Growth and metabolism of *Beauveria bassiana* spores and mycelia

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**Abstract**

**Background:** Fungi are ubiquitous in nature and have evolved over time to colonize a wide range of ecosystems including pest control. To date, most research has focused on the hypocrealean genera *Beauveria bassiana*, which is a typical filamentous fungus with a high potential for insect control. The morphology and components of fungi are important during the spores germination and outgrow to mycelia. However, to the best of our knowledge, there is no report on the morphology and components of *B. bassiana* spores and mycelia. In this work, the growth and metabolism of *Beauveria bassiana* spores and mycelia were studied. High performance liquid chromatography-mass spectrometry (HPLC-MS) was employed to study the metabolism of *B. bassiana* spores and mycelia. Principal component analysis (PCA) based on HPLC-MS was conducted to study the different components of the spores and mycelia of the fungus. Metabolic network was established based on HPLC-MS and KEGG database.

**Results:** Through Gompertz model based on macroscopic and microscopic techniques, spore elongation length was found to increase exponentially until approximately 23.1 h after cultivation, and then growth became linear. In the metabolic network, the decrease of glyoxylate, pyruvate, fumarate, alanine, oxaloacetate, dihydrothymine, ribulose, acetylcarbinimine, fructose-1, 6-bisphosphate, mycosporin glutamicol, and in spores indicated that the change of the metabolin can keep spores in inactive conditions, protect spores against harmful effects and survive longer.

**Conclusions:** Analysis of the metabolic pathway in which these components participate can reveal the metabolic difference between spores and mycelia, which provide the tools for understand and control the process of spores germination and outgrow to mycelia.

**Keywords:** *Beauveria bassiana*, Gompertz model, PCA, HPLC-MC, Metabolism
knowledge, there is no report on the morphology and components of *B. bassiana* spores and mycelia. The change of morphology and components can reveal the connection between spores and mycelia, and provide a systems-level understanding of the cell.

Despite its importance, only a limited number of methodologies have been developed for morphology and components analysis. This is primarily due to the characteristics of most components that display high polarity, nonvolatility, poor detectability, and overall similar properties [17]. Recently, high performance liquid chromatography - mass spectrometry (HPLC-MS) equipped with electrospray ionization (ESI) detection has been used for components analysis [18–21]. It is a robust, sensitive, and selective technique, and also has become popular for quantitative and qualitative analyses. In the present study, the morphology of *B. bassiana* spores and mycelia were studied by combining macroscopic and microscopic techniques. And then HPLC-MS coupled with PCA were used to distinguish different metabolites of mycelia and spores. In addition, metabolic pathway was established based on HPLC-MS and KEGG database. Tracking metabolite changes under different conditions not only provides direct information on metabolism but is also complementary to gene expression and proteome analysis [22, 23]. Metabolomics, which can be defined as the measurement of the level of all intracellular metabolites, has become a powerful new tool for gaining insight into cellular function. The aim of the study was to reveal the reason of keep survive longer and infective of spores by compare significant change in metabolites between spores and mycelia. And provide the tools for understand and control the process of spores germination and outgrow to mycelia.

**Results and discussion**

**Spore germination kinetics**

The germination of spores takes place when the spores are introduced into a proper environment, which requires proper nutrition and special conditions. The spore germination can be divided into three phases: spore swelling, germ tube emergence and germ tube elongation [9]. In the first phase, spores begin to swell to increase their dormant diameter significantly until a germ tube emerges (second phase). The two phases in early growth are supported by mobilization and utilization of storage compounds in the spores. In the third phase the elongation of the germ tube is observed, which contributes to biosynthesis and extension by uptake and metabolism of nutrients from the medium [15].

The spore germination kinetics was investigated in the study. The values for hyphal length were measured with the aid of Image-Pro Plus software in a series of images monitoring the growth of *B. bassiana* spores on PDA at 26 °C, and the duration of the germination phase was estimated. Until the 6th hour of the cultivation, no germ tubes could be spotted, although an increase in the mean diameter of spores due to swelling. (Fig. 1).

