





A Geno Technology, Inc. (USA) brand name

FOCUS™ Mitochondria

(Cat. # 786-022)



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INTRODUCTION

FOCUS[™] Mitochondria kit enables the fast and easy isolation of enriched mitochondrial fractions from animal cells and tissues. The majority of the isolated mitochondria obtained from this kit contain intact inner and outer membranes. This kit contains reagents for processing 50-80 preparations of 20 million cultured mammalian cells or 20-30 preps of 50-100mg tissue. The number of preparations varies depending on the protocol, preparation size and cell or tissue type.

ITEM(S) SUPPLIED (Cat. #: 786-022)

Description	Size
SubCell Buffer-I	60ml
SubCell Buffer-II [3X]	30ml
SubCell Buffer-IV	25ml
SubCell Buffer-V	15ml
Mitochondria Storage Buffer	10ml
Mitochondria Storage Component	1 vial

STORAGE CONDITION

The kit is shipped at ambient temperature. After receiving store the kit components at 4°C except Mitochondria Storage Component at -20°C. The kit is stable for one year when stored unopened. Use aseptic techniques when handling the reagent solutions.

ADDITIONAL ITEMS REQUIRED

Syringes and 20 gauge needles or Wheaton Dounce homogenizer, centrifuge and centrifuge tubes. Optional reagents: Delipidated BSA, Trypsin, PBS and protease inhibitor cocktail.

PREPARATION BEFORE USE

All buffers should be kept ice cold. Dilute appropriate volume of 3X SubCell Buffer-II to 1X with SubCell Buffer-I as needed (e.g. mix 2ml SubCell Buffer-I with 1ml SubCell Buffer-II). All centrifugation steps should be performed at 4°C.

Preparation of Working Mitochondria Storage Buffer

Pipette 0.5ml Mitochondria Storage Buffer to Mitochondria Storage Component vial. Pipette up and down a few times to dissolve all components completely. Transfer the solution of Mitochondria Storage Component to Mitochondria Storage Buffer bottle and mix well. The Working Mitochondria Storage Buffer should be kept frozen for long-term use.

PROTOCOLS

A. Isolation of Mitochondria from Animal Cells.

This protocol is for processing $20x10^6$ cells (or ~100 μ l wet cell pellet). It can be scaled up and down accordingly.

OPTIONAL: Add appropriate protease inhibitor cocktail (e.g. G-Biosciences' Protease Arrest, Cat# 786-108) to SubCell Buffer-I just before use.

- 1. Use fresh cells only. Pellet the harvested cells by centrifugation at \sim 800 x g for 1 minute. Carefully remove and discard the supernatant.
 - **OPTIONAL:** Wash the cell pellet with 1ml ice cold PBS, centrifuge it as above and discard the supernatant.
- Add 500µl of ice cold SubCell Buffer-I. Gently vortex to suspend the cells and incubate on ice for 10 minutes.
- 3. Perform this lysis step on ice. Using a narrow opening (20 gauge) syringe needle, gently pull the suspension up and down 10-30 times. Alternatively, transfer cell suspension to ice cold Dounce homogenizer. Homogenize the cells on ice using tight pestle. Perform 5 to 20 strokes to lyse the cells effectively. Transfer the lysate to a microcentrifuge tube. Rinse Dounce homogenizer with 200µl of SubCell Bufferland pool together. Invert the tube several times to mix.
 - **NOTE:** To check the cell lysis efficiency, spot 5µl of cell lysate onto a glass slide, add coverslip and view under a phase-contrast microscope. Pulling times or strokes in the above lysis step are only guidelines. Mechanical force to lyse cells depends on cell types, the total number of the cells and hands on experience. Insufficient force will not lyse all the cells, but will achieve cleaner mitochondrial fractions with less nuclear contamination. Excess force may damage some nuclei, but high yield mitochondria fractions will be obtained with some contamination from nuclei.
- 4. Add 250μl 3X SubCell Buffer-II (350μl if Dounce homogenizer is used) and mix by inverting. This generates a 1X final concentration of SubCell Buffer-II.
- 5. Centrifuge the tube at 700x g for 10 minutes to pellet the nuclei. Transfer the supernatant to a new tube.
- Centrifuge supernatant at 12,000x g for 15 minutes. The pellet contains mitochondria.
 - **NOTE:** To fractionate light and heavy mitochondria, and obtain more purified mitochondrial fractions, see Section B. For a crude mitochondrial fraction, continue with step 8.
- 7. Transfer the supernatant (cytosol fraction) to a new tube. Add 500μ l 1X SubCell Buffer-II to the pellet, and centrifuge again at $12,000 \times g$ for 5 minutes. Discard the supernatant.
- 8. Suspend the mitochondrial pellet in 50-100µl Working Mitochondria Storage Buffer and keep the suspension on ice before downstream processing. The suspension

may be stored on ice for up to 48 hours. Freezing and thawing may compromise mitochondria integrity.

B. Fractionation of light and heavy mitochondria by gradient cushion

- 1. Suspend the mitochondrial pellet in 100µl 1X SubCell Buffer-II.
- 2. Make a step gradient by adding 200 μ l SubCell Buffer-V to a centrifuge tube and then overlaying with 200 μ l SubCell Buffer-IV. Gently float the mitochondrial suspension on the surface of the step gradient.
- Centrifuge the gradient at 20,000x g for 20 minutes. Observe the two white bands.
 The band at the interface of SubCell Buffer-IV and V is the heavy mitochondria fraction, whereas the band above the heavy mitochondria band is the light mitochondria fraction.
- 4. Carefully remove each band to a separate tube. Dilute the mitochondrial suspensions with equal volume of 1X SubCell Buffer-II. Centrifuge the tubes at 12,000x g for 15 minutes and discard the supernatant.
- Suspend the mitochondrial pellets in 30-50μl Working Mitochondria Storage Buffer and keep the suspensions on ice before downstream processing. The suspensions may be stored on ice up to 48 hours. Freezing and thawing may compromise mitochondrial integrity.

