



NIBB CENTER OF THE INTER-UNIVERSITY BIO-BACKUP PROJECT (IBBP CENTER)

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In order to realize a life sciences community resilient to natural disasters and calamities, the National Institutes for Natural Sciences (NINS) and Hokkaido University, Tohoku University, University of Tokyo, Nagoya University, Kyoto University, Osaka University, and Kyushu University concluded an agreement on June 1st 2012 to launch a system to 'back up' the biological resources essential to the work being done at universities and research institutions nationwide, called the 'Interuniversity Bio-Backup Project (IBBP)'.

The IBBP Center was established as a centralized backup and storage facility at NIBB, while IBBP member universities set up satellite hubs and work closely with the IBBP center to put in place reciprocal systems for backing up important biological resources that have been developed by researchers residing in the area each university satellite hub is responsible for.



Figure 1. IBBP Center



Figure 2. Cryogenic storage system. Liquid nitrogen tanks are monitored 24 hours a day and are refilled

The IBBP Center includes earthquake proof structures capable of withstanding even very large scale quakes (equipped with emergency backup power generators), cryopreservation facilities equipped with automatic liquid nitrogen feeding systems, deep freezers, and refrigerated storage (mainly for seed stocks), as well as all manner of automated laboratory equipment, cell culture tools, and the latest equipment necessary to back up the genetic resources in a collaborative manner. The specific methods of preservation are freezing of the sperm and eggs of animals, cultured plant and animal cells, and gene libraries. Plant seeds are frozen or refrigerated.

University satellite hubs receive preservation requests of biological resources from researchers and report to the Managing Project Committee of IBBP (constituted of faculty of NIBB and satellite institutes), where the relevance of the request is reviewed. When the request is sustained, biological resources to be preserved will be sent to the IBBP Center by the requesting researcher, where they will be frozen (or refrigerated) and their particular information will be registered into a database. In the event of a disaster leading to the loss of a researcher's own biological resources, preserved samples will be promptly supplied back to the researcher so they can quickly resume their work.

Through the development of this backup system biological resources that had only been stored at individual research institutes can now be preserved at the state of the art facilities of the IBBP Center, and Japan's research infrastructure has been significantly strengthened.

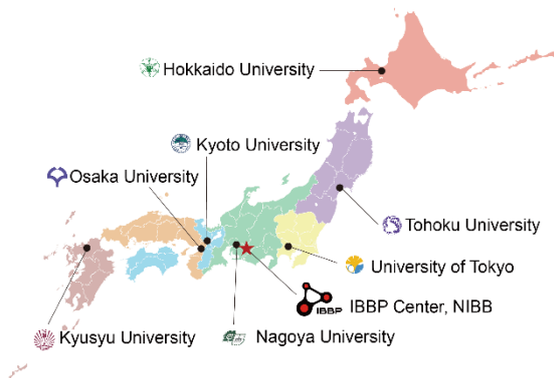


Figure 3. IBBP Satellite Hub

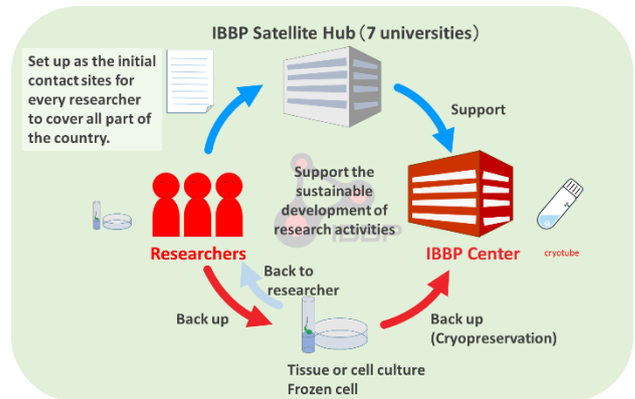


Figure 4. Backup system (IBBP)

Current status of back up for the biological resources

In 2013, IBBP Center stored 3,598 384-well plates consisting of 1,381,632 clones as cDNA/BACs clones, 1013 tubes for plant and animal samples and 618 tubes for microorganisms.

Collaborative Research Project for the development of new long-term storage technologies and cryo-biological study



As the IBBP Center can only accept biological resources which can be cryopreserved in liquid nitrogen, researchers cannot backup those biological resources for which cryopreservation methods are not well established, to increase the usability of IBBP, we started a collaborative research project for the development of new long-term storage technologies and cryo-biological study in 2013. This collaborative research includes two subjects 1) Establishment of new storage technology for biological resources for which long-term storage is unavailable. 2) Basic cryobiological research enabling us to improve low temperature storage for biological resources. In 2013 we had eleven applications and accepted nine proposals. We are also working to establish a research center for cryo-biological study through this Collaborative Research Project.