Figure 1 showed typical forms of *B. bassiana* spores and hyphae in their development. Tubes emerged from 8 h to approximately 11 h. About 10 h after cultivation, most of the spores had their tubes emerged. At that moment the spores entered the phase of tube elongation. The hyphae remained unbranched till the 20th hour. After 28 h, the objects became difficult to observe and analyse due to their increasing concentration and aggregation of hyphal elements.

Models were used to describe the behavior of microorganisms [24]. In the study, the modified Gompertz model was employed to detect the relation between spore elongation length and culture time, which can provide useful information about the variance of the germination time for individual spores. Figure 2 showed changes of the most important morphological parameters. It can be seen that the modified Gompertz equation described satisfactorily the length of spore elongation over time for *B. bassiana*, with coefficients of determination ($R^2$) of 0.9825. The three phases, spore swelling, germ tube emergence and germ tube elongation can be observed distinctly. In the phase of germ tube emergence and in the early hyphal development, the measured length appeared to increase exponentially until approximately 23.1 h after cultivation, when growth became linear. The above results are in agreement with the findings of Trinci [25, 26], who examined the kinetics of hyphal extension of several fungi, and considered the observed transition from exponential to linear growth can be attributed to the weakness of the hyphal tips to incorporate the increasing material that is being supplied or the deficiency of transporting material from distal hyphal regions.

**Principal component analysis**

PCA based on HPLC-MS data was used to study the metabolomic differences of mycelia and spores extracts from *B. bassiana*. PCA is an unsupervised pattern recognition method, which means that no prior knowledge concerning groups or tendencies within the data sets is necessary. PCA is usually employed to reduce the dimensionality of the data and extract essential information from large, mixed data sets [27]. A score plot is applied for the grouping of samples by reducing the dimensionality of the data. The complex data was reduced to two principal components PC1 and PC2, which can represent most of the components. The values of the two variables for the observations are called factor scores, two factors scores can be interpreted geometrically as the projections of the observations onto the
whole components. The data sets exhibiting similarities are clustered together, and those that are different are placed further apart [28].

Acid compounds were extracted in negative mode, while alkaline compounds were extracted in positive mode, to study the whole components extracted from spores and mycelia, both positive mode and negative mode must be employed to PCA. As shown in Fig. 3, from both positive mode and negative mode, PCA score plots employed in this study found that the mycelia were clearly separated from the spores of *B. bassiana*, and the spores or mycelia were grouped correctly together respectively. In the positive mode, both principle components were significant: PC1 accounts for 35.7 % of the total variance and PC2 accounts for 52.8 %. The first two principal components (PC1 and PC2) explain more than 80 % of the total variance. In the negative mode, PC2 leads to classification of the two groups and accounts for 58.3 % of the total variance, whereas PC1 denotes 29.5 % of the total variance. The high-resolution ESI-MS data for some metabolites are shown in Fig. 4. These findings were in good agreement with HPLC–MS based metabolic profiles (Tables 1 and 2). In Table 1, twenty-eight major components were identified from *B. bassiana* mycelia by HPLC-MS method, eleven of which were different from that in spores, such as glycerophosphocholine, palmitic acid, linoleic acid, phosphatidylethanolamine. For spores, thirty-six compounds were extracted, seventeen of which were distinguished from mycelia, such as succinic anhydride, dihydouracil, mannitol, sphinganine (Table 2).

**Metabolic pathway analysis**

Nutritional limitations or increase in cell density leads *B. bassiana* to produce dormant, environmentally resistant spores. The complex morphological changes that occur during sporulation are thought to be highly controlled by metabolic networks [29]. However, no comprehensive metabolite profiling approach has been used to demonstrate alterations in large-scale metabolites. Thus, to overall unravel the effects of these metabolites on the metabolic network, HPLC-MS described above was employed to detect the metabolic pathway of some metabolites identified in mycelia and spores.

*B. bassiana* cells produce spores under conditions of nutrition deprivation. Interestingly, most metabolites (pyruvate, fumarate and ribulose) in the glycolytic, tricarboxylic acid (TCA) cycle and pentose phosphate pathways were markedly decreased in the stage of spores (Fig. 5). In particular, the level of fructose-1, 6-bisphosphate, a key factor in catabolite repression, dropped more rapidly in spores than in mycelia. It was possibly because that the decrease in fructose-1, 6-bisphosphate resulted in slow of metabolism, which
kept spores in a dormant, metabolically inactive conditions and help them survive longer.