C. Isolation of mitochondria from soft tissues (liver or brain)

OPTIONAL: Delipidated BSA can be added to 1X SubCell Buffer-II to the concentration of 2mg/ml for removing free fatty acids prior processing. An appropriate amount of protease inhibitor cocktail also can be added to the 1X SubCell Buffer-II just before use.

- 1. Use a fresh tissue sample (obtained within one hour of sacrifice) kept on ice. <u>Do not freeze.</u>
- 2. Weigh approximately 50-100mg tissue. On a cooled glass plate, with the aid of a scalpel, mince the tissue into very small pieces.
- 3. Perform this step on ice. Transfer the minced tissue to an ice-cold Dounce tissue homogenizer. Add 10 volumes of 1X SubCell Buffer-II and using a loose-fitting pestle disaggregate the tissue with 5-10 strokes or until the tissue sample is completely homogenized. Using a tight-fitting pestle, release the nuclei with 8-10 strokes. Do not twist the pestle as nuclei shearing may occur.
- 4. Stand on ice for 2 minutes. Transfer the homogenate to a centrifuge tube and leave large chunks of tissue fragments in the homogenizer to be discarded. Centrifuge the lysate at 700x g for 5 minutes to pellet the nuclei.
- 5. Carefully transfer the supernatant into a new tube. Centrifuge supernatant at $12,000x \ q$ for 10 minutes.
- Remove the supernatant and resuspend the pellet in 10 volumes of 1X SubCell Buffer-II without BSA
- Centrifuge as in step 4. Repeat step 5 and remove the supernatant. The pellet contains mitochondria.

- **NOTE:** To fractionate light and heavy mitochondria, and obtain more purified mitochondrial fractions, see Section B.
- 8. Suspend the mitochondrial pellet in Working Mitochondria Storage Buffer (approximately 50μl for pellet from ~100mg tissue) and keep the suspension on ice before downstream processing. The suspension may be stored on ice for up to 48 hours. Freezing and thawing may compromise mitochondrial integrity.

D. Isolation of mitochondria from hard tissues (skeletal or heart muscle)

NOTE: For facilitating homogenization of the hard tissue, 0.25mg/ml Trypsin should be added to 1X SubCell Buffer-II. A concentrated BSA solution is needed to quench the proteolytic reaction after Trypsin treatment.

- Use a fresh tissue sample (obtained within one hour of sacrifice) kept on ice. <u>Do</u> not freeze.
- Weigh approximately 50-100mg tissues. On a cooled glass plate, with the aid of a scalpel, mince the tissue into very small pieces.
- Suspend the sample with 8 volumes of 1X SubCell Buffer-II containing 0.25mg/ml trypsin in a 2ml centrifuge tube.
- 4. Incubate on ice for 3 minutes and then spin down the tissue for a few seconds in the centrifuge.
- 5. Remove the supernatant by aspiration and add 8 volumes of 1X SubCell Buffer-II containing 0.25mg/ml Trypsin. Incubate on ice for 20 minutes.
- 6. Add BSA Solution to a final concentration of 10mg/ml and mix. Spin down the tissue at 1,000 x g for 5-10 seconds in the centrifuge.
- Remove the supernatant by aspiration. Wash the pellet with 8 volumes of 1X
 SubCell Buffer-II without Trypsin, and spin down the tissue for a few seconds in the centrifuge.
- Remove the supernatant by aspiration and add 8 volumes of the 1X SubCell Buffer-II without Trypsin.
- 9. Transfer the suspension to an ice-cold Dounce tissue homogenizer and using a loose-fitting pestle disaggregate the tissue with 5-15 strokes or until the tissue sample is completely homogenized. Using a tight-fitting pestle, release the nuclei with 8-10 strokes. Do not twist the pestle as nuclei shearing may occur.
- 10. Stand on ice for 2 minutes. Transfer the homogenate to a centrifuge tube and leave large chunks of tissue in the homogenizer to be discarded. Centrifuge the lysate at $700 \times q$ for 5 minutes to pellet nuclei.
- 11. Transfer the supernatant to a new tube. Centrifuge it at 12,000xg for 10 minutes and remove the supernatant. The pellet contains mitochondria.
 - **NOTE:** To fractionate light and heavy mitochondria, and obtain more purified mitochondrial fractions, see Section B.
- 12. Suspend the mitochondrial pellet in Working Mitochondria Storage Buffer (approximately 50μl for pellet from ~100mg tissue) and keep the suspension on ice

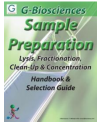
before downstream processing. The suspension may be stored on ice for up to 48 hours. Freezing and thawing may compromise mitochondrial integrity.

SOLUBILIZATION OF MITOCHONDRIAL FRACTIONS:

The fractionated mitochondria may be solubilized in any suitable buffer consistent with downstream procedures. For IEF/2D gel electrophoresis, the mitochondria fractions may be solubilized in a chaotropic extraction buffers. G-Biosciences offers a wide selection of buffers and reagents for IEF/2D gel electrophoresis. FOCUS/Extraction Buffer-VI (Cat. # 786-233) is suitable for solubilization of mitochondria fractions.

RELATED PRODUCTS

Download our Sample Preparation Handbook.



http://info.gbiosciences.com/complete-protein-sample-preparation-handbook/
For other related products, visit our website at www.GBiosciences.com or contact us.

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