

## Monitoring curli production in liquid culture using EbbaBiolight

This protocol describes how to monitor kinetics of *Salmonella* extracellular matrix (curli) production in liquid culture. The method described here is based on Choong et al. (2016) *npj Biofilms and Microbiomes*, 2, 16024 where isogenic mutants of *S. Enteritidis* were used to identify the extracellular matrix components curli and cellulose as targets for optotracer binding. When used as described, EbbaBiolight does not label *Salmonella* cell wall and does not influence biofilm formation.

**Note:** When adapting this technique, please make sure to include relevant controls to verify that EbbaBiolight does not affect biofilm formation, to confirm curli as EbbaBiolight binding target and to exclude pH effects. Please be aware that fixation might alter the staining pattern of EbbaBiolight.

### Materials:

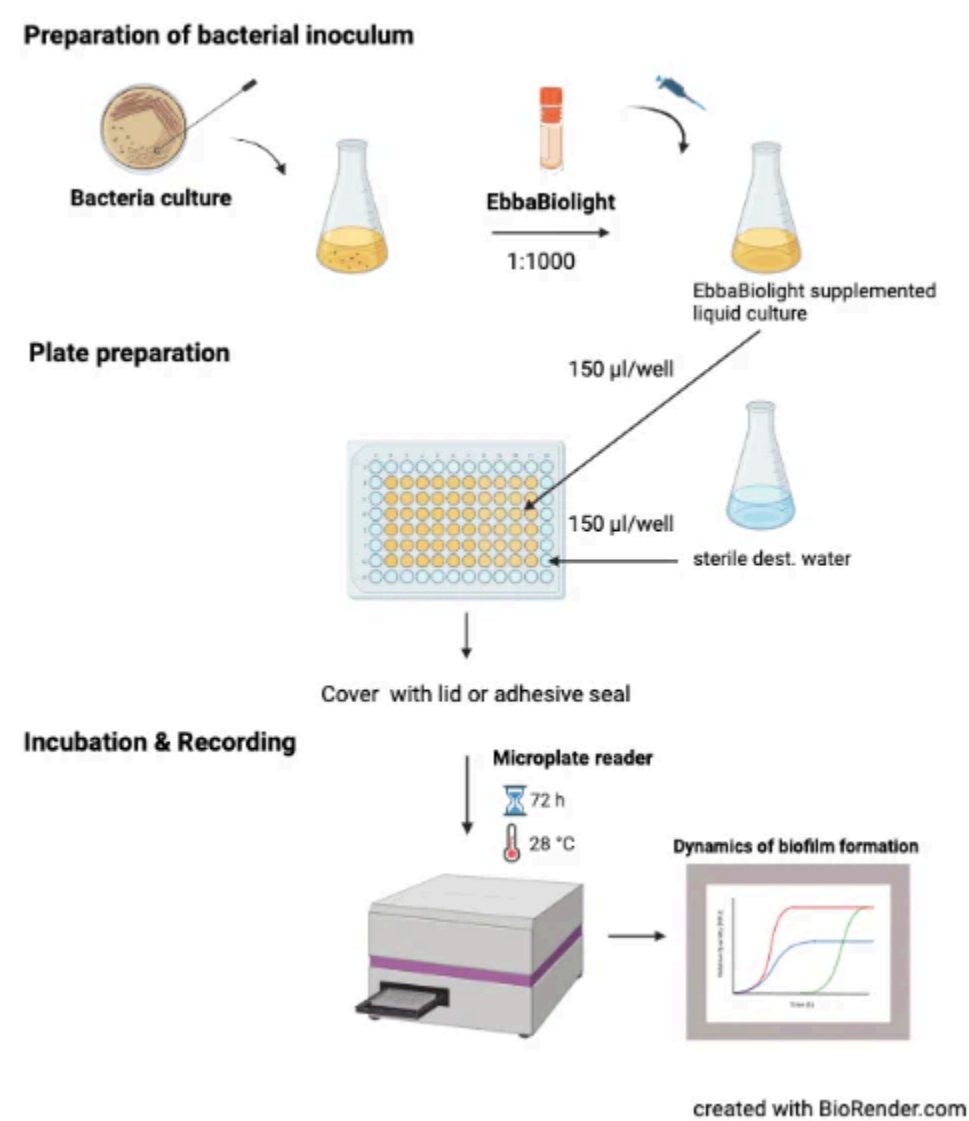
- EbbaBiolight
- Growth medium
- Bacteria on standard culture plate
- 96-well plate (round bottom) with cover
- Deionized water

