Analysis of Quality and Techniques for Hybridization of Medicinal Fungus *Cordyceps sinensis* (Berk.) Sacc. (Ascomycetes)*

*John C. Holliday, 1 Phillip Cleaver, 1 Megan Loomis-Powers, 1 and Dinesh Patel 2*

1 Aloha Medicinals Inc., Haiku, Hawaii, USA; 2 Integrated Biomolecule Corporation, Tucson, Arizona, USA

Address all correspondence to John C. Holliday, Aloha Medicinals Inc., PO Box 686, Haiku, Hawaii 96778, USA; jchms@spry.net

**ABSTRACT:** In the course of our research with cultivated medicinal fungus *Cordyceps sinensis*, we have noted a greater diversity of compounds from different strains of this single species than in almost any other organism we have analyzed. Because of this great difference in chemical composition, a wide range of quality is found in cultivated *C. sinensis*. This article details the unique methods used to develop hybridized strains of *C. sinensis* using rattlesnake venom to trigger somatic fusion of dissimilar mycelial strains. This results in the reproducible production of cultivated *C. sinensis* containing quantities of recognized bioactive compounds equal to or greater than the quantities found in wild collected strains. These techniques and methods offer great promise in allowing cultivators of *C. sinensis* and other *Cordyceps* species to take their artificially cultivated products to a higher and more consistent level of quality and provide protocols for analysis of *C. sinensis* quality using HPLC and GC techniques.

**KEYWORDS:** *Cordyceps sinensis*, snake venom hybridization, HEAA, Cordycepin, adenosine, hydroxyethyladenosine, deoxynucleosides, *Cordyceps* culture parameters, *Cordyceps* analysis

**INTRODUCTION**

*Cordyceps sinensis* (wild Chinese Tochukaso) is an insect colonizing Ascomycetous fungus belonging to the family Clavicipitaceae (Kirk et al., 2001). *C. sinensis* is a parasite living on lepidopterous larvac. The normal range of this fungus is above 2000 meters’ elevation, and it has been found as high as 6000 meters. It occurs in wild grassy plains in the interior of China, as well as in Nepal, Tibet, and Himalayas

**ABBREVIATIONS**

GC/MS: Gas chromatography with mass spectrometry detection; HEAA: hydroxy ethyl adenosine analogs; HPLC: high performance liquid chromatography; TBE: tris-borate-EDTA; TCM: traditional Chinese medicine

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*From the Editor-in-Chief: *Cordyceps* (Fr.) Link (1833) nom. cons., as an Ascomycetous genus, is not closely allied to mushrooms. However, because of the positive importance of *Cordyceps* species to the medicinal profession, they are discussed here. Important data about the medicinal effects of *Cordyceps* species are presented in articles published in the IJMM (T. Mizuno, Medicinal effects and utilization of *Cordyceps* (Fr.) Link (Ascomycetes) and *Iaria* Fr. (Mitosporic fungi) Chinese Caterpillar Fungi, "Tochukaso" (Review), 1999, 3.253-254; T. H. Liao and H. C. Lo, Biological activity of *Cordyceps* (Fr.) Link species (Ascomycetes) derived from a natural source and from fermented mycelia on diabetes in STZ-induced rats, 2002, 4, 2: 111-126).
(Mizuno, 1999). There are also many other species of the genus *Cordyceps*, which all seem to have potent biologically active compounds present. The genus has been shown to produce some potent antibiotics, immune stimulants, and antiviral and antitumor agents (Hobbs, 1995; Zhou et al., 1998a,b; Mizuno, 1999; Hsu and Lo, 2002).

*Cordyceps sinensis*, which is known to the Chinese as “Dong Chong Xia Cao” and to the Japanese as “Tochukasoo,” has long been used in medicine. The first written record on the medicinal uses of this herbal medicine was in the Ben-Cao-Cong-Xin (New Compilation of Materia Medica) by the author Wu-Yiluo. Written around the year 1757 AD, this early medical text lists the traditional usage of *Cordyceps* as going to the lung and kidney meridian and being useful as a “Lung Protectorate,” for “Kidney Improvement,” and as a “Yin/Yang double invigorant.” *Cordyceps* in traditional Chinese medicine (TCM) was, and usually still is, prepared by cooking the whole caterpillar/fruitbody combination in chicken or duck soup. It has been used this way for the treatment of many conditions, such as respiratory diseases, renal dysfunction, hyperlipidemia, and hyperglycemia (Zhou et al., 1998a,b).

This may seem to be a great number of medicinal claims for a single substance. But the observations of many generations of medical practitioners have validated and confirmed this belief. People tend to be very good observers of nature, especially when the observations are proven over a long period of time, as has been the case with *Cordyceps*; there is sufficient background for further research into this herbal substance for use in modern medicine.

Since the 1950s attempts have been made to cultivate *Cordyceps sinensis* and other *Cordyceps* species, and many of these artificially cultivated *Cordyceps* have been available in the international marketplace since the 1980s. In part as a result of this wide availability of *Cordyceps*, the worldwide demand for this previously rare health supplement has greatly increased in the last 20 years. This has led to over-harvesting of the wild stocks and a subsequent shortage of natural *Cordyceps*. With the increasing scarcity of *Cordyceps* in the wild (Chen et al., 2000), the prices for this popular health supplement are ever increasing, which has led to more and more companies producing artificially cultivated *Cordyceps* for the nutraceutical and pharmaceutical markets. This increase in demand and supply has lead to a wide variation in the purity and quality of *Cordyceps* products available. It has also lead to a large number of counterfeit and adulterated products being sold under the name *Cordyceps*.