The pentose phosphate pathway is a biochemical pathway parallel to glycolysis, which can generates NADPH and pentoses (5-carbon sugars). While it does involve in oxidation of glucose, its primary role is anabolic rather than catabolic [30, 31]. The pathway is a major source of reductant for biosynthetic processes such as fatty-acid synthesis and assimilation of inorganic nitrogen [32], and maintains redox potential necessary to protect against oxidative stress [33]. In the first stage of pentose phosphate pathway, glucose-6-phosphate was converted into ribose 5-phosphate, along with the energy production. And then ribose 5-phosphate was converted to phosphoribosyl pyrophosphate (PRPP). At the step, the metabolism was divided into three different pathways: histidine metabolism, purine metabolism and pyrimidine metabolism (Fig. 5). In the histidine metabolism, ergothioneine was extracted from B. bassiana as a metabolin, which is a naturally occurring amino acid and a thiourea derivative of histidine only produced by fungi and some prokaryotes. Ergothioneine has antioxidant properties in vitro [34]. Under laboratory conditions, it scavenges hydroxyl radicals and hypochlorous acid, inhibits production of oxidants by metal ions, and may participate in metal ion transport and regulation of metalloenzymes [35, 36]. In 2012, Bello identified an ergothioneine biosynthetic gene Egt-1 in Neurospora crassa and demonstrated that ergothioneine enhanced conidial survival and protected against peroxide toxicity during spore germination [37]. Thus, the high levels of ergothioneine in the spores play an antioxidant role, and keep active of spores.

Dimethyl guanosine is a product of purine metabolism. Charles and his co-workers have mentioned that dimethyl guanosine as a self-inhibitor serves to regulate spore development in Dictyostelium mucoroides [38]. Thus, the increased dimethyl guanosine in spores might diminish some gene transcription, which would make metabolism slower [39], and lead to improved shelf life of biological insecticide.

Dihydrothymine is a degradation product of thymine in the pyrimidine metabolism which is catalyzed by dihydropyrimidine dehydrogenase [40]. It associated with valine, leucine and isoleucine metabolism [41]. The decrease of dihydrothymine in spores might mean that the correlative metabolism network became slower than that in mycelia. A mannitol cycle was first proposed by Hult and Gatenbeck [42], which comprises four enzymes: mannitol 1-phosphate dehydrogenase (MPDH), mannitol 1-phosphate phosphatase (MPP), NADP+-mannitol 2-dehydrogenase (MDH), and hexokinase (HX), of which MPDH is the main synthetic enzyme and MDH is the main catabolic enzyme. The roles of mannitol vary in different fungi and might act as a scavenger of reactive oxygen species [43]. It was found only in spores (not found in mycelia) of B. bassiana, which implied that mannitol in spores might have an important impact on environment adaptability, germination, and virulence.
Glycerophospholipids, as glycerol-based phospholipids, are the main components of biological membranes, which play an important role in the generation of both extracellular and intracellular signals. As showed in Fig. 5, the increase of glycerophospholipids, esters and carnitine and the decrease of glyoxylate, pyruvate and acetylcarnitine in spores indicated that the hydrolysis of lipids and oxidation of fatty acids were depressed [44].

Beauverolides are insecticidal cyclodepsipeptides in *B. bassiana* [45]. The emergence of beauverolide Ka in mycelia (Table 1) may be related to the oxidative stress and/or oxylipin metabolism.

Sphingosine is a signaling lipid and may be involved in insect-fungi recognition, which can recognize the receptors on spores and induce the germination [46]. Sphinganine begins with the condensation of serine with palmitoyl-CoA to form 3-ketosphinganine, which is rapidly reduced to sphinganine [47]. Sphinganine is the biosynthetic precursor of sphingosines and sphingolipids, and its emergence in spores may affect fungal signaling and germination.
Fig. 4 MS spectrum of the eight compounds (a–d in mycelia extracts, e–h in spores extracts), a betaine, b carnitine, c glycerophosphocholine, d oxaloacetate, e dihydouracil, f succinic anhydride, g ergothioneine, h sphinganine
The TCA is a key metabolic pathway that unifies carbohydrate, fat and protein metabolism. The reactions of the cycle are carried out by eight enzymes that completely oxidize acetyl-CoA into two molecules of carbon dioxide. Through catabolism of sugars, fats and proteins, a two-carbon organic product acetate in the form of acetyl-CoA is produced which enters the citric acid cycle [48]. An increase of carnitine and decrease of acetylcarnitine, succinate, oxaloacetate and fumarate suggested that the TCA was depressed and lipid metabolism was enhanced in the spores. As a result, lipid was accumulated to protect spores against harmful effects of environment or others.