Research activity by D. Tanaka

Assistant Professor: *TANAKA, Daisuke*
Technical Assistant: *AKIMOTO-KATO, Ai*

Vitrification-based protocols are known to be effective for long-term, stable preservation of plant germplasm; this protocol can reduce the cost and manpower for maintaining a large number of germplasm lines and keep many valuable genetic lines for a long term under genetically stable conditions. However, it is still not widely employed as a reliable long-term preservation protocol due to the lack of basic knowledge on the cellular and water behavior in tissues when immersed in liquid nitrogen.

In the present study, electron microscopy combined with freeze-substitution was employed to examine the ultrastructure of cells of shoot apices of a Chrysanthemum plant that were cooled to the temperature of liquid nitrogen after exposure to various steps of the Cryo-plate protocol (Figure 5-6).

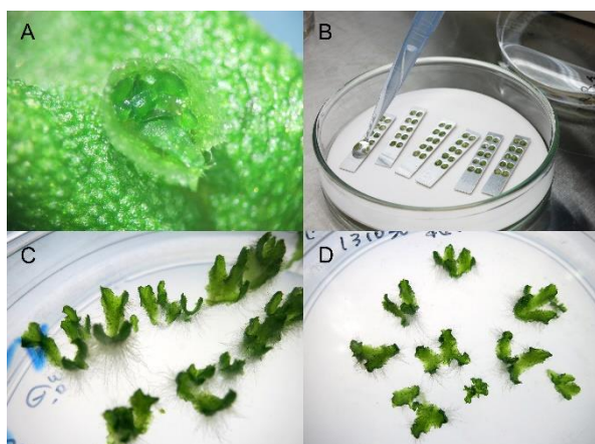


Figure 5. Cryopreservation protocol using aluminum cryo-plate.

A: Preparation of *in vitro* grown gemmae of liverwort. B: Placing precultured gemmae in a cryo-plate's wells. C: Regrowth of cryopreserved liverwort line 'Takaragaike-1' 30 days after rewarming. D: Control (without cooling in liquid nitrogen).



Figure 6. Plantlets regenerated from shoot apices of chrysanthemum cryopreserved by the D-Cryo-plate protocol. Shoot apices were on the post-thaw medium for 30 days. E, cryopreserved. F, control (without cooling in liquid nitrogen). G-H, Field preservation of chrysanthemum biological resources. About 2,000 races/lines are cultivated in the fields of Hiroshima University, Japan.

Access

From Central Japan International Airport (Centrair: NGO)

Take the Meitetsu bus bound for Okazaki Station and get off at Higashi Okazaki Station (approximately 1 hr from the airport). The Institute is a 7 min walk up the hill (Myodaiji-area) or 20 min walk (Yamate-area) on the south side of the station.



Cryopreservation Conference 2014 – NIBB

Cryopreservation Conference 2014

When: October 23 -24

Where: Okazaki Conference Center, Okazaki, Aichi, Japan

Who: NIBB

Abstract Deadline: September 19, 2014

Organizers: Tanaka D., Naruse K., Kawaguchi M., Fujikawa S.

Supported by NBRP Medaka, NIBB Spectrography and Bioimaging Facility



オーガナイザ: 田中大介, 成瀬 清, 川口正代司, 藤川清三
主催: 基礎生物学研究所 IBBPセンター
共催: 基礎生物学研究所 NBRPメダカ, 生物機能解析センター光学解析室

超低温保存研究の新たな挑戦

ゲノム編集技術の登場により非モデル生物でも容易に様々な変異体の作出が可能になりました。それにより、増え続ける変異体や絶滅危惧種などの遺伝資源の維持は研究の質や方向性に直接影響を与えます。新規超低温保存技術の開発やガラス化メカニズムに関する基礎研究の最新情報を共有し共同研究の輪を広げませんか?



講演者

10月23日

- ・藤川 清三 (北海道大学)
- ・菊地 和弘 (農業生物資源研究所動物発生分化研究ユニット)
- ・伴野 豊 (九州大学農学研究院附属遺伝子資源開発研究センター)
- ・柏木 昭彦 (広島大学大学院理学研究科附属両生類研究施設)
- ・今井 啓雄 (京都大学霊長類研究所)
- ・蛭田千鶴江 (自然科学研究機構岡崎統合バイオサイエンスセンター)
- ・大和 勝幸 (近畿大学生物理工学部)
- ・松村 和明 (北陸先端科学技術大学院大学マテリアルサイエンス研究科)
- ・白樫 了 (東京大学生産技術研究所機械・生体系部門)
- ・高松 洋 (九州大学大学院工学研究院機械工学部門)

