

CSI094Fe01

Primary Feline Splenic Fibroblasts (SF)

Organism Species: Felis silvestris catus (Feline)

Instruction manual

FOR RESEARCH USE ONLY
NOT FOR USE IN CLINICAL DIAGNOSTIC PROCEDURES

1st Edition (Revised in May, 2025)

[DESCRIPTION]

Cell Type: Fibroblasts

Synonyms: SF Strain: Feline Age: 3-4 Weeks

Tissue Source: Spleen

Disease: Normal **Size:** >5×10⁵cell/vial

[PROPERTIES]

Cell activity: >85% (Viability by Trypan Blue Exclusion).

Formulation: Frozen 1 mL or T25 flask.

Biosafety: Negative for HIV-1, HBV, HCV, mycoplasma, bacteria, yeast and fungi.

Applications: For research use only. It is not approved for human or animal use, or for application in

clinical diagnostic procedures. **Growth Properties:** Adherent

[CONTENTS]

Form & Buffer: Supplied as solution form in frozen stock solution, containing 90% FBS+10% DMSO.

[USAGE]

Upon receiving the cells in a T-25 flask at room temperature, immediately transfer the cells to 37°C, 5% CO₂ incubator; the cells in vials, directly and immediately transfer the cells from dry ice to liquid nitrogen.

Culture conditions:

DMEM+10%FBS+1% Fibroblasts growth supplement+1%Penicillin-Streptomycin Solution

Temperature: 37°C

Condition: 95% air, 5% carbon dioxide

Cell recovery:

After receiving the cells, shake at 37°C in a water bath until completely dissolved, transfer to a 15 ml centrifuge tube, add 3-5 times complete culture solution, 1000 rpm for 5 min, discard the supernatant, and place in a T25 flask for culture.

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Cell passage:

- Cell passage when cell growth at 85-95%.
- Discard the medium and wash with PBS 1-2 times.
- 3. Add 1 ml of Trypsin at 37°C, observe the cell under the microscope. If the cells are retracted and rounded, pat the culture flask to let the cells fall off. Stop digestion by adding 2 ml of complete medium containing 10% serum. Make it a single cell suspension.
- Add the fresh medium to resuspend the cells. Unless otherwise stated, the recommended ratio of primary cells is 1/2.

[Shipping]

Dry ice.

[STORAGE]

Upon receiving, directly and immediately transfer the cells from dry ice to liquid nitrogen and keep the cells in liquid nitrogen until they are needed for experiments.

[IMPORTANT NOTE]

The cell is for research use only, and we will not be responsible for any issue if the cell was used in clinical diagnostic or any other procedures.

[Figure]

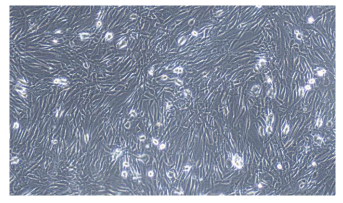


Figure 2

Figure 1

Figure 1 Morphology of Primary Feline Splenic Fibroblasts (Optical microscope, ×100)

Figure 2 Morphology of Primary Feline Splenic Fibroblasts (Optical microscope, ×200)

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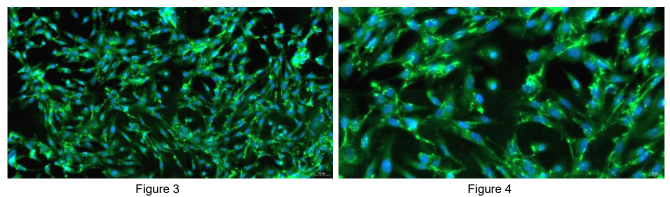


Figure 3 Immunofluorescence identification of Fibronectin specific antibody (×200)

Figure 4 Immunofluorescence identification of Fibronectin specific antibody (×400)

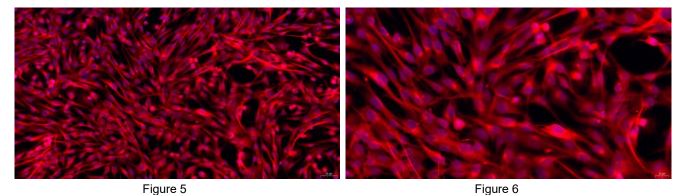


Figure 5 Immunofluorescence identification of Vimentin specific antibody (×200)

Figure 6 Immunofluorescence identification of Vimentin specific antibody (×400)