Conclusion

The proposed HPLC-MS methods enable the global determination of charged species, so that they can be used as universal tools for metabolome analysis. Metabolome data, along with macroscopic and microscopic techniques were used to provide important new information of the metabolism and growth of *B. bassiana* spores and mycelia. Gompertz model based on macroscopic and microscopic techniques was used to detect the relation between spores elongation length and culture time. Spore elongation length was found to increase exponentially until approximately 23 h after cultivation, and then growth became linear. The results of PCA displayed clear differences of the components in mycelia and spores of the fungus. Metabolic pathway of *B. bassiana* spores and mycelia was established based on HPLC-MS and KEGG database, which revealed the presence of twenty-eight major components in mycelia and thirty-six compounds in spores. In the metabolic network, the decrease of

**Table 1 Metabolites putatively identified by HPLC–MS in mycelia extracts**

| RT (min) | Detected mass | Metabolite | Ionization mode | Molecular formula | Theoretical mass | Δmass (mDa) | Relative concentration |
|---------|---------------|------------|-----------------|-------------------|-----------------|-------------|---------------------|
| 3.23    | 72.99348      | Glyoxylate | ESI (−)         | C₂H₂O₃            | 74.00039        | 1.460       | ↑                   |
| 64.92   | 87.00003      | Pyruvate   | ESI (−)         | C₃H₄O₃            | 88.01379        | −15.625     | ↑                   |
| 1.32    | 90.05431      | Alanine    | ESI (+) and ESI (−) | C₄H₆NO₂        | 89.0932         | −0.645      | ↑                   |
| 5.70    | 115.0154      | Fumarate   | ESI (−)         | C₆H₄O₄            | 116.07216       | 0.758       | ↑                   |
| 2.82    | 117.01956     | Succinate  | ESI (−)         | C₆H₄O₄            | 118.08804       | 1.415       | ↑                   |
| 1.45    | 118.08540     | Betaine    | ESI (+)         | C₆H₁₁NO₂          | 117.07704       | −0.855      | ↓                   |
| 1.17    | 129.13795     | Dihydrothymine | ESI (+)        | C₆H₁₄N₂O₂         | 128.12922       | 0.952       | +                   |
| 1.30    | 131.04636     | Oxaloacetate | ESI (−)        | C₆H₄O₅            | 132.07156       | −1.023      | ↑                   |
| 45.75   | 149.04712     | Ribulose   | ESI (−)         | C₆H₁₄O₅           | 150.55480       | 1.899       | ↑                   |
| 1.30    | 162.11154     | Carnitine  | ESI (+)         | C₆H₁₄NO₃          | 161.10318       | −0.930      | ↓                   |
| 1.45    | 204.12184     | Acetylcarnitine | ESI (+)       | C₆H₁₄NO₄          | 203.11348       | −1.195      | ↑                   |
| 24.58   | 230.11407     | Ergothioneine | ESI (+)      | C₆H₁₄N₂O₆         | 229.30031       | 0.569       | ↓                   |
| 41.61   | 255.23244     | Palmitic acid | ESI (−)       | C₁₈H₃₄O₂          | 256.24080       | 1.677       | +                   |
| 1.35    | 258.10883     | Glycerophosphocholine | ESI (+)   | C₆H₁₄NO₃P        | 257.10047       | −1.270      | +                   |
| 39.18   | 297.23239     | Linoleic acid | ESI (−)       | C₁₈H₃₄O₂          | 280.24075       | 0.533       | +                   |
| 29.99   | 295.2723     | 8-hydroxy-linoleic acid | ESI (−)    | C₁₈H₃₄O₂          | 296.23559       | −0.458      | +                   |
| 1.32    | 296.06451     | S-aminomimidazole ribonucleotide | ESI (+) | C₆H₁₄N₃O₆P | 295.0615       | 0.297       | +                   |
| 59.15   | 309.27618     | Linoleic acid ethyl ester | ESI (+)   | C₂₀H₃₆O₂          | 308.26782       | −0.080      | +                   |
| 28.48   | 312.30252     | Dimethyl guanosine | ESI (+)    | C₁₂H₁₄N₆O₇ | 311.29388      | 1.726       | ↓                   |
| 1.24    | 317.11334     | Mycosporin glutamicol | ESI (+)    | C₁₈H₂₃N₆O₇ | 316.10498      | 1.278       | ↑                   |
| 3.37    | 335.07434     | Fructose −1,6-bisphosphate | ESI (−)   | C₁₀H₁₄O₇P₂        | 336.08392       | 1.500       | ↑                   |
| 28.53   | 476.27719     | Phosphatidylethanolamine (18/2/0.0) | ESI (−) | C₂₀H₃₆NO₇P | 477.28555      | −0.024      | +                   |
| 30.57   | 478.29260     | Phosphatidylethanolamine (18/1.0) | ESI (−)   | C₂₀H₃₆NO₇P | 479.30096      | 0.095       | +                   |
| 35.26   | 522.35986     | Phosphotidycholine (18/1.0) | ESI (+)  | C₂₀H₃₆NO₇P        | 521.3515       | 0.154       | ↓                   |
| 36.15   | 524.29797     | Phosphatidyserine (18/1.0) | ESI (+)  | C₂₀H₃₆NO₇P        | 523.28961      | −0.082      | ↓                   |
| 36.80   | 595.28802     | Phosphatidylinositol (18/2.0) | ESI (−)   | C₂₀H₃₆O₁₃P        | 596.29638      | 0.230       | ↓                   |
| 40.74   | 597.30420     | Phosphatidylinositol (18/1.0) | ESI (−)   | C₂₀H₃₆O₁₂P        | 598.31256      | 0.760       | ↓                   |
| 16.52   | 631.38678     | Beauveriolide Ka | ESI (+)    | C₂₂H₃₈NO₃         | 630.378411      | −0.345      | +                   |