There have not been any widely accepted standards available on which to base the analysis of *Cordyceps* for quality purposes. Some manufacturers analyze for Adenosine, while others analyze for Cordycepin, while yet others analyze for polysaccharides or other compounds. This has lead to great confusion in the public's mind as to which *Cordyceps* is best for them to buy. In the course of our research into this quality issue, the first thing we had to do was to develop analytical protocols and choose prototypical compounds to test for, in order to establish a quality index by which different strains and commercial products could be graded. These protocols and compound descriptions are found in the section Materials and Methods.

There is debate among many scientists at present whether the species of the genus *Cordyceps* are in fact single organisms or if they are symbiotic colonies of more than one organism. Perhaps what we are calling *Cordyceps sinensis* today will one day be known as a fungal/bacterial symbiosis. DNA sequencing has proven inconclusive in this regard, because the DNA sequence tends to change with time, as if the fungus were incorporating some of the insect DNA into its own DNA code for the initiation of its fruitbody form, then losing the insect DNA when it goes back into its mycelial form.

Microscopic examination of growing *Cordyceps* mycelium often reveals some very interesting morphology, including the concurrent anamorphs of filamentous mycelium and rapidly moving single-celled yeast-like morphological forms. This has been seen in other *Cordyceps* species as well, such as *Cordyceps sobolifera* (Hill.) Berk. et Br. We have observed what appears to be inner-cellular symbionts—small, spherical, rapidly moving, apparently single-celled organisms living within the long, bamboo-like hyphal strands. When we observe this growth form, these secondary inner cellular objects occur in about 70–80% of the mycelial cells observed.
with anywhere from one to five individually moving spheres per mycelial cell. These are easily observed in the living culture under 1000-power magnification with normal back lighting.

MATERIALS AND METHODS

Modern Cultivation of Cordyceps sinensis

Because native Cordyceps sinensis is quite rare and very expensive, there has been a lot of research into methods for cultivation of this fungus. The strain presently known as CS-4 was one of the first commercial strains of C. sinensis isolated and brought into cultivation. Known by the name of Paecilomyces hepialii Chen, the aseptically fermented mycelium of this strain underwent extensive human testing and clinical trials during the 1980s and resulted in a commercial product with wide usage in China, known as Jin-Shui-Bao capsules. More than 2000 patients were involved in the clinical trials with CS-4 during the pharmaceuticalization of Jin-Shui-Bao, and the chemical composition, therapeutic activity, and toxicity are very well known for this strain. (Bau, 1995).

A number of other strains have been isolated from wild Cordyceps since then that are presently under commercial cultivation around the world. These strains are each so different from the original Cordyceps and from each other that they have been given many different genus and species names. Even though the parent fungus is the same in each case, the resultant asexual mycelial growth forms are characteristically different enough in morphology and chemistry to be considered different species by many taxonomists. Table 1 lists some of the different strains isolated from wild C. sinensis (Yin and Tang 1995; Zhao et al., 1999).

| TABLE 1. Characteristics of Some Commercialized Strains of Cordyceps sinensis |
|---|---|---|
| Latin binomial | Isolated by | Commercial product |
| Cephalosporium sinensis | Qing Hai Institute of Livestock and Veterinary sciences | NingXinBao |
| Paecilomyces sinensis Cn80-2 | FujianQingLiu County Hospital | various |
| Scyphialium sp. | Sanming Mycological Institute | SM1H8819 |
| Scyphialium sp. | Chinese Navy Institute of Medicine | 832 |
| Hirsutella sinensis | | various |
| Mortierella hepialii Chen Lu | | various |
| Topycladium sinensis | | various |
| Scyphalidium hepialii G.L.Li | | various |

Adulterated and Counterfeit Cordyceps sinensis

With the opening of China to business with the West during the 1970s, many people in countries far from China were exposed to the benefits found in TCM. Along with this exposure to traditional medical methods came a great demand for the various herbal compounds used in that medical system. This great worldwide demand for C. sinensis, and the enormous cost of the wild collected variety has led to many unscrupulous manufacturers and distributors providing adulterated and counterfeit C. sinensis in the world market (Hsu et al., 2002). Most of the Western world prefers their medicine to come in clean white bottles and neat capsules, rather than in the whole caterpillar form. This makes it even easier and more tempting for some suppliers to sell just about anything under the label of Cordyceps. Most consumers of natural Cordyceps already know that it is normal practice for collectors to insert small
segments of twigs or even pieces of wire into the body of the caterpillars to increase the weight (Wu et al., 1996). And most consumers of capsulated Cordyceps in the western countries do not even know what real Cordyceps tastes or smells like. We have analyzed some specimens of "Cordyceps" capsules which contained nothing but rice flour, and other samples which contained nothing but nutmeg.

