

**Tradename:** AC ExoTone

**Code:** 60194

**CAS #:** 7732-18-5 & 85251-63-4 & 123465-35-0 (or) 8002-43-5

**Test Request Form #:** 10226

**Lot #:** N230613C

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**Test Performed:**

Tyrosinase Inhibition Assay

**Introduction**

Tyrosinase is a key enzyme in melanin biosynthesis, involved in determining the color of mammalian skin and hair. Tyrosinase's main application is to identify new potent tyrosinase inhibitors in the cosmetic industry. Tyrosinase is a copper-containing monooxygenase that is widely distributed in nature. The enzyme catalyzes the first two reactions of melanin synthesis, the hydroxylation of L-tyrosine to 3,4-dihydroxyphenylalanine, L-dopa, and the oxidation of L-dopa to dopaquinone. This quinone is a highly reactive compound and can polymerize spontaneously to form melanin. Tyrosinase is one of the causes of hyperpigmentation, an over-production of dermal melanin pigment, leading to melasmas, freckles, age-spots, and liver spots.

A tyrosinase inhibition assay was conducted to determine the ability of **AC ExoTone** to inhibit tyrosinase, indicating a potential component to reduce hyperpigmentation. The key active ingredient in **AC ExoTone**, *Pyrus malus* (Apple) Fruit Extract, was tested to demonstrate the superior nature of bioauthentic exosomes as a delivery system.

**Assay Principle**

This assay is based on the conversion of L-tyrosine into a dopachrome complex by tyrosinase. This dopachrome complex has an absorbance at 510nm and can be quantitated through optical density measurements. The greater the inhibition exhibited by the sample, the lower the optical density value due to the lack of L-tyrosine conversion. This is a time course assay, after which the results are analyzed and compared to a known tyrosinase inhibitor, Kojic Acid.

## Materials

- A. Kit:** Tyrosinase Inhibition Screening Kit, Colorimetric (Sigma; MAK257)\*
- B. Equipment:** Synergy H1 Microplate reader (BioTek Instruments, Winooski, VT); Gen5 software (BioTek Instruments, Winooski, VT); Pipettes
- C. Buffers:** Tyrosinase Assay Buffer (MAK257A)\*
- D. Reagents:** Tyrosinase Substrate (MAK257B)\*; Tyrosinase (MAK257C)\*; Tyrosinase Enhancer (MAK257D)\*; Inhibitor Control Kojic Acid (0.01% (0.75mM), MAK257E)\*
- E. Preparation:** Synergy H1 Microplate reader
- F. Plates:** 96 Well Microtiter Plates; Multichannel sample wells

\*Or suitable alternatives, subject to change without notice based off vendor availability