↑ Relative concentration of the component was higher in mycelia than in spores; ↓, Relative concentration was lower in mycelia than in spores; +, Relative concentration was only found in mycelia, and not found in spores.
Table 2 Metabolites putatively identified by HPLC–MS in spores extracts

| RT (min) | Detected mass | Metabolite               | Ionization mode | Molecular formula | Theoretical mass | Δmass (mDa) | Relative concentrations |
|---------|---------------|--------------------------|-----------------|-------------------|------------------|-------------|------------------------|
| 2.95    | 72.9902       | Glyoxylate               | ESI (−)         | C₂H₂O₃            | 74.0039          | −1.530      | ↓                      |
| 6.487   | 87.00903      | Pyruvate                 | ESI (−)         | C₃H₄O₃            | 88.01739         | −1.360      | ↓                      |
| 1.30    | 90.05455      | Alanine                  | ESI (+) and ESI (−) | C₃H₇NO₂ | 89.0932 | −0.405 | ↓                      |
| 2.95    | 99.00914      | Succinic anhydride       | ESI (−)         | C₄H₆O₃            | 100.01751        | 1.470       | +                      |
| 5.79    | 115.07640     | Fumarate                 | ESI (−)         | C₄H₈O₄            | 116.07216        | −1.627      | ↓                      |
| 1.17    | 115.12245     | Dihydropuracil           | ESI (+)         | C₅H₈N₂O₂          | 114.10264        | 0.361       | +                      |
| 2.98    | 117.0748      | Succinate                | ESI (−)         | C₆H₆O₄            | 118.08804        | −0.002      | ↓                      |
| 1.34    | 118.08559     | Betaine                  | ESI (+)         | C₆H₇NO₂          | 117.07704        | −0.024      | ↓                      |
| 45.76   | 131.06079     | Oxalacetate              | ESI (−)         | C₆H₈O₃            | 132.07156        | 1.486       | ↓                      |
| 3.32    | 128.03557     | Pyroglutamic acid        | ESI (−)         | C₆H₇NO₃           | 129.04393        | 1.350       | +                      |
| 1.05    | 129.13797     | Dihydrothymine           | ESI (+)         | C₆H₈N₂O₂          | 128.12922        | −0.085      | ↓                      |
| 2.27    | 146.04608     | Glutamic acid            | ESI (−)         | C₆H₈NO₄           | 147.05444        | 1.296       | +                      |
| 64.65   | 149.04669     | Ribulose                 | ESI (−)         | C₇H₈O₄            | 150.05505        | 1.240       | +                      |
| 1.28    | 162.1182      | Carnitine                | ESI (+)         | C₇H₇NO₃           | 161.10318        | −0.930      | ↑                      |
