



## 细胞爬片荧光探针原位杂交（FISH）+免疫荧光（IF）实验报告

### 1.实验器材及试剂

#### 1.1.实验器材

| 名称      | 厂家           | 型号                  |
|---------|--------------|---------------------|
| 盖玻片     | 江苏世泰实验器材有限公司 | 10212432C           |
| 无酶离心管   | Servicebio   | EP-150-M            |
| 摇床(钟摆式) | Servicebio   | TSY-B               |
| 涡旋混匀器   | Servicebio   | MV-100              |
| 移液枪     | Dragon       | KE0003087/KA0056573 |
| 免疫组化笔   | Servicebio   | G6100               |
| 冰箱      | 青岛海尔股份有限公司   | BCD-192TGN          |
| 正置荧光显微镜 | 日本尼康         | NIKON ECLIPSE TI-SR |
| 成像系统    | 日本尼康         | NIKON DS-U3         |
| 恒温箱     | LABOTERY     | GSP-70              |
| 高压灭菌锅   | 松下健康医疗       | MLS-3751L-PC        |

#### 1.2.主要实验试剂

| 试剂             | 厂家         | 货号      | 稀释比 |
|----------------|------------|---------|-----|
| DEPC           | Amresco    | E174    |     |
| 4%多聚甲醛（DEPC 水） | Servicebio | G1113   |     |
| PBS            | Servicebio | G0002   |     |
| 20×SSC 洗脱液     | Servicebio | G3016-4 |     |
| BSA            | Servicebio | G5001   |     |
| 蛋白酶 K          | Servicebio | G1234   |     |
| DAPI           | Servicebio | G1012   |     |
| 抗荧光淬灭封片剂       | Servicebio | G1401   |     |
| 杂交缓冲液          | Servicebio | G3016-3 |     |
| 一抗             | ⑦          |         |     |
| 二抗             | ⑧          |         |     |

### 2.细胞爬片荧光探针原位杂交+免疫荧光实验步骤

**2.1.细胞爬片固定：**细胞爬片置于 4%多聚甲醛（DEPC）固定 20min，于 PBS（PH7.4）中在脱色摇床上晃动洗涤 3 次，每次 5min。

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**2.2.消化:** 基因笔画圈, 根据不同组织不同指标特性, 滴加蛋白酶 K (20ug/ml) 消化②min。纯水冲洗后 PBS 洗 3 次×5min。.

**2.3.预杂交:** 滴加预杂交液 37°恒温箱 1h。

**2.4.杂交:** 待爬片完全冷却后, 滴加杂交液 (含探针 ③ 浓度 ④), ⑤度杂交过夜。

**2.5.杂交后洗涤:** 洗去杂交液, 2×SSC, 37°C 洗 10min, 1×SSC, 37°C 洗 2×5min, 0.5×SSC 37°洗 10min。若非特异杂交体较多, 可以增加甲酰胺洗涤。

**2.6.滴加封闭液:** 滴加封闭血清 ⑥。室温 30min。

**2.7.孵育一抗:** 滴加一抗 ⑦, PBS 稀释比 ⑨。4°过夜。后 PBS 洗 3×5min。

**2.8.孵育二抗:** 滴加相应二抗 ⑧, 室温孵育 50min。后 PBS 洗 3×5min。

**2.9.DAPI 复染核:** 切片滴加 DAPI 染液, 避光孵育 8min, 冲洗后滴加抗荧光淬灭封片剂封片。

**2.10.镜检拍照:** 切片于尼康正置荧光显微镜下观察并采集图像。(紫外激发波长 330-380nm, 发射波长 420nm, 发蓝光; FAM(488)绿光激发波长 465-495nm, 发射波长 515-555 nm, 发绿光; CY3 红光激发波长 510-560, 发射波长 590nm, 发红光。)

### 3.细胞爬片荧光探针原位杂交+免疫荧光实验结果判读

DAPI 染出来的细胞核在紫外的激发下为蓝色, 阳性表达为相应荧光素标记的荧光。FAM(488)为绿光, cy3 为红光。mRNA 原位杂交显示结果理论为胞浆阳性, 少数核阳性属正常。micRNA 与 lncRNA 不同指标表达定位不同。免疫荧光结果根据不同指标, 定位不同。根据表达量不同荧光亮度有强弱。