In an attempt to establish a reliable method for identifying "real" Cordyceps, we started analyzing all of the available cultivated Cordyceps products, both consumer packaged products and bulk raw material products, grown by nearly all of the commercial cultivators and suppliers worldwide. What we found was shocking. Nearly all of the commercially available Cordyceps products available in the United States (nearly all of which were imported from China) contained no detectable amounts of Cordyceps whatsoever. The results of testing on American-produced Cordyceps were a little better. With American Cordyceps, in every case we were able to recognize the characteristic analytical signature of Cordyceps, (refer to Fig. 1 to see the characteristic analytical signature used in proving specimens as being Cordyceps or not), but in none of the American samples was there any significant amount of active

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**FIGURE 1.** Wild and hybrid Cordyceps total HEAA content in this sample of wild Cordyceps is about .091% w/w. The HEAA content of the hybrid Cordyceps is .323% w/w. NOTE: This plot of the wild Cordyceps is the basic signature of Cordyceps sinensis that we use for comparative purposes when determining whether a sample is actually Cordyceps or not. An unknown sample is considered as "authentic" Cordyceps sinensis if the plot of the unknown sample matches most points (>90%) on this natural Cordyceps plot when one is overlaid on the other.
ingredients. The American-grown Cordyceps products consisted almost entirely of unconverted grain substrate upon which the Cordyceps is grown.

Test Methods and Standards of Quality

The methods for analyzing Cordyceps species quality have not yet become standardized throughout the world. Every lab that is conducting this type of testing uses its own methods and standards. So when we first began our analysis of Cordyceps sinensis in 1999, we had to develop our own test methods. We tried many different test protocols before settling on the following two as being accurate, repeatable, and relatively economical.

Gas Chromatography with Mass Spectrometry Detection

Trimethylsilyl Derivative Method. Starting with well dried and finely ground powder of the raw test sample, 20 mg is added to 0.3 mL of derivatizing agent (BSTFA) and 0.3 mL of acetonitrile. This mixture is heated for 20 minutes at 60°C, which yields a Trimethylsilyl derivative, which carries the material through gas chromatography (GC) for detection with a mass spectrometry (MS) detector. This test method is simple and quick, and it yields a positive/negative answer as to whether the test sample is actually C. sinensis or not. This method can also be used for quantification of the target compounds, although the next method is more accurate and more suitable for complete target compound quantification.

High Performance Liquid Chromatography/ Mass Spectrometry Detection

Powdered samples (2.0 g) were defatted by decanting with hexane (3 × 50 mL) and dried in vacuum. Samples were dissolved in 0.1M TBE (Tris-borate-EDTA) buffer (pH 8.5 with 0.1N NH₄OH) (100 mL) and sonicated for 30 minutes at 40°C. An aliquot (10 mL) of the sample was then passed through a C-18 Sep-Pak that had previously been pre-equilibrated with 0.1 M TBE buffer (pH 8.5 with 0.1 N NH₄OH). The eluent was collected and the Sep-Pak further washed with the equilibration buffer to give a final eluent volume of 20 mL. After thorough, mixing the sample was filtered through a 0.45 micron PVDF membrane and placed into suitable vials for HPLC-MS analysis. The chromatography was performed on a Waters 2695 separation module using a Wako Wakosil-II 5C18 HG column (5 μm, 15 cm × 4.6 mm i.d.) at 45°C with gradient elution of H₂O:methanol (1 mL/min) from 22:3 to 77:23 in 19 minutes, then to 18:7 at 24 minutes and 27:23 at 39 minutes. The chromatographic eluent was passed into a Vestec particle-beam interface for solvent removal and particle atomization and then via Teflon transfer line into the mass spectrometer using a helium carrier gas. Detection was performed on a Finnigan TSQ7000 triple-quadrupole mass-spectrometer in positive ion mode with full scan centroid data collection (50–1000 m/z). MS/MS experiments using an argon collision gas were used to verify the identity of unusual nucleotides for which no primary standards were available.

Standards of Quality, or "Quality index"

Quality determination in Cordyceps sinensis for the health supplement industry has been haphazard up until now, because there have been no universally recognized test methods for analyzing this particular supplement. Each company producing or supplying Cordyceps has used different tests or tested for different substances in order to show that its product stands above the rest. Some analyze for adenosine, some for Cordycepin, some for cordycepic acid, and some for particular sugars or total polysaccharides. While all of these tests have some usefulness in determining relative quality, none by itself is in any way meaningful when it comes to whether the product in question will yield good results when used as medicine for humans.

Almost all of the samples of wild collected Cordyceps analyzed very similarly in chemical composition, but we quickly realized that there was a tremendous variation—greater than we had found
in any other organism—in the secondary metabolite compounds present in cultivated *Cordyceps sinensis* and other *Cordyceps* species. So our first step was to determine which compounds we could use as targets to equate with medicinal potency. After a thorough review of the literature, the nucleosides, and specifically the deoxy-nucleosides, were determined to be the most reliable indicators of potency. This class of compounds has been carefully studied and reported in the scientific literature and was the class of compounds showing the most variation in different samples of cultivated *Cordyceps*. Furthermore, many of the deoxy-nucleosides are found in no other organism, or at best, a very limited number. We chose the compound N<sup>6</sup>-(2-hydroxyethyl)-Adenosine as our indicator compound, because we found this in all specimens of *Cordyceps* tested but not in any other organism. This compound, along with Adenosine and 3’-deoxyadenosine (Cordycepin) were used in summation as the quality indicator for comparison of different strains and production methods of *Cordyceps*. In other words, the quantities of the three compounds (in percentage w/w) were added together to come up with a numerical quality index for *Cordyceps* (Furuya et al., 1983). Structures for these three compounds are shown in Table 2.

