



### Challenge

Studying the generation of reactive oxidative species in response to inflammation.

### Solution

Low light luminescence imaging using the iBox Scientia.

## Low Light Imaging of Reactive Oxidative Species within Wound Sites Using Luminol Using the iBox Scientia

### Introduction

Luminol is a standard tool in forensic science. Typically, crime scenes are analyzed for blood samples by applying a solution of luminol and oxygen radicals onto surfaces. The iron-laden hemoglobin molecules present in blood serve as catalysts for the reaction which can then be captured by photograph. When combined with an oxidizing agent and a catalyst, luminol emits blue light.

For routine applications, as in forensic testing or in a laboratory setting, the luminol molecule is combined with a hydroxide salt to generate a dianion. When in the presence of a superoxide radical, the dianion molecule produces unstable organic peroxide. Upon relaxing to a ground state, the dianion emits a photon within the visible spectrum, primarily in the blue light range (peak emission is at 424 nm). The reaction typically lasts for a short time, on the order of seconds, as the luminol molecule is consumed in the generation of light.

In addition to forensics, the use of luminol extends into preclinical research and has been used when studying the generation of reactive oxidative species (ROS). Normal cellular respiration produces oxide radicals. Additionally, ROS are generated during acute inflammation as well as in response to environmental stimulants. Therefore, inflammation can be studied using luminol due to the generation of substrate for the luminescent reaction to occur.

Recent studies have examined the use of luminol within in vivo models. Specifically, the role of inflammatory cells has been under study due to the response of the immune system to stress. Typical inflammatory processes involve a cellular phase in which inflammatory cells are recruited to the site of trauma to mount a response. This response takes the form of an "oxidative burst" in which ROS are generated by a host of enzymes within the phagocytic cell to both destroy

pathogens as well as to recruit and activate other inflammatory cells.

Luminol was used *in vivo* to monitor the generation of ROS, specifically due to the inflammatory response mounted by phagocytic cells. Two mice, a control and a mouse

deficient in a factor that clears cells from the wound, were wounded at a specific time point to determine the degree of inflammation of a knockout mouse and imaged 12 hours later.

## Materials and Methods

### Samples and reagents

- Imaging System: iBox Scientia
- Mice
- Luminol

### Samples preparation

Two mice were wounded at 12 hours prior to imaging. The mice were briefly anesthetized with xylazine/ketamine and injected with luminol intraperitoneally. The mice were then placed in the prone position on the iBox<sup>®</sup> Scientia<sup>™</sup> Small Animal Imaging System's imaging stage and images were captured with a highly sensitive CCD camera immediately after injection of the luminol.

### Imaging

Each mouse was imaged for 20 minutes at 4x4 binning (Fig 1). To ensure euthermia, the imaging stage/warming plate maintained the temperature of the mouse at 37°C. Two images were acquired at each time point using a bright field (white light) and a luminescent channel. These images were then selected for overlay via the composite feature using VisionWorks software. The luminescent image was pseudo colored with an intensity map. The pseudo colored luminescent image was then overlaid on the bright field image (Fig 2).

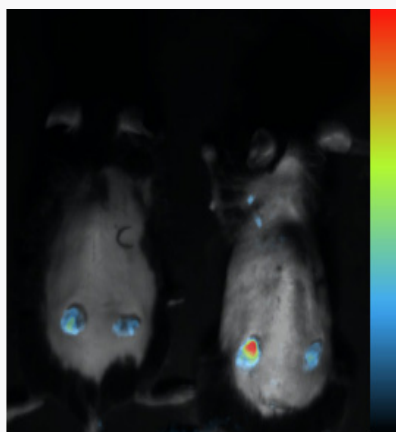


Figure 1. Luminescence at a site of local inflammation. Both mice were imaged 12 hours after wounding and immediately after luminol injection. Note that the intensity in the control mouse (left) was significantly lower than the experimental mouse (right).

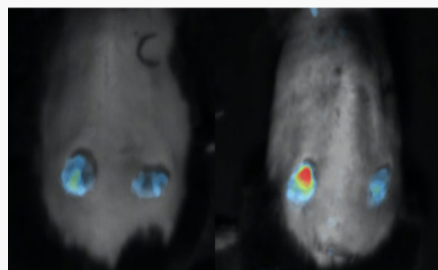


Figure 2. Luminescence at a site of local inflammation. Note the greater intensity of the experimental mouse lesions (right) as compared to control (left).

## Results and Discussion

A key biological catalyst in the generation of ROS in the inflammatory response is myeloperoxidase (MPO), an enzyme that converts hydrogen peroxide to hypochlorous acid in addition to several other reactive species. This enzyme is present in phagocytic cells, inflammatory cells which engulf pathogens and cellular debris.

The acute inflammatory process includes the recruitment of phagocytic cells, such as polymorphonuclear leukocytes or macrophages. These cells generate ROS during the phagocytic process to both destroy pathogens as well as to clear the wound of debris from damaged tissue. Acute wounding, either through mechanical or chemical means, generated intense signals upon exposure to luminol after 12 hours. The knockout mouse showed the greatest generation of luminescence, suggesting that this mouse has impairment in clearing of inflammatory cells.

## Conclusion

The iBox Scientia is a powerful tool for imaging disease processes within an in vivo model. The Scientia couples a light-tight darkroom, a highly sensitive CCD camera and sophisticated optics for low-light imaging, ideal for luminescence related applications. Through the use of luminescence, real-time imaging of ROS generation by inflammatory cells can be observed evolving within the wound of a mouse.

## References

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