**注:** 上述涉及到的所有试剂, 仪器等在 RNA 原位杂交实验时都需使用 DEPC 处理后的 Rnase free 环境。

#### 附表 1 探针信息



## Cell Climbing–fluorescence Probe-FISH and Immunofluorescence Protocol

### 1. Apparatus and Reagents

#### 1.1 Apparatus

| Name                        | Producer   | Model               |
|-----------------------------|------------|---------------------|
| Coverslip                   | Citotest   | 10212432C           |
| Enzyme-free centrifuge tube | Servicebio | EP-150-M            |
| Shaker                      | Servicebio | TSY-B               |
| Vortex                      | Servicebio | MV-100              |
| Pipettor                    | Dragon     | KE0003087/KA0056573 |
| Liquid Blocker PAP Pen      | Servicebio | G6100               |
| Refrigerator                | Haier      | BCD-192TGN          |
| Microscopy                  | Nikon      | NIKON ECLIPSE CI    |
| Imaging system              | Nikon      | NIKON DS-U3         |
| Incubator                   | LABOTERY   | GSP-70              |
| Autoclave                   | PANASONIC  | MLS-3751L-PC        |

#### 1.2 Major reagents

| Reagent                                     | Manufacturer | Article Number |
|---------------------------------------------|--------------|----------------|
| DEPC                                        | Amresco      | E174           |
| 4% of paraformaldehyde (DEPC water)         | Servicebio   | G1113          |
| Ethanol                                     | SCRC         | 100092683      |
| PBS                                         | Servicebio   | G0002          |
| 20×SSC solution                             | Servicebio   | G3016-4        |
| BSA                                         | Servicebio   | G5001          |
| Proteinase K                                | Servicebio   | G1234          |
| DAPI                                        | Servicebio   | G1012          |
| Anti-fluorescence quenching sealing tablets | Servicebio   | G1401          |
| Hybridization buffer                        | Servicebio   | G3016-3        |
| First antibody                              |              |                |
| Second antibody                             |              |                |

### 2. The Steps of The Experiment

#### 2.1. Cell climbing fixation: cell climbing was fixed in 4% paraformaldehyde (DEPC) for 20 min,

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wash 3 times with PBS (pH 7.4) in a decolorizing shaker for 5 min each time.

2.2. **Digestion:** mark the objective tissue with liquid blocker pen, according to the characteristics of different tissues and different indicators, Add proteinase K (20 ug/ml) to cover tissues and incubate at 37°C for \_\_\_\_\_ minutes. wash in sterilized water, then washed three times in PBS, 5 min each.

2.3. **Prehybridization:** add hybridization buffer onto specimen and incubate at 37°C for 1h.

2.4. **Hybridization:** remove the pre-hybridization solution, add the \_\_\_\_\_ probe hybridization solution with concentration of \_\_\_\_\_, and incubate the section in a humidity chamber and hybridize overnight at \_\_\_\_\_°C.

2.5. **Washing:** remove the hybridization solution. Wash sections in 2×SSC for 5 min at 37°C , wash sections in 1×SSC two times for 5 min each at 37°C, and wash in 0.5×SSC for 10 min at room temperature. Formamide washing can be added if there are more non-specific hybrids.

2.6. **Blocking:** add blocking serum to the section and incubate at room temperature for 30 min.

2.7. **Incubate first antibody:** PBS solution containing a \_\_\_\_\_ dilution of primary antibody were added and incubated at 4°C overnight. Samples were then washed with PBS three times for 5 min each at RT.

2.8. **Incubate second antibody:** after washing, the section was incubated for 50 min with second antibody at RT. Samples were then washed with PBS three times for 5 min each at RT.

2.9. **Stain cell nuclei (counter stain):** incubate with DAPI for 8min in the dark, and then mounting.

2.10 **Microscopic examination and photography:** to take photos with positive fluorescence microscope. DAPI glows blue by UV excitation wavelength 330-380 nm and emission wavelength 420 nm; FAM glows green by excitation wavelength 465-495 nm and emission wavelength 515-555 nm; CY3 glows red by excitation wavelength 510-560 nm and emission wavelength 590 nm.

### 3. Interpretation of the results

The nuclear stained by DAPI were blue under ultraviolet excitation, and the positive expression was a kind of fluorescence labeled by corresponding luciferin. FAM (488) appears green, cy3 appears red. The results of mRNA in situ hybridization were cytoplasmic positive and a few nuclear positive were normal. MicRNA and lncRNA were expressed differently. According to the expression, Different fluorescence brightness is strong or weak.

Note: All reagents, instrument need RNase free.

Attached table 1 probe information.