| 1.23    | 173.10434     | Arginine                 | ESI (−)         | C₇H₇N₂O₂          | 174.1127         | 1.038       | +                      |
| 1.34    | 181.07187     | Mannitol                 | ESI (−)         | C₇H₈O₄            | 182.08023        | 1.205       | +                      |
| 1.43    | 204.12219     | Acetylarnitine           | ESI (+)         | C₇H₈N₂O₄          | 203.11348        | −0.845      | ↓                      |
| 1.38    | 230.09485     | Ergothioneine            | ESI (+)         | C₇H₈N₂O₃S         | 229.30031        | 1.203       | ↑                      |
| 1.37    | 236.14837     | Ulvaline                 | ESI (+)         | C₈H₈N₃O₄           | 235.14003        | −0.859      | +                      |
| 33.64   | 299.25864     | 3-hydroxy-stearic acid   | ESI (−)         | C₈H₁₆O₃           | 300.267          | 0.569       | +                      |
| 27.63   | 302.30411     | Sphinganine              | ESI (+)         | C₈H₁₆N₂O₄          | 301.29575        | −1.246      | ↑                      |
| 2.38    | 312.12906     | Dimethyl guanosine       | ESI (+)         | C₉H₁₄N₂O₃          | 311.1207         | −0.025      | ↑                      |
| 3.32    | 315.11945     | Mycosporin glutamicol    | ESI (−)         | C₉H₁₄N₂O₄          | 316.12781        | 0.776       | ↓                      |
| 24.46   | 318.29892     | 2-amino-6-methyl-1,3,4-heptadecanetriol | ESI (+) | C₁₃H₂₀NO₃ | 317.29056 | −1.351 | + |
| 30.58   | 330.33508     | 2-amino-1,3-eicosanediol | ESI (+) | C₁₃H₂₂NO₄ | 329.32672 | −1.576 | + |
| 2.81    | 335.07434     | Fructose –1,6-bisphosphate | ESI (−) | C₁₃H₁₄O₄P₂ | 336.08392 | 3.0 | ↓ |
| 27.79   | 346.32990     | 2-amino-1,3,4-eicosanetriol | ESI (+) | C₁₃H₂₄NO₄ | 345.32154 | −1.671 | + |
| 41.75   | 433.23291     | Phosphatidic acid (18:2/0:0) | ESI (−) | C₁₃H₂₄O₃P | 434.24127 | 2.056 | + |
| 43.20   | 435.25119     | Phosphatidic acid (18:1/0:0) | ESI (−) | C₁₃H₂₄O₃P | 436.25955 | −2.939 | + |
| 31.79   | 520.33716     | Phosphotidylcholine (18:2/0:0) | ESI (+) | C₁₃H₂₄NO₄P | 519.32880 | 1.589 | + |
| 34.49   | 522.35303     | Phosphotidylcholine (18:1/0:0) | ESI (+) | C₁₃H₂₄NO₄P | 521.34467 | −2.386 | ↑ |
| 25.76   | 524.29108     | Phosphatidyserine (18:1/0:0) | ESI (+) | C₁₃H₂₄NO₄P | 523.28272 | 1.724 | ↑ |
| 45.33   | 538.31598     | Phosphatidyserine (19:0/0:0) | ESI (−) | C₁₃H₂₄NO₄P | 539.32434 | 2.501 | + |
| 16.13   | 566.34436     | Phosphatidyserine (21:0/0:0) | ESI (−) | C₁₃H₂₄NO₄P | 567.35272 | 2.515 | + |
| 11.48   | 595.28656     | Phosphatidylinositol (18:2/0:0) | ESI (−) | C₁₃H₂₄O₃P | 596.29492 | −1.230 | ↑ |
| 67.34   | 597.30200     | Phosphatidylinositol (18:1/0:0) | ESI (−) | C₁₃H₂₄O₃P | 598.31036 | −1.440 | ↑ |

