



分子イメージングセミナー RIKEN Molecular Imaging Seminar

生物試料超薄切の課題および電子顕微鏡・ウルトラマイクロトームの関連技術紹介

日時 2018年4月19日(木) 13:00~17:30
場所 理化学研究所 生命機能科学研究センター
大会議室 (神戸 MI R&D センター 2F)
神戸市中央区港島南町 6-7-3

アクセス <http://www.clst.riken.jp/access.html>

言語 日本語、英語

スケジュール

13:00-13:15 生物試料超薄切の課題
講演者: 久米 慧嗣 (理研 BDR、理研 RC)
13:15-13:25 電子顕微鏡関連製品の紹介
講演者: 工藤 大樹 (日新イーエム株式会社)
13:25-14:15 バイオ分野における超薄切片法
講演者: Helmut Gnaegi (DiATOME)
14:15-14:30 ダイヤモンド単結晶の製造技術とその応用
講演者: 藤森直治 (株式会社イーディーピー)
14:30-15:00 ブレーク & ディスカッション
15:00-17:30 ウルトラマイクロトーム実技講習 (DiATOME) ※

※希望者5名程度で実施予定。希望者多数の場合には事務局で選考を行います。

主催 理化学研究所 生命機能科学研究センター
共催 理研エンジニアリングネットワーク、健康“生き活き”羅針盤リサーチコンプレックス

セミナー参加の事前申込が必要です。以下の参加登録フォームよりご登録ください。

<https://goo.gl/jyPkz7>

お問い合わせ: 理化学研究所 生命機能科学研究センター
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分子イメージングセミナー RIKEN Molecular Imaging Seminar

Issues of Ultra-Thin Sectioning for Biological Samples and Introduction of Technologies and Products Related to Electron Microscope and Ultramicrotome

Date&Time 13:00~17:30 on 19th April 2018 (Thu.)
Venue Conference Room, RIKEN Center for Biosystems Dynamics Research
6-7-3 Minatojima-minamimachi, Chuo-ku, Kobe
Access <http://www.clst.riken.jp/en/about/access/>
Language Japanese, English

Schedule

13:00-13:15 Issues of ultra-thin sectioning for biological samples
Speaker : Satoshi Kume (Research scientist, RIKEN BDR, RIKEN RC)
13:15-13:25 Introduction of products related to electron microscope
Speaker : Hiroki Kudo (Nisshin EM Co., Ltd.)
13:25-14:15 Ultramicrotomy in biology (including serial sectioning)
Speaker : Helmut Gnaegi (Managing Director, DiATOME)
14:15-14:30 Manufacturing of artificial single crystal diamond and its application
Speaker : Naoji Fujimori (CEO, EDP corp.)
14:30-15:00 Break & Discussion
15:00-17:30 Practical training course for Ultramicrotome (DiATOME) ※

※We plan to implement it with about 5 participants. In the case of a large number of applicants, we will make a selection at the secretariat.

Organizer RIKEN Center for Biosystems Dynamics Research
Co-organizer RIKEN Engineering Network Project, The “Compass to Healthy Life”
Research Complex program

Advance application for seminar participation is necessary. Please register from the below form.

<https://goo.gl/jyPkz7>

Inquiry Laboratory for Cellular Function Imaging,
RIKEN Center for Biosystems Dynamics Research
(TEL:078-304-7160, E-mail: riken-cfi-en-seminar@ml.riken.jp)

Special lecturer

Helmut Gnaegi (Managing Director, DiATOME)

Ultramicrotomy in Biology



Abstract

Serial ultrathin sectioning for 3D reconstruction gains increasing interest in the life science community. Sample trimming, sectioning, pick-up on TEM grids, glass slides or silicon wafers is presented. Sections as thin as 30nm are requested to get better resolution in TEM or SEM. The compression is a major obstacle to achieve such thin sections. The thinner the sections, the higher the compression. A nominal setting of 30nm at the ultramicrotome results in sections with a thickness of approx. 50-60nm. The compression factor also depends on the resin type. We discuss the compression and how to reduce or avoid.

Cryo-ultramicrotomy is used to obtain ultrathin cryo-sections from cryo-fixed or aldehyde-fixed cryo-protected samples. Samples which were soaked in 2.3M sucrose over night are removed from the sucrose, cut into small pieces and placed on the aluminium pins of the UC7/FC7 cryochamber. The pins with the samples are immersed in liquid nitrogen, then transferred to the cryochamber. Trimming is performed preferably with the so-called trim diamond blades. The size of the sample block depends on the desired section thickness. Good trimming results in long section ribbons. Section ribbons are cut with a cryo 35° diamond knife. For both the trimming and the sectioning the use of an antistatic device is mandatory. The ribbons are picked up with the aid of a loop and a sucrose/methyl cellulose droplet, then placed onto the grids at room temperature. The use of sucrose/methyl-cellulose led to a better structural preservation (W. Liou et al., *Histochem. Cell Biol.* 1996). Excellent structure preservation is mandatory for precise localisation of proteins.

Cryo-ultramicrotomy of frozen hydrated samples is performed at 120K. The sample preparation by high pressure freezing, cryoultramicrotomy, cryo-transfer and cryo-TEM preserve living matter close to the native state (A. Al-Amoudi et al., *EMBO Journal* 2004). Ultrathin cryosections serve for high resolution imaging and 3D reconstruction (C. E. Hsieh et al., *Journal of Structural Biology* 2006. A. Al-Amoudi et al., *Nature* 2007). Special 25° and 35° cryo diamond knives allow the cutting of cryo-sections with reduced compression. A crucial part in cryo-sectioning is the section pick-up. An ionizer/charger eliminates electrostatic charging in the cryo-chamber and improves the gliding of the cryo-sections (J. Pierson et al., *Journal of Structural Biology* 2010). In addition the ionizer/charger electrostatically fixes cryo-sections on the carbon film of the grid. A crucial part is the section pick-up. A micromanipulator system composed by two parts was developed (Studer et al., *Journal of Structural Biology* 2014) for pulling the section ribbons and holding the grid. A holy carbon film keeps the sections flat during tilting and imaging (J. Quispe et al., *Microscopy & Microanalysis* 2007). An obstacle for successful electron tomography of vitreous sections (TOVIS) is the compression and the crevasses. Possible scenarios for reducing these artifacts are discussed.