There are a number of other interesting deoxy-nucleosides produced by *Cordyceps sinensis*, such as the compound 2’,3’-deoxyadenosine, which is marketed in the United States as a drug for the treatment of AIDS under the trade name Didanosine. There are also several varieties of deoxy-uridines present, but there is not much literature on these compounds yet, so we decided not to use these as quality index markers. In time we will learn what effect these unique compounds have in the human body and may find that these compounds are as important a group as the adenosine class of compounds, which we refer to in this paper as HEAA (hydroxy ethyl adenosine analogs).

We find that the HEAA content is a much more reliable indicator of *Cordyceps* species quality than could be determined through testing of the polysaccharides. The reason for this has to do with the methods of analysis for polysaccharides, which is usually done by wet-chemistry methods—breaking the polysaccharide bonds through acid hydrolysis or enzymatic activity and then measuring the quantity of simple sugars present after cleavage. We find this method unreliable as a quality indicator, because the residual sugars tell nothing of their source or their linkage characteristics (i.e., alpha-bonded or beta-bonded), and the test results are easily altered through the addition of starch or other polysaccharides to the raw material at the manufacturing stage. Furthermore, the resultant simple sugars are not unique to the *Cordyceps* in question. Rather, by using the HPLC/MS or GC/MS method and testing for relatively rare nucleosides, any sample of *Cordyceps* from any source is quickly and easily analyzed and a numerical quantity index can be applied to it. When this method becomes universally applied in the health supplement industry, producers will be forced to address the quality issue of their

Table 2. Hydroxy Ethyl Adenosine Analog Structures

<table>
<thead>
<tr>
<th>Structure</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1.png" alt="Structure 1" /></td>
<td>N&lt;sup&gt;6&lt;/sup&gt;-(2-hydroxyethyl)-Adenosine</td>
</tr>
<tr>
<td><img src="image2.png" alt="Structure 2" /></td>
<td>Cordycepin (3’-deoxyadenosine)</td>
</tr>
<tr>
<td><img src="image3.png" alt="Structure 3" /></td>
<td>Adenosine</td>
</tr>
</tbody>
</table>
products, and the industry as a whole will benefit. There will be much less adulterated or counterfeit product placed on the market and consumer confidence will grow as a result. It is hoped that by the application of the test methods given here, such deceptive production practices will stop.

**Culture Methods and Substrates**

In looking at the variations in quality from different strains and producers of Cordyceps, one must wonder what causes this to be so. After all, a tomato is a tomato, no matter where it is grown. Yet with Cordyceps sinensis, even the same strain (CS-4) grown by different cultivators turns out to be entirely different from a standpoint of active ingredients.

In looking into this question, it is first important to realize that there are two different methods used today in the cultivation of Cordyceps. There is the method primarily used in China, known as liquid culture or fermentation, in which the organism is introduced into a tank of sterilized liquid medium, which has been formulated to provide all of the necessary nutritional components for rapid growth of the mycelium. After growth in the liquid medium, the mycelium is harvested by straining it out of the liquid broth and dried, after which it can be used as is or further processed. Generally in this method the extracellular compounds, which were exuded by the fungus during the growth cycle, are discarded with the spent broth. This represents a major loss of bioactive compounds, because many of the active ingredients are extracellular in nature and are found only in small concentrations in the mycelium.

The second cultivation method is the solid/substrate method followed by most growers in Japan and America. In this cultivation system the mycelium is grown in plastic bags or glass jars full of sterilized medium, which is almost always some type of cereal grain. This grain is usually rice, wheat, or rye, although many different types of grain have been used. After some period of growth, the mycelium is harvested along with the residual grain. While this is an easily mastered and low-capital-investment cultivation technique, the negative side of this method is that the grain content is usually greater than the mycelium content. In many cases the solid/substrate-grown mycelium we tested contained greater than 80% residual grain. However, a bonus to this method is that the extracellular compounds are harvested along with the substrate and mycelium.

Cordycepin is an example of one of the compounds that are primarily extracellular in nature. Many tests have been done on cultured Cordyceps mycelium for the presence of Cordycepin, and these show that Cordycepin is generally present in solid/substrate grown Cordyceps but not in liquid-cultured Cordyceps. The presence or absence of Cordycepin is dependent upon, among other factors, by which method the mycelium was grown and harvested.

We can see from this that the culture method itself has an effect on the quality of the resultant Cordyceps product. Beyond the methodology itself, the next most important factor in the production of particular secondary metabolites (or target medicinal compounds) is the nature and composition of the substrate itself (Zhang et al., 1992). While it would seem that a substrate that favors rapid and strong growth of the mycelium would be an ideal substrate to use, this is not necessarily the case. Substrates are chosen on availability and price or on historical usage or preference in handling. But rarely have they been chosen on the basis of the end compounds produced. In fact, the only way to determine whether the substrate being used is the best choice is to compare the resultant product after growth on that particular substrate with some standard. If the end goal of production is Cordycepin or didioxyadenosine (or some other specific compound)—as it is with some of the pharmaceutical companies—then the analysis is fairly straightforward. Just look for the amount of Cordycepin or didioxyadenosine present and adjust substrate composition accordingly. However, business in the health supplement industry is rarely so simple. First we have to assume that we know what we are looking for. Because natural products such as Cordyceps are chemically very complex, the truth is that we do not really know all of the components that are bioactively important.