1. Relative concentration of the component was higher in spores than in mycelia; 2. Relative concentration was lower in spores than in mycelia; +, Relative concentration was only found in spores, and not found in mycelia.

glyoxylate, pyruvate, fumarate, alanine, succinate, oxalacetate, dihydrothymine, ribulose, acetylcarnitine, fructose-1,6-bisphosphate, mycosporin glutamicol, and the increase of betaine, carnitine, ergothioneine, sphingosine, dimethylguanosine, glycerophospholipids, and in spores indicated that the change of the metabolin can keep spores in inactive conditions, protect spores against harmful effects and survive longer. The study provided the tools for understand and control the process of spores germination and outgrow to mycelia.
**Methods**

**Microorganisms and culture medium**

*B. bassiana* Bb0062 provided by the Anhui Provincial Key Laboratory of Microbial Control, Anhui Agricultural University, was stored at −20 °C in sterilized cryovials containing 10 % glycerol (in 0.02 % Tween 80 solution). It was cultured on potato dextrose agar (PDA) slants (potato 200 g/L, glucose 20 g/L and agar 20 g/L) at 26 °C for 7 days, and then stored at 4 °C until use.

**Preparation of inocula**

*B. bassiana* was cultured on PDA medium mentioned above at 26 °C for 7 days to obtain heavily sporulating cultures. Spores were then suspended in sterile distilled water containing 0.02 % (v/v) Tween 80 by gently scraping the agar surface with a sterile spatula, and then filtered through two layers of gauze to remove any debris (mostly mycelial fragments). The final spore concentration was adjusted to $1 \times 10^6$ spores/mL and used as quickly as possible.

**Assessment of germination and outgrowth**

Portions (100 μL) of the inoculum (*B. bassiana*), containing approximately $10^5$ spores, were surface plated aseptically on PDA. After inoculation, plates were sealed with parafilm to prevent moisture loss, and stored at
26 °C for 2 days. To monitor the kinetic behavior of the *B. bassiana* spores for prolonged periods (from germination to mycelium formation), computer morphometry (Leica Microscopy System Ltd, DMI 4000B, Germany) was employed to examine spore germination and mycelia outgrowth. Germination time was defined as the time at which the length of the germ tube was equal to the diameter of the swollen spore. Samples were measured every 2 h until 32 h. Images was analysed by Image-Pro Plus image analysis software version 6.3 (MediaCybernetics Inc., Bethesda, United States) and an auto-focus system.

For the germination study, length of germ tube over time was fitted to the modified Gompertz equation (Eq. (1)) [49] for the estimation of the germination kinetic parameters (ke and c):

\[
L = L_{\text{max}} \exp \left( -\exp \left[ \frac{ke}{L_{\text{max}}} (c-t) + 1 \right] \right)
\]

where *t* (h) is the time, *L* indicates length of spores elongation at time *t*, *L*_{max} represents length of hypha *t* \(\to\) \(+\infty\), ke signifies the slope of tangent line through the inflection point, *c* is the time when hyphal extention reach to maximum speed.

**Mycelia and spores preparation for HPLC-MS analysis**

Mycelia were harvested after 32 h of growth by a scoop, suspended in sterile water, and then filtered through two layers of gauze to remove debris, subsequently frozen in liquid nitrogen to terminate metabolism, and kept at −80 °C. Spores were collected at 7 days post-inoculation by scraping the colony into 0.01 % Tween 80 solution, and the contents were vortexed and filtered through a 10 μm microfiltration membrane to remove any debris. The filtrate was transferred to a 50 mL centrifuge tube and centrifuged for 10 min at 8000 rpm. The precipitate was resuspended in 2 mL of distilled water and transferred to a 5 mL centrifuge tube, centrifuged for 10 min at 10,000 rpm, and then frozen in liquid nitrogen to terminate metabolism. Finally, the spores were kept at −80 °C. Three independent biological replicates were measured per assay.

