



EDULAB FOR STUDENTS: MAGNETTECH ESR5000

# EPR Guide to Beer Freshness

## EPR of Beer

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**Experiment Hashtag: #beer, #shelf-life, #antioxidants #Educate2Resonate**

### Keywords:

Beer, free radicals, antioxidants, shelf-life

### Target group:

Advanced Undergraduate or Graduate,  
General Chemistry, Analytical Chemistry,  
Food Chemistry, Food Safety and Control  
Laboratory, General Life Sciences

### Objectives:

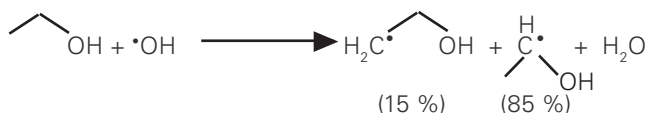
This laboratory experiment utilizes an exploration of beer's storage stability to introduce basic aspects of Electron Paramagnetic Resonance (EPR) spectroscopy. Radicals formed upon aerobic forced aging of beer samples are detected by using a spin trap. Students are introduced to basic principles of EPR spectroscopy as well as food and radical chemistry with a simple example taken from everyday life. The methodology presented provides students with invaluable insights into EPR spectroscopy and the role of free radicals in food chemistry.

## Background of the Experiment:

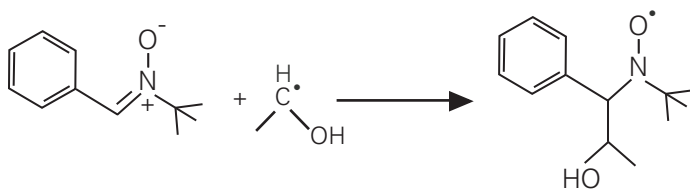
EPR spectroscopy detects free radicals in the degradation processes of beer. The role of oxidation processes in the aging of beer and the accompanying degradation in flavor is well known. EPR spectroscopy has been employed to verify that short-lived radicals are present as intermediates during the aging of beer. It has been shown that Fenton-like reactions can occur in beer and that oxygen acts as a precursor for the formation of hydrogen peroxide. This hydrogen peroxide, in a subsequent reaction, reacts with metal ions to form the highly reactive hydroxyl radical ( $\cdot\text{OH}$ ).



The hydroxyl radical reacts with several compounds in beer and forms an array of radical follow-up products. The most important of these is the reaction between the hydroxyl radical and ethanol.



Radical follow-up products such as the 2-hydroxyethyl-radical can be “captured” and detected using spin trapping reagents.



Even though the hydroxyl radical is highly reactive, the oxidation of beer still is a slow process. This is attributed to the low concentrations of hydrogen peroxide and metal ions in beer. Therefore, for a positive control, additional Fe(II) and hydrogen peroxide is added to a beer sample to observe immediately the effect of hydroxyl radical burst on beer oxidation. This “lag-time measurement” provides direct information on the amount of antioxidants in beer – the more antioxidants present, the longer oxidation processes are prevented. This time is called the “endogenous antioxidative potential”. Only after all antioxidants in the beer are consumed, the signal of the radical adduct is detected. To additionally accelerate this process, “aerobic forced aging” is often employed. Here, the beer is heated under ambient atmosphere to consume the antioxidants in the sample faster.

Students are going to determine the “endogenous antioxidative potential” of beer using EPR spectroscopy. Students’ task is to evaluate the time after which all antioxidants in beer are consumed using the method of “aerobic forced aging”. Additionally, students determine the influence of adding Fenton reagent (hydrogen peroxide + iron(II)sulphate) to the sample.

## Preparation:

The instructor needs to remove the carbon dioxide from the beer by placing the beer in an ultrasonication bath for 15 min and subsequently decanting it the day before the experiment. The removal of  $\text{CO}_2$  is necessary because bubbles in the capillaries could disturb the EPR measurements. The experiment is designed to be carried out by students working in pairs for approximately three hours. In addition to protective eyewear, the use of nitrile gloves is required. The spin trap *N-tert-Butyl- $\alpha$ -phenylnitrone* (PBN) is an irritant. The laboratory experiment should be carried out in a well-ventilated lab space. After completing the experiments, an additional 1 – 1.5 hours will be needed to write a report. It is assumed that students have already covered introductory concepts of EPR and have a basic understanding of instrumental parameters.

## Glossary

**EPR:** Electron paramagnetic resonance or electron spin resonance spectroscopy is a method for studying materials that have unpaired electrons. The basic concepts of EPR are analogous to those of nuclear magnetic resonance, but the spins excited are those of the electrons instead of the atomic nuclei.

**Free radicals:** An atom, molecule, or ion that has at least one unpaired valence electron.

### Spin trapping:

An analytical technique employed in chemistry and biology for detection and identification of short-lived free radicals through the use of EPR spectroscopy.

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For comprehensive information on EPR basics and optimizing instrumental parameters, students can consult the Magnettech ESR5000 educational kit, which is provided along with the benchtop EPR spectrometer.

To perform the measurements, an installed Magnettech ESR5000 spectrometer is required.

### Experimental Setup:

#### Materials:

- *N-tert*-Butyl- $\alpha$ -phenylnitrone (PBN) (MW = 177.24 g/mol)
- Hydrogen peroxide (30 % in H<sub>2</sub>O) (MW = 34.01 g/mol)
- Iron(II)sulphate heptahydrate (MW = 278.01 g/mol)
- Lager beer (Bud Light, Modelo, Budweiser) – 1 bottle or 1 can
- Pipettors and tips
- 50 ml beakers
- 50  $\mu$ L capillaries
- Capillary sealant
- Eppendorf self-lock tubes
- Small glass vials (4 – 5 ml) with screw caps
- Drying oven

### Sample Preparation:

1. The following stock solutions need to be prepared:
  - 50 mM PBN in beer (2 ml)
  - 5 mM FeSO<sub>4</sub> in beer (2 ml)
2. The PBN stock solution is stored at 60 °C in the drying oven. The FeSO<sub>4</sub> solution is stored at room temperature.
3. Every 15 minutes, PBN solution is measured by EPR to evaluate the endogenous antioxidative potential over time (10 points in total).
4. The peak-to-peak amplitude is plotted versus time in minutes using Excel (or similar software) to evaluate the “lag-time”.
5. In addition, the following solution is prepared and measured:
  - 0.2 ml PBN stock solution + 0.2 ml FeSO<sub>4</sub> stock solution + 0.6 ml beer + 2  $\mu$ l H<sub>2</sub>O<sub>2</sub> (30 %)

### Abbreviations

#### EPR:

Electron paramagnetic resonance





## References:

1. Andersen, M.L. and Skibsted, L.H., *Electron spin resonance spin trapping identification of radicals formed during aerobic forced aging of beer*, J. Agric. Food Chem. 46(4) (1998) 1272-1275
2. Kaneda, H. et al., *Detection of free radicals in beer oxidation*, J. Food Sci. 53(3) (1988) 885-888
3. Schmallegger M. and Gescheidt G., *Antioxidant activity of beer: an EPR experiment for an undergraduate physical-chemistry laboratory*, J. Chem. Educ. 95 (2018) 2013-2016

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