### Equipment:

- Incubator (28°C)
- Shaking Incubator (37°C)
- Fluorescence plate reader

### Assay Procedure:

- Prepare bacterial inoculum:
  - Pick a colony from a standard culture plate.
  - Transfer colony to growth medium.
  - Prepare an overnight or exponential culture under continuous shaking at 37°C.
  - Dilute bacterial culture 1:100 in fresh growth medium.
  - Add EbbaBiolight (1:1000) and mix gently.
- Prepare 96-well plate:
  - Use 100-200  $\mu$ l liquid culture supplemented with EbbaBiolight per well.
  - Fill unused wells with sterile water to avoid drying during incubation.
  - Seal the plate with a cover.
- Incubation & Readout:
  - incubate the 96-well plate directly in the plate reader under static conditions.
  - Record excitation- and emission at regular time points during biofilm growth.



## Optotracing with EbbaBiolight

EbbaBiolight fluorescent tracer molecules are optotracers. Unlike conventional fluorescent dyes, optotracers bind promiscuously to a range of targets with repetitive motifs. EbbaBiolight has been shown to bind to curli and cellulose in *Salmonella* extracellular matrix<sup>[1,2]</sup>, peptidoglycan and lipoteichoic acids in the cell envelope of *Staphylococci*<sup>[3]</sup>,  $\beta$ -glucans from *S. cerevisiae* and Chitin in *C. albicans*<sup>[4]</sup>. Upon binding, the fluorescence intensity of the optotracer increases. This property makes it possible to use EbbaBiolight for live fluorescent tracking of microorganisms, without the need to wash away unbound molecules. It is possible to read out fluorescence intensity at the emission maximum ( $Em_{max}$ ) when excited at or close to the excitation maximum ( $Ex_{max}$ ). This is useful for microscopy or fluorescence spectroscopy when straight-forward data analysis is required. Yet, due to the unique properties of the optotracers, a unique optical fingerprint is produced reflecting the specific nature of the target (sample composition) and environment (pH, osmolarity, polarity of the medium). This means that depending on the specific properties of the sample,  $Ex_{max}$  or  $Em_{max}$  can shift, or the appearance of double peaks or shoulders might indicate binding to multiple targets. We therefore recommend acquiring fluorescence excitation and emission spectra whenever possible within experimental limitations. EbbaBiolight excitation- and emission spectra can be accessed [here](#).

**Table:** EbbaBiolight spectral properties with maximum excitation ( $Ex_{max}$ ) and emission ( $Em_{max}$ ) when bound and recommended range for acquisition of excitation- and emission spectra as well as recommended filter sets for microscopy.

	$Ex_{max}$	$Em_{max}$	Excitation spectrum (detect at $Em_{max}$ )	Emission spectrum (excite at $Ex_{max}$ )	Recommended filter-sets
<b>EbbaBiolight 480</b>	420 nm	480 nm	300 - 450 nm	450 - 800 nm	DAPI
<b>EbbaBiolight 520</b>	460 nm	520 nm	300 - 490 nm	490 - 800 nm	FITC, GFP
<b>EbbaBiolight 540</b>	480 nm	540 nm	300 - 510 nm	510 - 800 nm	FITC, GFP, YFP
<b>EbbaBiolight 630</b>	520 nm	630 nm	300 - 600 nm	550 - 800 nm	PI, Cy3, TxRed, mCherry, Cy3.5
<b>EbbaBiolight 680</b>	530 nm	680 nm	300 - 650 nm	660 - 800 nm	PI, mCherry, Cy3.5



### Read More:

1. Choong FX et al. (2016) Real-Time optotracing of curli and cellulose in live *Salmonella* biofilms using luminescent oligothiophenes. *npj Biofilms and Microbiomes*, 2, 16024
2. Choong FX et al. (2021) A semi high-throughput method for real-time monitoring of curli producing *Salmonella* biofilms on air-solid interfaces. *Biofilm*, 3, 100060
3. Butina K. et al. (2020) Optotracing for selective fluorescence-based detection, visualization and quantification of live *S. aureus* in real-time. *npj Biofilms and Microbiomes*, 6(1), 35
4. Kärkkäinen, E. et al. (2022) Optotracing for live selective fluorescence-based detection of *Candida albicans* biofilms. *Frontiers in Cellular and Infection Microbiology*, 12, 2235-2988

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