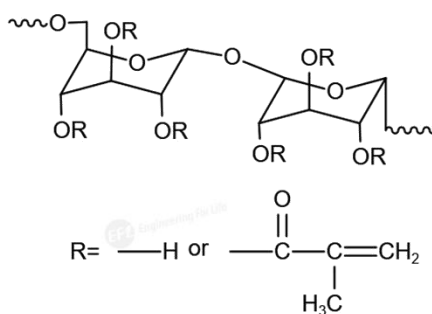


## Dextran Methacryloyl (DexMA)

### Product component

Item	Character	Package Size	Remark
A: DexMA	White spongy	1 g/bottle	Keep in dark
B: Photoinitiator LAP	White powder	0.05 g/bottle	

This instruction applies to EFL-DexMA



DexMA molecular structure

### Product introduction

Dextran methacryloyl (DexMA) is a double bond modified dextran, which can be crosslinked and solidified into a gel by UV and visible light under the action of photoinitiator. Due to the excellent water solubility, good biosafety and anti-nonspecific protein adsorption of DexMA, DexMA-based material systems have been widely used in biomedical fields, such as reducing thrombosis in blood vessels, reducing blood viscosity and drug delivery.

### Applications

Cell culture, biological 3D printing, tissue engineering, etc.

### Storage

**Dry kit:** room temperature, 3 months; 4°C, 12 months; -20°C, 18 months. **Sterile solution:** 4°C (in dark), 7 days; -20°C (in dark), 6 months. **Please note that repeated freezing and thawing of the solution will affect the performance of the product, so it is best to prepare it when using it.**

### Period of validity



企业微信公众号  
扫描右侧二维码  
获取更多信息

The date of manufacture is shown in the package.

## Solution preparation

### 1. Prepare 0.25% (w/v) standard solution of initiator

- (1) Add 20ml PBS into the brown bottle containing initiator LAP (containing 0.05g LAP);
- (2) Heat and dissolve the solution in a water bath at 40-50°C for 15 minutes, shaking several times.

The LAP standard solution can be stored for 12 months at 4°C in dark.

### 2. Prepare DexMA solution (5-15% (w/v) is recommended)

- (1) Take the required mass of DexMA into the centrifugal tube;
- (2) Add the initiator standard solution into the centrifuge tube;
- (3) Dissolve at room temperature for 30 minutes, oscillating several times during the period;
- (4) Sterilize the DexMA solution immediately with a 0.22μm sterile needle filter.

## Suggestions for 2D cell culture

- Injecting DexMA solution into the well plate immediately;  
(96-well plate: 50-100 μL/ well, 48-well plate: 100-300 μL/ well, 24-well plate: 300-500 μL/ well);
- Irradiate the wells with 405 nm light for 10-30 seconds to gelate, the gel strength can be adjusted by the time and intensity of the light;
- Add medium to the wells to cover the gel. Place the well plate in a 37°C incubator for 5 minutes. And then wash the sample and remove the medium;
- Add the cell suspension to the well plate. Change medium, observe, and photograph according to experimental design. (No special requirements for operation procedures).

## Suggestions for 3D cell culture

- Cells were collected and resuspended in DexMA to prepare the cell suspension;
- Add cell suspension into the well plates;  
(96-well plate: 50-100 μL/ well, 48-well plate: 100-300 μL/ well, 24-well plate: 300-500 μL/ well)
- Irradiate the wells with 405 nm light for 10-30 seconds to gelate, the gel strength can be adjusted by the time and intensity of the light;
- Add medium to the wells. Place the plate in a 37°C incubator for 5 minutes. And



企业微信公众号  
扫描右侧二维码  
获取更多信息

then wash the sample and remove the medium;

- Add fresh medium and incubate for a long time. Change medium, observe, and photograph according to experimental design. (No special requirements for operation procedures).

**Tips: Do not look directly at the light source.**



企业微信公众号  
扫描右侧二维码  
获取更多信息