

Oxygen Radical Absorbance Capacity (ORAC) Assay

info@activeconceptsllc.com • Phone: +1-704-276-7100 • Fax: +1-704-276-7101

Tradename: BiEau® Actif Tri-Mushroom

Code: 16908

CAS #: 223751-82-4 & 2055734-24-0 & 1174745-80-2

Test Request Form #: 5638

Lot #: N190923B

Sponsor: Active Concepts, LLC; 107 Technology Drive Lincolnton, NC 28092

Study Director: Maureen Danaher

Principle Investigator: Jennifer Goodman

Test Performed:

Oxygen Radical Absorbance Capacity (ORAC)

Introduction

Reactive oxygen species (ROS) are generated by normal cellular processes, environmental stresses, and UV irradiation. ROS are dangerous to cellular structures and functional molecules (i.e DNA, proteins, lipids) as they act as strong oxidizing agents or free radicals. The oxygen radical absorbance capacity (ORAC) assay is a standard method used to assess antioxidant capacity of physiological fluids, foods, beverages, and natural products. The assay quantitatively measures a sample's ability to quench free radicals that have the potential to react with and damage cellular components.

Oxygen Radical Absorbance Capacity (ORAC) assay was conducted to assess the antioxidant capacity of **BiEau® Actif Tri-Mushroom**.

Assay Principle

This assay is based upon the effect of peroxyl radicals generated from the thermal decomposition of 2, 2'-azobis-2-methyl-propanimidamide dihydrochloride (AAPH) on the signal intensity from the fluorescent probe, fluorescein, in the presence of an oxygen radical absorbing substance. The degree of change is indicative of the amount of radical damage and the presence of antioxidants results in an inhibition in the free radical damage to the fluorescein. The antioxidant protection of the sample can be calculated by comparing it to a set of known standards. Trolox®, a water soluble vitamin E analog, with known antioxidant capabilities is used in this ORAC assay as the standard for measuring the antioxidant capacity of unknown substances. ORAC values, expressed in µM of Trolox® equivalents (TE), are calculated using the area under the curves (AUC) of the test product, Trolox®, and the control materials. Trolox® equivalency is used as the benchmark for antioxidant capacity of mixtures since it is difficult to measure individual components.

Information contained in this technical literature is believed to be accurate and is offered in good faith for the benefit of the customer. The company, however, cannot assume any liability or risk involved in the use of its chemical products since the conditions of use are beyond our control. Statements concerning the possible use of our products are not intended as recommendations to use our products in the infringement of any patent. We make no warranty of any kind. expressed or implied, other than that the material conforms to the applicable standard specification.

Page 1 of 3 Version#2/02-11-20/Form#56



Oxygen Radical Absorbance Capacity (ORAC) Assay

info@activeconceptsllc.com • Phone: +1-704-276-7100 • Fax: +1-704-276-7101

Materials

A. Equipment: Synergy H1 Microplate reader (BioTek Instruments, Winooski, VT); Gen5

software (BioTek Instuments, Winooski, VT); Pipettes

B. Buffers: 75mM Potassium Phosphate (pH 7.4); Deionized H₂O

C. Reagents: 2,2'-Azobis(2-methylpropionamidine) dihydrochloride (AAPH) (153mM);

6-Hydroxy-2,5,7,8-tetramethylchromane-2-carboxylic acid

(Trolox®);Fluorescein Sodium Salt (4nM)

D. Preparation: Pre-heat (37°C) Synergy H1 Microplate reader; Prepare Trolox®

standards, sample dilutions, fluorescein solution, and AAPH.

E. Microtitre Plates: Corning 96 Well Black Side/Clear Bottom Microplates

Methods

Solutions of **BiEau® Actif Tri-Mushroom** and Trolox® (positive control) were prepared in 75mM potassium phosphate buffer. Materials were prepared at three different concentrations/dilutions. Trolox® was used as a reference for antioxidant capacity and prepared at a concentrations ranging from 12.5 μ M to 200 μ M in 75mM potassium phosphate buffer.