10月24日

- ・津田 栄 (産業技術総合研究所生物プロセス研究部門)
- ・櫻井 実 (東京工業大学バイオ研究基盤支援総合センター)
- ・石川 雅也 (農業生物資源研究所植物生産生理機能研究ユニット)
- ・田上 貴寛 (畜産草地研究所家畜育種研究グループ)
- ・楠 比呂志 (神戸大学農学部)
- ・校重 圭祐 (高知大学農学部)
- ・高橋 恒夫 (京都大学再生医科学研究所幹細胞研究部門)
- ・森 史 (地球・人間環境フォーラム, 国立環境研究所)
- ・上村 松生 (若手大学農学部附属寒冷バイオフロンティア研究センター)
- ・中桐 昭 (鳥取大学農学部附属菌類きのこ遺伝資源研究センター)
- ・稲葉 重樹 (製品評価技術基盤機構バイオテクノロジーセンター)
- ・藤江 昭彦 (医薬基盤研究所創薬支援戦略室)
- ・岩本 まり (熊本大学生命資源研究・支援センター)
- ・田中 大介 (基礎生物学研究所 IBBPセンター)



大学連携バイオバックアッププロジェクト (IBBP) は、全国の大学や公設試験研究機関に所属する研究者が利用できる生物遺伝資源バックアップシステムです。IBBPセンターは基礎生物学研究所に集中バックアップ保管施設として設置され、平成25年より全国の研究者からお預かりしたサンプルのバックアップ保管を開始しています。また、共同利用研究の公募を行い研究者と共同で新規超低温保存法の開発を行っています。



日程 2014年 10月23日(木), 24日(金)

場所 自然科学研究機構 岡崎コンファレンスセンター
愛知県岡崎市明大寺町字伝馬8-1 東岡崎駅から徒歩15分

参加費 無料 (ポスター発表の登録は9月19日まで)

その他 平成27年度 IBBP共同利用研究の公募説明会を行います

連絡先 Cryopreservation Conference 2014 事務局
TEL : 0564-59-5930 email : cryo2014@nibb.ac.jp



詳細は web サイトをご覧ください
<http://www.nibb.ac.jp/ibbp/cryoconf2014/>

IBBP 検索

Research activity by T. Kimura

Assistant Professor: *KIMURA, Tetsuaki*

Analysis of median fin-rays development

The vertebrate body plan has evolved by the acquisition of structures projecting from the body axis. The original structures have transformed, as an environmental adaptation, into appendages such as fins, limbs, and wings. Median fins are the oldest of such evolved structures. In order to better understand the mechanisms by which the median fins developed, we crossed two inbred lines of medaka (*Oryzias latipes*). From the results, we found that the number of anal fin-rays was determined by two genetic traits, the anteroposterior length of the anal fin and interval between the anal fin-rays. The 19-ray fish has a longer anal fin than the 17-ray fish (Figure 4A). The 19-ray fish has the same anal fin length as the 17-ray fish (Figure 4B). This indicates that the 19-ray fish has narrower intervals between fin-rays than the 17-ray fish.

Further, the pattern of rays was independent of the pattern of the somites and vertebrae. Thus, the pattern formation of fin-rays proposes a new model of bone patterning.

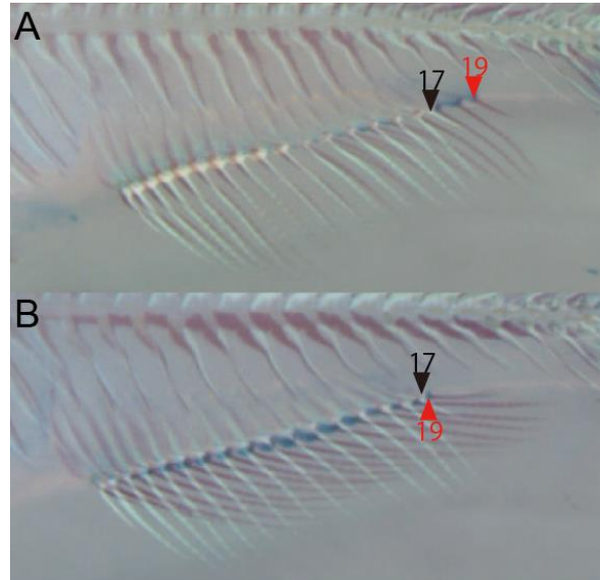


Figure 1. Superimposition of F₂ anal fin-rays. (A) Superimposed images of F₂ fish with 17 and 19 rays. (B) Superimposed images of F₂ fish with 17 (the same fish as in A) and 19 rays (a different fish from that in A). All three fish have 29 vertebrae. Note the pattern of the vertebrae is the same for both A and B. Arrowheads indicate the posteriormost ray. Numbers indicate total number of fin-rays.