With this realization in mind, we set out on a mission to produce the best Cordyceps possible. What is the best? Because we did not know the answer to
that question, we decided to try to copy the natural, wild collected *Cordyceps* as closely as possible. We attempted this by altering the substrate composition and analyzing the resultant mycelial product for known bioactive compounds. Then we altered the substrate repeatedly. We did this through several hundred different substrates and through many thousands of kilograms of resultant product. What we found was that there was not any single method, strain, or substrate we could use that would yield the results desired.

**Substrate**

The substrate of choice for most Chinese growers is a liquid medium based upon silkworm residue, with added sugar, carbohydrates, and minerals. This seems a logical choice, because this mushroom is found in nature growing on insects. Dried silkworm bodies are the byproduct of an existing industry and have little other use. They are therefore readily available and cheap. This silkworm-based substrate seems to yield a relatively high-quality product. The only problem with silkworm-residue-based substrate is that in the United States, the FDA requirements are for mycelial products to be produced on a normally consumed human food source. Silkworms do not fit into that category. They are also not available as a raw material source to most of the worlds *Cordyceps* cultivators.

The most usual substrate for Japanese and American growers is rice. It was determined in our trials that rice is not a suitable substrate for *Cordyceps* production if the target medicinal compounds are considered. Rice does not allow the full range of secondary metabolites to be expressed by the fungus, and rice grown *Cordyceps* has tested inferior in all of our analyses of active ingredient. There is rarely any appreciable amount of adenosine or Cordycepin present in rice-grown *Cordyceps*. Furthermore, there are growth-stunting metabolites that build up in the substrate when *Cordyceps* is grown on rice, limiting the growth stage to only about 22–24 days and allowing no more than about 40% of the rice to be converted into mycelial mass. This figure of 40% represents the high end of conversion, and is usually around 25–30%. This means that when *Cordyceps* is grown on rice then dried and powdered, the resultant product is actually between 60–75% unconverted rice flour.

Rye grain is another substrate often used for solid culture, and it yields a higher quality product than rice, as long as some vegetable oil as an amendment is added to the growth medium at the time of substrate makeup. The oil provides necessary nutrients, which the organism uses for bioactive compound production. Rye has other disadvantages, however. The compounds in rye, which give it that characteristic rye smell and taste, are not broken down by the *Cordyceps* and concentrate in the final product. This rye taste and smell overrides the characteristic *Cordyceps* taste and smell, and even though the resultant product is of better quality than the rice-grown mycelium (as determined through analysis), there are certain perceptual problems that must be overcome by the buyer to make this an economical substrate alternative. Rice-grown *Cordyceps* may seem like a better product to the average buyer because the rice does not mask the characteristic *Cordyceps* smell and taste. Most buyers in the health supplements industry tend to purchase bulk products on perception and faith rather than requiring an independent analysis. Rye also has growth-limiting factors, which cause *Cordyceps* growth to stunt at about 28–30 days, although this can be overcome to a slight degree with the addition of about 1% ground oyster shell buffer to the medium at time of make-up. We tried many other sources of calcium, but none seemed to work as well as the oyster shell calcium.

Millet is a very good choice of substrate when it is available. It has no strong taste or smell, it does not stunt the growth to any significant degree, and it allows for the full expression of the secondary metabolites by the organism. It has another problem though, which is the high ratio of chitinous outer husk layer to starch. This outer husk is not broken down and represents a large portion of the final product weight, about 15%. The chitinous husk cannot be removed from the grain ahead of time, because doing so causes the grain to become too sticky during sterilization, and a high degree of anaerobic contamination follows. The husk can be removed from the final product through mechanical
means such as a time-of-flight separation process, or the product can be used for hot water extractions or other processing. *Cordyceps* does not grow as quickly on millet as it does on other grains, but the end product quality is higher.

White milo grain, also known as white kaffir corn or white sorghum, is an excellent choice of substrate. The red variety of milo does not work nearly as well as the white variety as a substrate. White milo has all of the best characteristics—it is cheap, it has a high starch to husk ratio, it does not stunt the growth, it allows the full expression of bioactive compounds to be expressed, and it has no strong odor or taste of its own to compete with the taste and smell of the resultant *Cordyceps* product. White milo, when used alone, however, lacks some essential ingredients required for optimum growth by the *Cordyceps*. The addition of some portion of millet to the white milo speeds up the growth by a factor of 6 times. The millet to milo ratio is optimum at 1 part millet to 4 parts white milo.

Many farmers grow both white and red milo in the same fields, or store them in the same silos, or otherwise do not keep the white and red separated. This is to be avoided when using the milo as a *Cordyceps* substrate, because a small proportion of the red mixed in with the white can drastically reduce the growth rate and overall quality of the final product.

So from our substrate testing it was determined that the ideal medium for solid substrate growth of *Cordyceps* is as follows: 1 part white proso millet (husk on) to 4 parts of white milo (husk on), with the addition of 0.8% w/w of ground oyster shell and 1% w/w vegetable oil (peanut oil or soybean oil). Add water to equal 50% total moisture in the sterilized substrate. Precooking the grain mixture for 4 to 6 hours prior to sterilization tends to trigger a much faster growth response from the *Cordyceps*. On this medium, *Cordyceps* can be grown for long periods of time, allowing nearly complete conversion of the substrate to mycelium (96%-+) and the full expression of secondary metabolites from the *Cordyceps*. The resultant *Cordyceps* grown on this substrate is about 3-4% residual grain, or about 96-97% pure mycelium. The real benefit to this method of growing is the capture of the entire compliment of extracellular metabolites produced throughout the entire growth process. With the addition of certain growth-triggering compounds to this mixture, *Cordyceps sinensis* is easily induced to fruit in culture without any insect material being present. However, the formation of the fruitbody on this medium does not result in any significant change to the analytical chemistry profile. See Table 3 for substrate summary.

