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# HOW DO YOU VIEW?

Illuminating biological processes and structures through (bio)chemical detection.

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# Can your imager do this?



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## Illuminating biological processes and structures through (bio)chemical detection.

Ever since Antonie van Leeuwenhoek first described the microbial world seen through his microscope, scientists have been looking for new ways to watch biology unfold. These days much of the imaging that goes on in a lab occurs beyond the oculars and takes advantage of sophisticated chemistry to "see" biological processes and structures. As we approach the 345th anniversary of van Leeuwenhoek's first publication, let's take a moment to celebrate these powerful techniques that have shed so much light on how life works.





Invaluable for their ability to deliver direct quantitation at the single-molecule evel, radiolabeled chemicals were once the only method for following dynamic biochemical processes. However, radioisotope usage has gradually declined from a peak towards the end of the last century, as safer alternatives have been developed. Still, for some applications, the advantages of radiolabeled compounds outweigh the risks, which can be ameliorated through the use of safety procedures.

ne widespread in the 1950s in rts by the Atomic Energy Commiss

lsotope	Half-life	Decay Mode	Emission (MeV)	Shielding	Maximum Range in Air	Maximum Range in Tissue
³Н	12.3 y	β-	0.0186	None	0.25 in (0.6 cm)	Negligible
<sup>14</sup> C	5730 y	β-	0.156	None	10.0 in (24 cm)	0.012 in (0.28 mm)
<sup>18</sup> F	1.8 h	β+, EC	0.63 (β+), 1.66 (EC)	Plastic	62 in (1.6 m)	0.09 in (0.23 cm)
<sup>32</sup> P	14.3 d	β-	1.71	Plastic	20 ft (6.1 m)	0.33 in (0.76 cm)
<sup>35</sup> S	87.3 d	β-	0.17	None	10.2 in (26 cm)	0.015 in (0.32 mm)
125	59.5 d	EC, y	0.19 (EC), 0.35 (γ)	Lead	N/A	N/A



1. A.N.H. Creager. Life Atomic: A history of radioisotopes in science and medicine. Chicago. IL. The University of Chicago Press. 2013. 2. M.H. England and E. Majer-Reimer. "Using chemical tracers to assess ocean models." Rev Geophys 39(1): 29-70 cal and Non-clinical Application and Future Uses," Appl Biochem Biotechnol 173(2): 333-355, 2014. 5. G.H. Thorpe, "Phenols as enhancers of i n 31(8):1335-1341, 1985. 6. G.G. Stokes, "XXX. On the change of refrancibility of light," Phil Trans R Soc Lond 142, 463-562, 1852. 7. A. Stirbet and Govindiee, "On the relation between the Kautsk stforms for fluorescence imaging," Chem Soc Rev 42(2):622-661, 2013. 9. S.R. Tsai and M.R. Hamblin. "Biological effects and medical applications of infrared radiation." J Photochem Photobiol B 170:197-207, 20" rescence Imaging in Humans with Indocyanine Green: A Review and Update," Open Surg Oncol J2(2): 12-25, 2010. 11. E.A. Owens, et al., "Tissue-Specific Near-Infrars 49(9)·1731-1740 2016

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PLICATIONS Comm r: imaging tissue samples, m blots, northern blots, and

### **U KNOW?** Nuclear testing during the early Cold War period mount of <sup>3</sup>H on the Earth's surface by several orders of <sup>3</sup>H a useful tracer for atmospheric pa

Highly specific, extremely sensitive, safer than radioisotope imaging, and more amenable to signal amplification than fluorophores,<sup>3</sup> chemiluminescent assays such as ELISAs and Western blots are almost ubiquitous in life science laboratories. However, because chemiluminescence harnesses biological reactions, its reproducibility is affected by experimental and environmental factors such as reaction duration, reactant amount, light exposure, and environmental conditions (e.g., temperature, buffers).<sup>2</sup>



APPLICATIONS



## The Enzyme-Luminol Reaction

Luminol generates chemiluminescence when the dianion form is oxidized, creating excited intermediates which produce light during stabilization to the ground state

**Complexities** Apart from the oxidizer and catalyst, chemiluminescence intensity is also affected by experimental conditions such as pH (alkaline is best but not too alkaline), heat (luminol is thermally unstable), luminol concentration. and protein concentration

**ECL** Phenols and other compounds can enhance the magnitud and stability of catalyzed luminol oxidation-derived light emission by as much as 1.000-fold.<sup>5</sup> leading to the advent of



**Oxidizers** Commonly used oxidizing agents include **Enzymes**, including horseradish peroxidase (HRP) **Detection** The emission range of luminol is ozone, halogens, singlet oxygen, hydrogen peroxide, and alkaline phosphatase (Alk Phos), as well as metal typically ~370-490 nm with a maximum at ~420 nm, and hypochlorites. Of these, hydrogen peroxide is most ions serve as catalysts for popular because it augments the luminescent intensity the luminol-oxidant reaction of luminol.<sup>4</sup>

but can vary significantly. For example, the presence of iron has been documented to cause a spectrum shift resulting in a maximum emission of 455 nm.<sup>4</sup>

Often associated with microscopy, fluorescence-based techniques have been rapidly expanding from the microscope slide and into the heart of the lab. Another alternative to radioactivity, non-microscopy uses for fluorescence became more widespread in part as a result of the human genome sequencing project and the development of dye-based sequencing. From there, rapid advances in detector technology and powerful computer processors have facilitated fluorescence use for protein detection, such as in Western blots, where the ability to use several fluorophores at once (multiplexing) can greatly enhance experimental efficiency.

