Additional file 1

Fungal artificial chromosome mining of the fungal secondary metabolome

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Supplementary Figures

26

27 a

pSMART® BAC pyrGAMA1
full size
19301 bp

pSMART® BAC pyrGAMA3
rev full size
19301 bp

pSMART® BAC pyrGAMA2
full size
19301 bp

pSMART® BAC pyrGAMA4
rev full size
19301 bp
Figure S1. Panel a: FAC vectors: pSMARTBACpyrGAMA1~4, each has two Not I sites (N) flanking the cloning site (B, Bst XI) within the BAC end sequencing primers SP6 and T7. In addition to the chloramphenicol resistance gene (camR), loxP and cos sites, pyrGA represents the pyrG gene from Aspergillus parasiticus, AMA1 is the replication origin of fungal artificial chromosome, genes parA and parB are for active partitioning and gene sopC is to ensure that each daughter cell gets a copy of the shuttle BAC plasmid, gene repE is for BAC plasmid replication and regulation of copy number, and oriV is the BAC replication origin. Panel b: A. nidulans FAC transformants using pSMARTBACpyrGAMA3 vector.
Figure S2. Preparation of HMW genomic DNA from *A. terreus* and random shear FAC cloning results. Panel a: *A. terreus* HMW genomic DNA ranging from 20~200kb. Panel b: CHEF gel electrophoresis and *NotI* digestion of random selected FAC clones, the average insert size was estimated at ~110kb. M, Lambda ladder Marker.
Figure S3. Three additional CHEF gels of *E. coli-Aspergillus* shuttle FACs that were successfully transferred from transformed strains of *A. nidulans* back into *E. coli*. The examples of recovered FAC clones shown here include 9O3 (cluster 30, ~100 kb), 9A23 (cluster 25, ~80 kb), and 7A10 (cluster 56, ~90kb) from top to bottom panel. The first and last lanes are DNA Lambda ladder Markers, the 2nd and 3rd lane(s) on the left hand side of the gels is the control FAC used to transform *A. nidulans*, and all of other lanes are randomly selected FAC clones recovered. All control and recovered FACs were digested with *Not I*. 
Figure S4. Antibiotic activity test of 14 FAC clones. Ten μl out of 150 μl methanol extract from FAC transformants cultured on GMM plate for 7 days at 37 °C was loaded on small disc (diameter: 1cm) for antimicrobial activity test against *Aspergillus* spp., *Candida albicans*, *Bacillus cereus*, *Micrococcus luteus* and *Pseudomonas aeruginosa*. Antibiotic activity was observed against *Bacillus cereus* with two FAC extracts.