**Culture Parameter Modification:**

**Low Temperature Hypoxia**

Using the above-described substrate, the complete chemical profile of the cultivated *Cordyceps* still will not approach that of the wild collected *Cordyceps* unless it is grown under very specific conditions. *C. sinensis* produces a relatively large amount of free adenosine when grown at normal atmospheric oxygen levels and room temperatures. It will also produce a large quantity of uridine and guanidine. But there is very little if any Cordycepin produced, and virtually no hydroxyethyl adenosine. For the organism to produce these compounds, it needs to be growth stressed through the absence of oxygen, a drop in temperature, and the total absence of light. Just growing it under cold and anaerobic conditions from the start will not do the trick, because when *C. sinensis* is grown under those conditions, it forms a yeast-like anamorph that has a very different chemical profile. It must first be grown hot and fast, then tricked into converting its "summertime" metabolites into the target medicinal compounds we are looking for. To get these target compounds, we found that we needed to follow a strict growth protocol: After inoculation on to the millet/milo substrate, the *C. sinensis* is grown at 20-22 °C, in diffuse light and at sea level atmospheric oxygen for 28-30 days. It is then moved into a specially controlled environmental chamber, where the oxygen content is dropped to 50% atmospheric. The remainder of the growth atmosphere is made up of nitrogen, carbon monoxide, and carbon dioxide. The temperature is dropped to 3 °C, and all light is excluded. It is held under these conditions for 15-20 weeks. This results in much of the adenosine being converted to Cordycepin, di-
TABLE 3. Comparison of Different Substrate Compositions on Cordyceps Quality as Detailed in Text

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Speed of growth</th>
<th>% Residual substrate in finished product</th>
<th>Substrate quality index(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silkworm / Sugar (Chinese)</td>
<td>Fast</td>
<td>15–20%</td>
<td>Not FDA allowable(^a)</td>
</tr>
<tr>
<td>Millet</td>
<td>Medium-fast</td>
<td>15%</td>
<td>90</td>
</tr>
<tr>
<td>White milo</td>
<td>Medium</td>
<td>2–5%</td>
<td>90</td>
</tr>
<tr>
<td>Millet &amp; white milo 1:4 ratio</td>
<td>Fast</td>
<td>2–5%</td>
<td>100</td>
</tr>
<tr>
<td>Rye</td>
<td>Medium-slow</td>
<td>40–50%</td>
<td>60</td>
</tr>
<tr>
<td>Oats</td>
<td>Medium</td>
<td>20–30%</td>
<td>80</td>
</tr>
<tr>
<td>Wheat</td>
<td>Medium-slow</td>
<td>50–80%</td>
<td>60</td>
</tr>
<tr>
<td>Brown rice</td>
<td>Medium-fast</td>
<td>60–80%</td>
<td>60</td>
</tr>
<tr>
<td>Soy beans</td>
<td>No to slow</td>
<td>90–99%</td>
<td>Not a good substrate(^c)</td>
</tr>
</tbody>
</table>

\(^a\) FDA Regulations cpg 585.525

\(^b\) This substrate quality index is based upon the analytical profile of *Cordyceps sinensis* grown on each substrate, and to a lesser degree it is subjective and is based upon the authors’ experience of the growth characteristics on each, such as speed of growth, tendency to contaminate, ease of sterilization, and ease of handling. The best substrate found was the millet/milo mix, which was assigned a Quality index value of 100. All the other substrates are judged in relationship to this.

\(^c\) Soy beans are not an acceptable substrate in the authors’ experience. We tried to grow a number of different strains of *Cordyceps sinensis*, *C. militaris* (L.:Fr. Link, *C. apoiiognoioides* (Ehrh.) Fr., and *C. sobolifera* on different soy based compositions but had virtually no success. We are aware that there are commercial products on the market that advertise being grown on soy, however we can only guess what tricks those cultivators have developed to insure good growth on this medium. We mention soy-based substrates here only as a guide for the beginning cultivator, so they do not waste too much time trying to follow a route that we have found unacceptable.

deoxyadenosine, and hydroxyethyl-adenosine. Many other unique nucleosides are also produced, with a final chemical profile identically matching that of the wild Cordyceps, as can be clearly seen in Figure 1.

Hybridization

Once we had developed the substrate and growth parameters to optimize the target compounds, we started looking into the chemical profile differences from different strains of *C. sinensis*. Because there were so many strains of Cordyceps, and each strain has its own unique chemical profile, we tested all of the strains we were able to obtain. None of the known strains was shown to produce nearly the quantities of active ingredients found in the wild Cordyceps. So we started experimenting with ways to quantitatively increase the target compound production through the hybridization of Cordyceps strains—to crossbreed them in order to gain greater production of target compounds. This was quite a challenge. Because spore collection and separation is very time-consuming and results in entirely too many unknown variations, we felt this method would take too much time before we had reliable results.