Mycelia and spores were lyophilized until a constant weight was attained. Mycelia were then crushed into a fine powder and then kept at 4 °C until extraction. Ten milligrams of sample was extracted with 2 mL of 80 % methyl alcohol (Tedia company, USA, HPLC grade), followed by 1 min of vortexing and subsequent sonication (12-KHz, 8-s exposure followed by a 4-s rest interval) for 1 h. Samples were further kept at 4 °C for 12 h in the dark. After centrifugation at 8000 rpm for 10 min, 1.8 mL of supernatant was collected and dried with a centrifugal concentrator. All samples were stored at −80 °C until analysis. The dried extracts were redissolved ultrasonically in 300 μL of 80 % methanol and filtered through a 0.22 μm polyvinylidene fluoride membrane filter before HPLC–MS analysis.

**HPLC – MS conditions**

In order to determine retention time and to obtain extract HPLC profiles, HPLC was performed on a Thermo-Fisher UPLC system (Thermo-Fisher, SanJose, CA, USA) coupled with an LTQ XL mass spectrometer. HPLC analyses were conducted on a C18 reversed-phase (RP) column (5 μm, 3 mm × 150 mm, 100A, Luna PFP Phenomenex, Torrance, CA, USA). The parameters were as follows: injection volume, 5 μL; column temperature, 40 °C; flow rate, 0.3 mL/min; and the eluates were monitored by full-length scan from 200 to 600 nm. The mobile phase was (A) 0.1 % formic acid (Anaqua Chemical Supply, USA, HPLC grade) in water and (B) 0.1 % formic acid in acetonitrile (Merck, Germany, HPLC grade), and gradient elution was carried out: 5 % B for 0–3 min, 5–100 % B for 3–50 min, and 100 % B for 50–60 min. The mass spectrometer parameter settings used for the measurement were as follows: ionization mode, for both positive and negative; gas temperature, 350 °C; drying gas, 12 L/min; nebulizer pressure, 45 psi; capillary voltage, 4000 V in positive mode and 3500 V in negative mode; fragmentor voltage, 215 V in positive mode and 170 V in negative mode; skimmer voltage, 60 V; and OCT 1 RF, 250 V. Data acquisition was performed in the m/z range of 50–1100 Da.

**Data processing and statistical analysis**

All data were processed using the Xcalibur software provided by the manufacturer. After all of the detected peaks were subjected to noise-reduction in both the HPLC and MS domains, the analytical peaks were processed by the software. A list of peak intensities with retention times and m/z data pairs was generated. The intensity of each peak was normalized by the sum of all of the peak intensities. Peaks with signal-to-noise (S/N) ratios lower than 5 were rejected. PCA were performed by SPSS v18.0 (IBM SPSS Statistics, Ontario, Canada) to envisage the different components of *B. bassiana* spores and mycelia.

**Metabolite identification**

The small-molecule inventory (SMI) or metabolome is a pattern of molecules that reflects the cell’s status. The molecular formula calculated by the Xcalibur software was predicted, and based on a general understanding of fungi metabolism pathways by searching web databases (Dictionary of Natural Products, METLIN, PUBCHEM, and CHEMSPIDER). The exact monoisotopic masses of possible metabolites were calculated based on their elemental compositions. Putative biomarkers were verified by its elution order (polarity) and structure characteristics. The ambiguous metabolites were identified by
comparison to authentic compounds available or referring to the published literature about fungi, especially entomopathogenic fungi.

Abbreviations
ESI: Electrospray ionization; HPLC-MS: High performance liquid chromatography-mass spectrometry; HK: Hexokinase; KEGG: Kyoto Encyclopedia of Genes and Genomes; MDH: Mannitol 2-dehydrogenase; MPDH: Mannitol 1-phosphate dehydrogenase; MPP: Mannitol 1-phosphate phosphatase; PCA: Principal component analysis; PEP: phosphopyruvate; PRPP: Phosphoribosyl pyrophosphate; SMI: Small-molecule inventory; TCA: Tricarboxylic acid.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
Conceived and designed the experiments: HXL. Performed the experiments: HXL, MXG. Analyzed the data: HXL, ZM2. All authors have read and approved the final manuscript.

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