMAA_071450 hypothetical protein, PMAA_065680 nucleoside-diphosphate-sugar epimerase, PMAA_053710 hypothetical protein and PMAA_018570 allergen Asp F3. Asexual development specific; PMAA_082030 conidial pigment biosynthesis 1,3,6,8-tetrahydroxynaphthalene reductase Arp2, PMAA_082040 conidial pigment biosynthesis scytalone dehydratase Arp1, PMAA_010220 glutaminase GtaA, PMAA_075300 C2H2 type conidiation transcription factor BrlA and PMAA_097290 hypothetical protein. Yeast specific; PMAA_057450 ferrooxidoreductase Fet3, PMAA_040300 sodium P-type ATPase, PMAA_018640 cytochrome P450 monoxygenase, PMAA_031950 4-hydroxypyruvate dioxygenase and PMAA_091310 hypothetical protein.
Figure S2  Synteny across the ystA region is only conserved in species closely related to *P. marneffei*. Diagrammatic representation of the genomic region surrounding ystA in *P. marneffei*, *T. stipitatus* and *A. fumigatus* showing the predicted genes and gene product. This region is syntenic in *P. marneffei* and *T. stipitatus* but disrupted in *A. fumigatus* where each side occurs on a different chromosome. There is no homology to ystA in *A. fumigatus.*
Figure S3  The ystA genomic region encodes a single transcript. (A) Diagrammatic representation of the three reading frames with predicted ORFs longer than 50 aa in an 850 bp region identified by the 1E11 microarray probe. The DD11 and DD12 arrows represent oligonucleotides used to generate single stranded probes of this region to identify transcripts. The shaded regions (yellow) with arrows define the three ORFs in this region that are 68, 65 and 69 aa in length. The CC51-LL21, CC51-poly(T) and DD1-CC52 oligonucleotide primer pairs were used in RT-PCR experiments to define the boundaries of the transcript in this region. (B) Single stranded radiolabelled probes were generated using DD11 and DD12 and these were used to probe a northern blot with total P. marneffei RNA isolated from yeast cells at 37°. A single transcript of approximately 850 bases was detected with the DD12-derived probe only. To check the quality of the probes each was tested in a Southern blot hybridisation using total genomic DNA digested with BamHI. The predicted 6.5 kb genomic DNA fragment was identified with both probes. (C) RT-PCR using the primer pairs indicated in A was performed with total RNA isolated from yeast cells grown at 37°. To assess the amount of contaminating genomic DNA in the samples both reverse transcriptase (+) and no reverse transcriptase (-) reactions were performed.
Tables S1-S7
Available for download at http://www.g3journal.org/lookup/suppl/doi:10.1534/g3.113.006809/-/DC1

Table S1  Oligonucleotides used in this study
Table S2  Relative expression data for the three cell states.
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Table S4  MeV phase-specific curated clusters.
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Table S7  GO associations for early and late expression clusters.