

**石蜡切片 DAB 显色原位杂交 (CISH) 实验报告****1. 实验器材及试剂****1.1. 实验器材**

名称	厂家	型号
脱水机	DIAPATH	Donatello
载玻片	Wanwu	
盖玻片	江苏世泰实验器材有限公司	10212432C
病理切片机	上海莱卡仪器有限公司	RM2016
无酶离心管	Wanwu	EP-150-M
摇床(钟摆式)	Wanwu	TSY-B
涡旋混匀器	Wanwu	MX-F
移液枪	Dragon	KE0003087/KA0056573
Gene tech pen	Gene tech	GT1001
冰箱	青岛海尔股份有限公司	BCD-192TGN
显微镜	CIC	XSP-C204
正置光学显微镜	日本尼康	NIKON ECLIPSE CI
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
包埋机	武汉俊杰电子有限公司	JB-P5
冻台	武汉俊杰电子有限公司	JB-L5
组织摊片机	浙江省金华市科迪仪器设备有限公司	KD-P

**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)	Wanwu	G1113	
石蜡	Sakura		
无水乙醇	国药集团化学试剂有限公司	100092683	
二甲苯	国药集团化学试剂有限公司	10023418	
甲醇	国药集团化学试剂有限公司	40030561	
双氧水	国药集团化学试剂有限公司	10011208	
PBS 缓冲液 (DEPC)	Wanwu	G0020	
20×SSC 洗脱液	Wanwu	G3016-4	

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BSA	Wanwu	G5001
蛋白酶 K	Wanwu	G1205
苏木素染液	Wanwu	G1004
中性树胶	国药集团化学试剂有限公司	10004160
杂交缓冲液	Wanwu	G3016-3
anti-DIG-HRP	jackson	200-032-156
DAB 显色剂	Wanwu	G1211

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## 2.石蜡切片 DAB 显色原位杂交实验步骤

- 2.1. 组织固定:** 组织取出洗净后立即放入固定液 (DEPC 水配制) 中固定 12h 以上。
- 2.2. 脱水:** 组织固定完成后经梯度酒精脱水后浸蜡, 包埋。
- 2.3. 切片:** 石蜡经切片机切片, 摊片机捞片, 62°烤箱烤片 2h。
- 2.4. 石蜡切片脱蜡至水:** 依次将切片放入二甲苯I15min-二甲苯II15min-无水乙醇I5min-无水乙醇II5min, 风干, DEPC 水浸泡。
- 2.5. 消化:** 根据组织固定时间长短, 切片于修复液中煮沸 10-15 分钟, 自然冷却。后基因笔画圈, 根据不同组织不同指标特性, 滴加蛋白酶 K(20ug/ml) 37°消化\_\_\_min。纯水冲洗后 PBS 洗 3 次×5min。
- 2.6. 阻断内源性过氧化物酶:** 滴加 3%甲醇-H<sub>2</sub>O<sub>2</sub>, 室温避光孵育 15min, 将玻片置于 PBS (PH7.4) 中在脱色摇床上晃动洗涤 3 次, 每次 5min。
- 2.7. 预杂交:** 滴加预杂交液, 37°C 孵育 1h。
- 2.8. 杂交:** 倾去预杂交液, 滴加含探针\_\_\_杂交液, 浓度\_\_\_, 恒温箱\_\_\_度杂交过夜。
- 2.9. 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 室温洗 10min。若非特异杂交体较多, 可以增加甲酰胺洗涤。
- 2.10. 滴加封闭液:** 滴加封闭血清\_\_\_正常兔血清\_\_\_。室温 30min。
- 2.11. 滴加鼠抗地高辛标记过氧化物酶 (anti-DIG-HRP):** 倾去封闭液, 滴加 anti-DIG-HRP。4 度 孵育过夜, 后 PBS 洗 4 次×5min。
- 2.12. DAB 显色:** 切片稍甩干后, 在圈内滴加新鲜配制的 DAB 显色液, 显微镜下控制显色时间, 阳性为棕黄色, 纯水冲洗切片终止显色。
- 2.13. 复染细胞核:** Harris 苏木素复染 3min 左右, 自来水洗, 1%盐酸酒精分化数秒, 自来水洗, 氨水返蓝, 流水冲洗。
- 2.14. 脱水封片:** 将切片依次放入 75%酒精 6min-85%酒精 6min—100%酒精 I 6min—100%酒精 II 6min-正丁醇 6min—二甲苯透明, 将切片从二甲苯中拿出稍晾干后, 中性树胶封片。
- 2.15. 显微镜检, 图像采集分析。**

## 3.石蜡切片 DAB 显色原位杂交实验结果判读

苏木素染细胞核为蓝色, DAB 显出的阳性表达为棕黄色。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。根据表达量显色有深浅。

