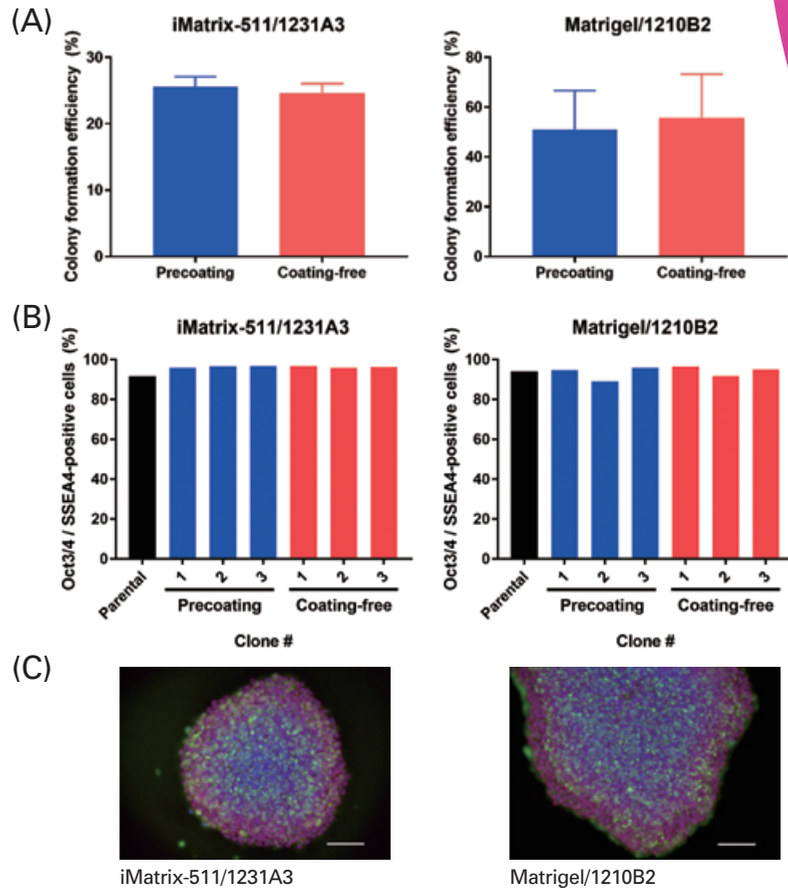
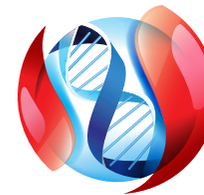


● Cloning efficiency with coating-free protocol and pluripotency analysis of isolated hiPSC clones



1231A3 and 1210B2 (Human episomal iPSC lines established by CiRA) were cloned on iMatrix-511 or Matrigel[®] by the precoating or coating-free method.
(A) Comparison of the cloning efficiency of hiPSCs by two different methods. Bars represent the means \pm S.D. (n=3). (B, C) Analysis of pluripotent markers in isolated hiPSC clones. Expression levels of pluripotent markers were evaluated by (B) FACS and (C) ICC (Blue: DNA, Green: Tra1-60, Red: Oct3/4). Scale bars: 100 μ m.

Feeder-free medium for ES/iPS cells



StemFit[®] Technical tips

Key Points for *single-cell cloning* with *coating-free method*

Benefit 1

Superior colony-forming efficiency

Enables efficient single-cell cloning

Benefit 2

Coating-free protocol

No coating process, No incubation

Eat Well, Live Well.



For further information, please contact

✉ stemfit@ajinomoto.com

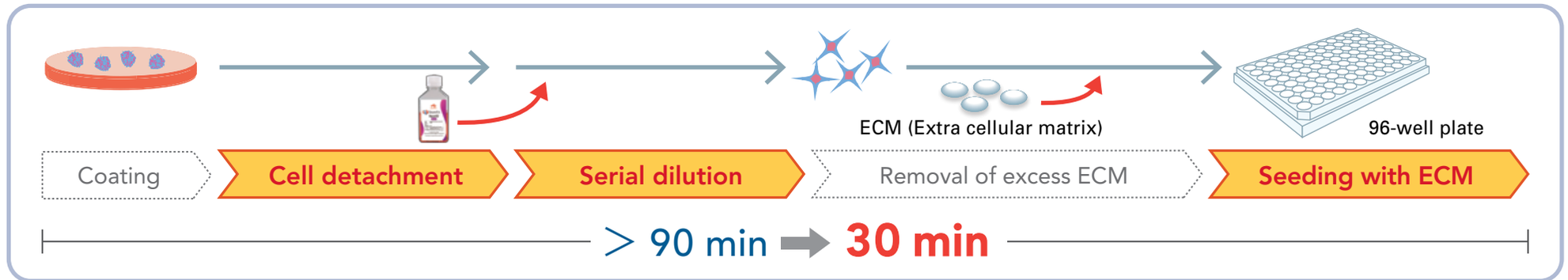
AJINOMOTO CO., INC. AminoScience Division

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Eat Well, Live Well.



Brief protocol for single-cell cloning by the coating-free method



1 Detach cells and resuspend in culture medium supplemented with 10 μ M Y-27632.

** Also see our Technical tips: Key points for successful single-cell passage*

2 Prepare 10 mL of 10 cell/mL cell suspension by serial dilution with culture medium with 10 μ M Y-27632.

3 Add iMatrix-511 or Matrigel® to the prepared cell suspension and mix thoroughly.

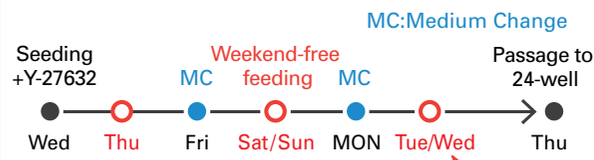
Point-1
Concentration of ECMs !

ECM	Amount	Final conc.
iMatrix-511 (0.5 mg/ml)	35 μ l	1.75 μ g/ml
Matrigel®	100 μ l (x1/100)	10 μ l/ml

4 Plate 100 μ L (= 1 cell) in each well of the 96-well plate immediately.

5 Replace medium with fresh culture medium without Y-27632 at least every three days. Around day 8, select single colonies to be passaged to a 24-well plate.

<Medium Change Schedule Example>



Point-2
Extend or shorten culture period for colonies to be appropriate sizes

6 After washing the colonies with 100 μ l of PBS, detach the cells with 50 μ l of cell detaching solution and incubate at 37 °C for 10 min.

** Accutase or TrypLE™ can be used*
** Incubation times may vary*

7 Carefully remove cell detaching solution.

Point-3
Remove very carefully because the colonies easily detach

8 Dissociate colonies by pipetting with 100 μ l of culture medium with 10 μ M Y-27632. Transfer resuspended cells to ECM-coated 24-well plate with 400 μ l of culture medium containing 10 μ M Y-27632 immediately.

Point-4
Detach the colonies one by one as reattachment can occur soon after adding the culture medium

9 Change the medium to fresh culture medium without Y-27632 at least every three days.