## Methods

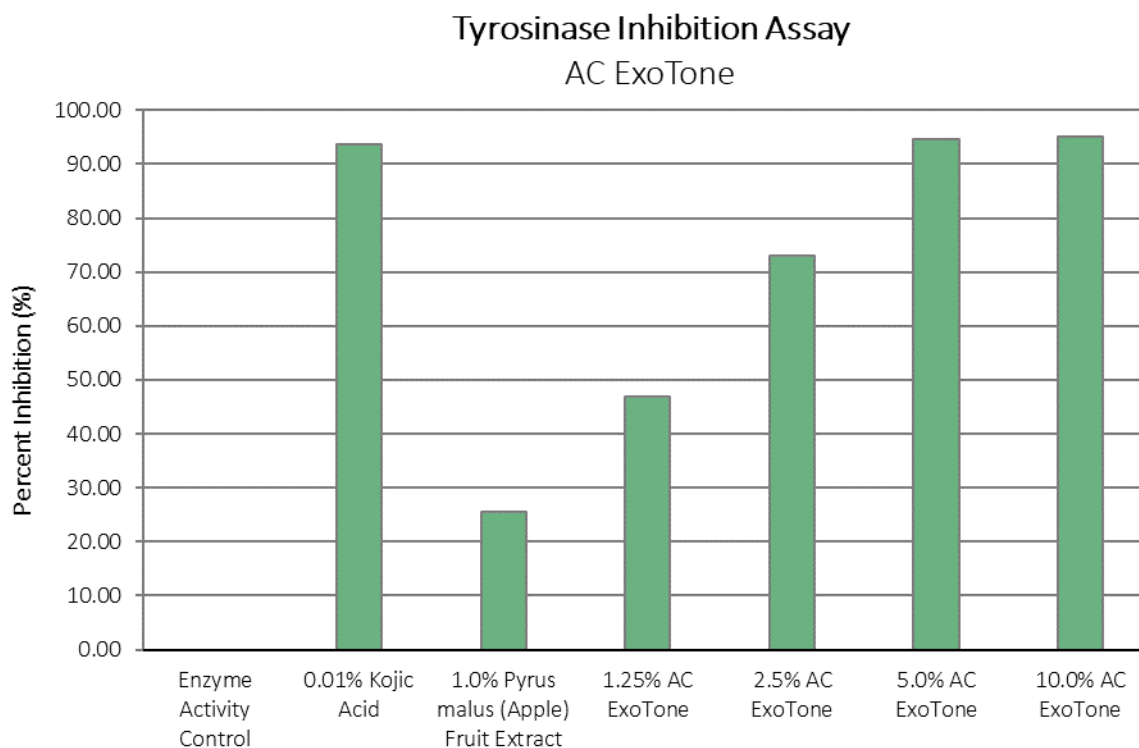
Solutions of **AC ExoTone** (1.25%, 2.5%, 5.0%, 10.0%), 0.01% Kojic Acid (positive inhibitor control), tyrosinase substrate solution, and tyrosinase enzyme solution were prepared in tyrosinase assay buffer. Additionally, a 1.0% solution of *Pyrus malus* (Apple) Fruit Extract was prepared in tyrosinase assay buffer. This concentration of *Pyrus malus* (Apple) Fruit Extract is equivalent to the amount present in 5.0% **AC ExoTone**. For the negative control, tyrosinase assay buffer was used and is labeled as the tyrosinase enzyme control (EC). 20 µL of test material and controls were combined with 50 µL of the tyrosinase enzyme solution and incubated at 25°C for 10 minutes. Next, 30 µL of the tyrosinase substrate solution was added to each respective test well. The plate was placed in the Synergy H1 reader and optical density measurements were then taken every minute for 60 minutes at 510 nm.

The slope of each sample was calculated by dividing the net  $\Delta Abs$  ( $Abs_2 - Abs_1$ ) values by the  $\Delta Time$  ( $T_2 - T_1$ ). The percent of tyrosinase inhibition was calculated by the below equation:

$$\% \text{ Inhibition} = \frac{Slope_{EC} - Slope_{Sample}}{Slope_{EC}} \times 100$$

## Results

The data obtained from this study met criteria for a valid assay and the control performed as anticipated. Regarding the experimental conditions, 1.0% of the *Pyrus malus* (Apple) Fruit Extract inhibited tyrosinase by 25%, whereas all concentrations of **AC ExoTone** inhibited tyrosinase in a dose dependent fashion.



**Figure 1:** Tyrosinase Inhibition.

## Discussion

As shown in Figure 1, Kojic Acid, inhibited tyrosinase by 94%. This data demonstrates the cosmetic industry standard skin lightening active ingredient (Kojic Acid) provides skin lightening benefits by inhibiting tyrosinase. **AC ExoTone** was able to inhibit tyrosinase comparably to the positive control, Kojic Acid (Figure 1). The inhibition activity of **AC ExoTone** increased as the concentration increased, confirming the ability to decrease tyrosinase activity is dose dependent (Figure 1).

Similarly, **AC ExoTone** at 1.25%, 2.5%, 5.0%, and 10.0% inhibited tyrosinase by 47%, 73%, 95%, and 95%, respectively. Conversely, 1.0% of the *Pyrus malus* (Apple) Fruit Extract only inhibited tyrosinase by 25%. Together, these data demonstrate the inhibition activity of **AC ExoTone** improved with increasing concentrations, illustrating the ability of **AC ExoTone** to decrease tyrosinase activity is dose dependent. Additionally, all concentrations of **AC ExoTone** outperformed the *Pyrus malus* (Apple) Fruit Extract in tyrosinase inhibition, highlighting the superior nature of bioauthentic exosomes as a delivery system.

Collectively, these data indicate **AC ExoTone** inhibits tyrosinase activity and can provide skin lightening benefits to counteract challenges associated with hyperpigmentation.