Rather we took a novel approach. We experimented with various ways to get different strains of the fungi to perform their own nuclear fusion. There are several chemicals known to trigger this exchange of genetic material between unlike cells. Nicotinic acid, for instance, can be used to create hybridized mycelium. This compound is difficult to use and yields unreliable results. After trying several different compounds to trigger this fusion, we settled on snake venom.
Snake Venom as Hybridization Agent

We used purified snake venom from the Western diamondback rattlesnake (Crotalus atrox) (Sigma Scientific, St Louis, Missouri) for our hybridization techniques. The snake venom is added to the agar medium in quantities that alter the growth but do not prove toxic to the strain in question. This range of snake venom is from 10 to 30 mg per 300 mL of agar medium. The venom is not heat stable and must be added aseptically after sterilization of the medium. The agar used for this hybridization is an Aloha Medicinals' proprietary agar named R7 Agar, consisting of malt extract, activated carbon, minerals, and humus—the carbon-rich ash residue from a coal-burning industrial process. For the exact recipe see Table 4. Other agars could probably be used as well. This just happens to be our production agar that we use everyday, and once we found that it also worked with the snake venom for hybridization, we found no reason to experiment with any other agar.

Hybridization Technique

Petri dishes of this R7 agar medium are inoculated with mycelium from two strains of the Cordyceps genus. These are usually two varieties of C. sinensis, although we have also crossbred C. sinensis with other Cordyceps species such as C. militaris, C. sobolifera, and C. ophioglossoides. These different strains when inoculated together onto one petri dish will normally grow toward each other until they almost meet, at which point they form a zone of inhibition, where neither strain can grow. Eventually, one strain may prove stronger than the other and overgrow the plate, but they will remain genetically distinct: two different cultures residing in the same Petri dish.

With the addition of a sufficient quantity of snake venom to the agar, we found that the two cultures grow toward each other until they meet and form their mutual zone of inhibition. This period of inhibition is short-lived however, for in only about 2 or 3 hours the colonies each start sending out mycelial strands into this no man’s land, the zone of inhibition. These strands grow together and exchange nuclear material through their venom-weakened cell walls. They form a hybrid strain at this point of mutual contact—a new strain, one that is distinctly different from either of the parent strains. Within about 4 hours after first forming the zone of inhibition, the hybridization is complete and the colonies resume rapid growth toward each other. They become three colonies, rather than the original two. Then there exist in the same plate the original two colonies and a genetically distinct third—the hybrid.

A section of the newly formed hybrid is carefully removed from the original zone of inhibition at the precise time that the colonies begin to fuse. That is during hour 3–4 after the initial meeting of the colonies. The hybrid is transferred to a new petri dish containing normal (non-snake venom) agar. Our quick method of determining hybridization is to inoculate a new dish containing normal agar with tissue samples from all three strains, the original two and the suspected hybrid (see Fig. 2). If hybridization has failed to occur, then the suspected hybrid will readily fuse with either one or the other of the original colonies, proving that our suspected hybrid is not genetically distinct from the original, and we start again. But if the hybridization has in fact taken place, these are now three distinct colonies that will form a mutual three-way zone of inhibition. These can be clearly seen in Figure 3, where there are distinct zones of inhibition shown between three different colonies in the plate.

Once a hybrid is confirmed, it is tested for growth parameters. If it appears to be a vigorous and hard grower on our substrate of choice, we grow out a quantity of mycelium, harvest it, and analyze it for active ingredients. Through repeated testing in this

<table>
<thead>
<tr>
<th>TABLE 4. Snake Venom / R7 Agar Recipe</th>
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</thead>
<tbody>
<tr>
<td>2.1 L</td>
</tr>
<tr>
<td>50 g</td>
</tr>
<tr>
<td>34 g</td>
</tr>
<tr>
<td>10 g</td>
</tr>
<tr>
<td>5 g</td>
</tr>
<tr>
<td>1 g</td>
</tr>
<tr>
<td>10 mL</td>
</tr>
<tr>
<td>As required</td>
</tr>
</tbody>
</table>
FIGURE 2. Snake venom hybridization. Agar shown is proprietary R7 agar with .0001% Crotalus atrox venom added. Cultures are (A) Cordyceps sinensis CX4 and (B) Cordyceps sinensis SC1007 strains.

The way we were able to create the hybrid strain shown in Figure 1, which is easily grown in solid substrate culture, with a potency greater than any other cultivated strain and at least equal to the highest quality wild Cordyceps. We are referring to this new strain as Cordyceps sinensis Alohaensis. We are presently continuing this hybridization work with other species of Cordyceps, for the production of very specific target compounds. Top-quality Cordyceps is no longer a health supplement only for the very rich. By using these new methods of cultivation, the best quality Cordyceps is now within economic reach of even the common man throughout the world.

RESULTS

Analytical Results

We conducted analysis over the course of 4 years on samples from what we believe to be every commercial producer of cultivated Cordyceps. The results obtained through testing these approximately 100 samples from producers in four countries are shown in Table 5.

It can easily be seen from this table that there is quite a difference in the quality of Cordyceps from different producers. These quality differences become even more pronounced when seen in graphic format. Figure 1 shows the HPLC analytical profile of the hybrid C. sinensis strain overlaid on a plot of wild C. sinensis to show the comparison in qualitative similarities as well as the quantity of active ingredients in the cultivated hybrid strain versus the wild strain of Cordyceps. The secondary metabolites produced are very nearly identical in these two specimens, as revealed by these overlaid analytical signature.