**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

**Paraffin -DIG -DAB -ISH protocol**
**1. Apparatus and reagents**
**1.1 Apparatus**

Name	Producer	Model
Dehydrator	WHJJ	JJ-12J
Paraffin embedding machine	WHJJ	JB-P5
Pathologic microtome	Leica	RM2016
Frozen flat	WHJJ	JB-L5
Water Bath-Slide	Kedee	KD-P
RNase-freeGlass microscope slides	Wanwu	
Micro-centrifuge	Wanwu	D1008E
Rocker	Wanwu	TSY-B
Vortex	Dragon	MX-F
Pipettor	CIC	KE0003087/KA0056573
Microscopy	Gene tech	XSP-C204
Liquid blocker pen	HAIER	GT1001
Refrigerator	LABOTERY	BCD-192TGN
incubator		GSP-70
autoclave	PANASONIC	MLS-3751L-PC

**1.2 Major reagents**

reagent	manufacturer	article number
Ethanol	SCRC	100092683
Xylene	SCRC	10023418
PBS solution	Wanwu	G0002
Proteinase K	Wanwu	G1205
4% of paraformaldehyde (DEPC water)	Wanwu	G1113
Methanol	SCRC	
Hydrogen peroxide	SCRC	
DEPC	Amresco	E174
20×SSC solution	Wanwu	G3016-4
BSA	Wanwu	G5001
Hematoxylin dye	Wanwu	G1004
neutral balsam	SCRC	
hybridization buffer	Wanwu	G3016-3

 网址：<http://www.wanwusw.com>

 细胞网址：<http://www.hfwanwu.com>

地址：合肥市蜀山区长江西路248号11层

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邮箱：2028438226@qq.com

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anti-DIG-HRP	Jackson	
DAB reagent	Wanwu	G1211

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## 2. The steps of the experiment

2.1 **Organization fixed:** Take out the organization, Wash clean, then Immediately put in the fixed fluid (DEPC) to fix 2-12h.

2.2 **Dehydration:** The tissue is dehydrated by gradient alcohol, paraffin, Embedding.

2.3 **Section:** The paraffin is sliced through the slicer, the piece of the slice machine and the 62 - degree oven roast for 2 hours.

2.4 **Dewaxing and dehydration:** Soak sections in 2 changes of xylene, 15 minutes each. Dehydrate in 2 changes of pure ethanol for 5 minutes each. Then, followed respectively by dehydrating in gradient ethanol of 85% and 75% ethanol 5 minutes each. Wash in DEPC dilution.

2.5: **Digestion:** According to the tissue fixation time, the slices are boiled in the retrieval solution for 10-15 minutes and naturally cooled. Mark the objective tissue with liquid blocker pen, according to the characteristics of tissues, Add proteinase K(20ug/ml) working solution to cover objectives and incubate at 37°C for \_\_\_min. Washing with pure water, then wash three times with PBS (pH 7.4) in a Rocker device, 5 min each.

2.6 **Block endogenous peroxidase:** add 3% methanol-H<sub>2</sub>O<sub>2</sub> , incubate in dark at room temperature for 15min. Wash slides in PBS(PH7.4) three times for 3 min each, with gentle agitation and protected from light.

2.7 **Pre-hybridization:** Add Pre-hybridization solution to each section and incubate for 1 h at 37°C.

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

2.9 **Washing:** Remove the hybridization solution. Wash sections with 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.

2.10 **Blocking:** add blocking solution (Rabbit serum) to the section and incubate at room temperature for 30 min.

2.11 Add the mouse anti-digoxigenin-labeled peroxidase (anti-DIG-HRP): Remove the blocking solution and add anti-DIG-HRP. Incubate at 37 °C for 40 min, then Wash sections in PBS four times for 5 min each.

2.12 **DAB developing:** dry sections slightly, add fresh prepared DAB chromogenic reagent to marked tissue. Manage reaction time by observing in microscopy until positive expression appears brown-yellow. Then stop developing reaction by wash in running tap water.

2.13 Counterstain in nucleus with Hematoxylin staining solution for 3 min and wash in tap water.

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Treat with the differentiate solution for a few seconds, wash in running tap water. Back to blue by bluing solution, wash in running tap water.

2.14 Dehydrate successively in gradient ethanol of 75%, 85%, and 2 changes of pure ethanol, respectively, 6 min each. Clear in xylene for 6 min and mount with resin mounting medium.

2.15 Image the results using a brightfield fluorescence microscope. Store slides at RT.

### **3. Interpretation of the results**

The nucleus stained with hematoxylin are blue, and the positive expression of DAB was brownish yellow. The results of in situ hybridization of mRNA were cytoplasmic positive and a few nuclear positive were normal. MicRNA and lncRNA were expressed differently. According to the expression of color depth.

Note: All reagents, instrument need RNase free.

**Attached table 1 probe information.**