For the ORAC assay, $25\mu L$ of test material and Trolox® were combined with 150 μL of fluorescein in 75mM potassium phosphate buffer and incubated in the Synergy HT Microplate reader at 37°C for 30 minutes. At the end of the incubation period, $25\mu L$ of AAPH (153mM in 75mM potassium phosphate buffer) were pipetted into each well. Fluorescent measurements were then taken every 2 minutes for approximately 2 hours at an excitation wavelength of 485nm and an emissions wavelength of 520nm.

The AUC and Net AUC values of the standards and samples were determined using Gen5 2.0 Data Reduction Software (BioTek Instruments) using the below equations:

$$AUC = 0.5 + \frac{R2}{R1} + \frac{R3}{R1} + \frac{R4}{R1} + \dots + \frac{Rn}{R1} \rightarrow Where R is fluorescence reading$$

$$Net\ AUC = AUC_{sample} - AUC_{blank}$$

The standard curve was obtained by plotting the Net AUC of different $Trolox^{@}$ concentrations against their concentration. ORAC values of samples were then calculated automatically using the Gen5 software to interpolate the sample's Net AUC values against the $Trolox^{@}$ standard curve. ORAC measurements for the test material were expressed in micro molar $Trolox^{@}$ equivalents (μ MTE), where 1 ORAC unit is equal to 1 μ MTE.

ORAC values are also calculated in Units/milliliter (U/mL). The equation used for the calculation is shown below:

$$ORAC\ (U/mL) = (50 \times Dilution\ Factor) \times \left(\frac{AUC_{Sample} - AUC_{Blank}}{AUC_{Tralox} - AUC_{Blank}}\right)$$

Information contained in this technical literature is believed to be accurate and is offered in good faith for the benefit of the customer. The company, however, cannot assume any liability or risk involved in the use of its chemical products since the conditions of use are beyond our control. Statements concerning the possible use of our products are not intended as recommendations to use our products in the infringement of any patent. We make no warranty of any kind. expressed or implied, other than that the material conforms to the applicable standard specification.



Oxygen Radical Absorbance Capacity (ORAC) Assay

info@activeconceptsllc.com • Phone: +1-704-276-7100 • Fax: +1-704-276-7101

Results

BiEau® Actif Tri-Mushroom showed very potent antioxidant activity at 5.0% and 2.5% concentrations.

The ORAC value expressed in U/mL for 5.0% BiEau® Actif Tri-Mushroom is 198.

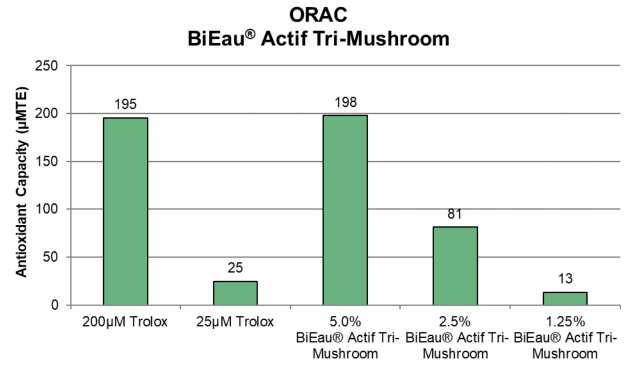


Figure 1: Antioxidant capacities

Discussion

As shown in figure 1, **BiEau® Actif Tri-Mushroom** (16908) exhibited greater antioxidant activity than 200µM Trolox®. The antioxidant capacity of **BiEau® Actif Tri-Mushroom** increased as the concentration increased, as a result we can assure that its ability to minimize oxidative stress is dose dependent. Maximizing the antioxidant capacity on a cellular level allows for ROS to be dealt with at a rate that provides protection from cellular damage. This cellular damage can be seen as physical signs of aging such as wrinkles, loss of elasticity, unwanted pigmentation, and skin unevenness with slow regeneration.

It can therefore be concluded that **BiEau® Actif Tri-Mushroom** is capable of providing antioxidant properties and aids in the anti-aging process through protection at the cellular level.

Information contained in this technical literature is believed to be accurate and is offered in good faith for the benefit of the customer. The company, however, cannot assume any liability or risk involved in the use of its chemical products since the conditions of use are beyond our control. Statements concerning the possible use of our products are not intended as recommendations to use our products in the infringement of any patent. We make no warranty of any kind. expressed or implied, other than that the material conforms to the applicable standard specification.