We further showed through HPLC and GC analysis that Cordyceps is by no means consistent in its profile of secondary metabolites, but rather the analytical profile varies greatly, according to the combination of all the differing parameters laid out in this article. While different strains seem to have the greatest influence in exactly which secondary metabolites are produced, all factors involved in the entire cultivation process come into play in this regard. Even subtle changes in light cycle can drastically influence the resulting chemical profile.
TABLE 5. Quality Index of Cultivated Cordyceps sinensis

<table>
<thead>
<tr>
<th></th>
<th>American mycelium</th>
<th>Oriental mycelium</th>
<th>Standardized extracts</th>
<th>Wild Cordyceps</th>
<th>Aloha Medicinals hybrid Cordyceps</th>
</tr>
</thead>
<tbody>
<tr>
<td>HEAA range</td>
<td>0.00-0.041%</td>
<td>0.00-0.071%</td>
<td>0.050-0.073%</td>
<td>0.071-0.100%</td>
<td>0.190-0.323%</td>
</tr>
<tr>
<td>HEAA avg</td>
<td>0.005%</td>
<td>0.025%</td>
<td>0.070%</td>
<td>0.090%</td>
<td>0.303%</td>
</tr>
</tbody>
</table>

Note: Test results through the HPLC/MS methods listed here

DISCUSSION

We found great differences in final cultivated Cordyceps product quality, dependent on several factors, such as strain, substrate, and culture parameters followed. The milo- or milo/millet-based substrates proved to be the best of all that we tested, based on the quantities of the secondary metabolites we were measuring (HEAA). If other end products are desired, then perhaps other substrate compositions will prove more valuable. The only way to determine the end result is to try different substrates and analyze the resulting mycelium. Parameters that are commonly used in farming edible mushrooms such as speed of growth, density, or color were found to be generally unreliable indicators of Cordyceps quality, if quality is to be defined in terms of concentration of bioactive compounds. These “target compounds desired” should be used as the basis for formulating substrate composition, with composition used as one of the three arms of strategic husbandry to insure that medicinal properties are optimized. The other two arms of optimum cultivation—culture parameters and strain—are no less important, and all three factors must be evaluated together.

The HPLC analytical protocols presented in this article, and to a lesser degree the GC protocols, allow quick and relatively inexpensive methods of analysis to guide the cultivator in their search for quality. Until better protocols are developed or further bioactive compounds are identified, we urge all cultivators of medicinal Cordyceps to adopt these test methods, so that all the different Cordyceps on the market will be evaluated consistently. By taking the approach of adding together the quantities of several different compounds, the total Quality Index of dissimilar products can equate, while specific levels of this or that compound still vary due to strain or growing conditions. Furthermore, if and when customers call for greater quantities of any specific compound(s) in their product, these test methods offer rapid means of developing new cultivation strategies, again based around the target compounds desired.

These same comments of repeated trial and analysis apply equally to the specific cultivation parameters, such as high temperature for the production of higher quantities of Adenosine or low temperature hypoxia for production of greater quantities of the deoxygenated nucleosides. This work in modifying the culture parameters suggests that other parameters may yield some interesting compounds, such as high sulfur content in the substrate or atmosphere for the production of sulfated polysaccharides, such as ones that have shown promise as antifungal and antiviral agents (Wasser and Weis, 1999).

It is proposed by these authors that the simple addition of components to the substrate, coupled with carefully controlled growing conditions, could result in the production of some wonderfully unique compounds by this fungus. We have even grown Cordyceps sinensis on a substrate consisting primarily of chocolate milk, and one composed solely of ground squid. These substrates resulted in some very unique compounds produced, the usefulness of which has yet to be determined.

Hybridization of fungal strains is a relatively new practice, and the simple low-cost methods shown here allow the average cultivator a way to develop their own strains, suited to their particular requirements. Perhaps the cultivator has access only to some substrate material on which the Cordyceps grows but poorly. Here then is a method that can be applied to “create” a new strain, which will fulfill the intended function, growing on this substandard.
medium. This is only one of nearly limitless uses for this hybridization technique. We have found that in some cases, we are even able to hybridize between different species, such as crosses between Cordyceps sinensis and C. sobolifera. This may indicate that these two species are closer together in the evolutionary tree than was previously thought, or it could mean that we have a method by which to trick nature into performing that which does not take place naturally. But whatever the explanation, it certainly allows us to create unique new strains with which to work.

CONCLUSIONS

There is a large and growing market worldwide for Cordyceps as a medicine and as a health supplement. This large market demand coupled with the high prices of wild Cordyceps has given rise to an ever-increasing cultivation attempt by mushroom cultivators in many countries. Until now, there has been no standard of quality and no well-understood cultivation protocols with which to consistently produce high quality Cordyceps. Compounding this has been the lack of any standardized test method by which to analyze and ascertain whether the cultivated Cordyceps was of a good quality or not. Here we have shown how hybridization results in higher quality Cordyceps and have presented an overview of cultivation techniques, bioactive target compounds and quality index references, analytical protocols, substrate composition, cultivation methodology, and hybridization techniques for production of the cultivars necessary in accomplishing these ends. This should serve as an outline for the production of high-quality, economical Cordyceps, suitable for pharmaceutical and nutraceutical use, at a consistently high level of bioactive ingredients. This is a safe, natural, and traditional medicine with a long history of use that is now available and ready to be used on an ever-increasing scale in the treatment of modern diseases.

REFERENCES


