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Effect of Potassium Bromate on *OXR1* Gene Expression and Cell Growth in *Tetrahymena thermophila*

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### Introduction

- The unicellular, eukaryotic ciliate *Tetrahymena thermophila* (*T. thermophila*) is a useful model organism in molecular biology.
- The protein encoded by *OXR1* is required to prevent oxidative damage resulting from energy production.
- Potassium Bromate (*KBrO₃*) is a strong oxidizing agent that is used as a leavening agent in baked goods.
- Previous studies have shown that *KBrO₃* is toxic to living organisms.

**Hypothesis:** If *Tetrahymena* are exposed to *KBrO₃* in their environment, oxidative damage will be induced and, as a result, expression of *OXR1* will increase while cell growth rate decreases.

### Methods

- **Primer synthesis:** IDT software was used to design primers for the *OXR1* gene.
- **Culturing:** *T. thermophila* cultures were maintained in NEFF media. During treatment, experimental cultures were transferred to culture dishes with SPP media containing 0.5mM *KBrO₃*. Control cultures were grown in pure SPP media.
- **RNA extraction:** RNA was extracted using Qiagen’s RNeasy Mini Kit.
- **Reverse transcription:** RevertAid was used to synthesize cDNA from extracted mRNA.
- **PCR** was performed using GoTaq master mix, and *BTU1* gene expression was used as a positive control.
- **Gel electrophoresis** was performed on RT-PCR reactions. Relative PCR signal intensities were analyzed using ImageJ.
- **Cell growth rate** was determined by counting cell number with a hemocytometer twice a day over a 72 hour period.

### Results

- **Figure 1:** Round 1 *OXR1* PCRs. Both control (lanes 3-4) and experimental (lanes 5-6) groups show *OXR1* expression.
- **Figure 2:** Round 1 *BTU1* PCRs. Three control (top) groups show *BTU1* expression, while negative controls do not.
- **Figure 3:** Round 2 *OXR1* PCRs. The control (lanes 3-4), one experimental (5) and three negative control groups show (b) *OXR1* expression.
- **Figure 4:** Round 2 *BTU1* PCRs. Three positive controls (top) and all negative controls (bottom) show *BTU1* expression.
- **Figure 5:** Growth curve of control and *KBrO₃* treated *Tetrahymena* cultures measured over a 72 hour period. From 0-72hrs, p-values: 1.097 0.71, 0.18, 0.56, 0.38, 0.80
- **Figure 6:** Image of *Tetrahymena* cells fixed in glutaraldehyde for counting with a hemocytometer.

### Conclusions

- Both experimental trials indicated that exposure to *KBrO₃* did not induce any major change in expression of *OXR1*, as seen in Figures 1 and 3.
- There was no significant increase in the quantitative relative RT-PCR signal of *OXR1*, as shown in Figure 7 (p=0.25).
- *KBrO₃* exposure had no significant effect on cell growth as seen in Figure 5. (p-values = T0:1, T12: 0.97, T24: 0.71, T36: 0.18, T48: 0.56, T60: 0.38, T72: 0.80).
- The results disagreed with our hypothesis in regard to both gene expression and cell growth rate.
- **Future Directions:** Optimize the concentration of *KBrO₃*, and increase the number of experimental trials.

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