## **NIBB CORE RESEARCH FACILITIES**



*Head* YOSHIDA, Shosei

The NIBB Core Research Facilities were launched in 2010 to support basic biology research in NIBB. They consist of three facilities that are developing and providing state-of-theart technologies to understand biological functions through functional genomics, bioimaging, and bioinformatics.

The NIBB Core Research Facilities also act as an intellectual hub to promote collaboration among the researchers of NIBB and other academic institutions.



The Functional Genomics Facility is a division of the NIBB Core Research Facilities and is organized jointly by NIBB and NIPS for promoting DNA and protein studies. The facility maintains a wide array of core research equipment, from standard machinery like ultracentrifuges to cutting edge tools such as next generation DNA sequencers, which amount to 60 different kinds of instruments. The facility is dedicated to fostering collaborations with researchers both of NIBB and other academic institutions worldwide by providing these tools as well as expertise. Our current focus is supporting functional genomics studies that utilize mass spectrometers and DNA sequencers. We also act as a bridge between experimental biology and bioinformatics.

We recently largely renovated the building of the Functional Genomics Facility. For example, the Visitors Lab and the Visitors Office were newly designed so that we can promote collaboration projects. Indeed, in 2016, more than 200 researchers visited to use our new facility and developed active collaborations.

# **Representative Instruments** *Genomics*

The advent of next-generation sequencing (NGS) technologies is transforming today's biology by ultra-high-throughput DNA sequencing. Utilizing HiSeq2500, HiSeq1500, and MiSeq (Illumina), and PacBio RS II (PacificBio Sciences), the Functional Genomics Facility is committed to joint research aiming to explore otherwise inaccessible new fields in basic biology.

During 2016 we carried out 52 NGS projects in collaboration with NIBB laboratories as well as the researchers of other academic institutions. These projects cover a wide range of species (bacteria, animals, plants, and humans) including both model and non-model organisms, and various applications such as genomic re-sequencing, RNA-seq and ChIP-seq.



Figure 1. Next-generation sequencer

### **Proteomics**

Three different types of mass spectrometer and two protein sequencers, as listed below, are used for proteome studies in our facility. In 2016, we analyzed approximately 700 samples with mass spectrometers and protein sequencers.

- LC-MS (AB SCIEX TripleTOF 5600 system)
- LC-MS (Thermo Fisher SCIENTIFIC Orbitrap Elite)
- MALDI-TOF-MS (Bruker Daltonics REFLEX III)
- LC-Q-TOF MS (Waters Q-TOF Premier)
- Protein sequencer (ABI Procise 494 HT; ABI Procise 492 cLC)

#### **Other analytical instruments** (excerpts)

- Cell sorter (SONY SH800, BD FACS Aria II)
- Bioimaging Analyzer (Fujifilm LAS 3000 mini; GE FLA9000)
- Laser Capture Microdissection System (Arcturus XT)
- DNA Sequencer (ABI 3130xl)
- Real Time PCR (ABI 7500)
- Ultra Centrifuge (Beckman XL-80XP etc.)



Figure 2. Triple TOF LC/MS/MS System

## **Genome Informatics Training Course**

We organize NIBB Genome Informatics Training Courses every year. These courses are designed to introduce basic knowledge and skills of bioinformatics to biologists who are not familiar with bioinformatics. In 2016, we provided two courses on RNA-seq data analysis and on BLAST sequence analysis, where a total of 107 researchers and students joined from all over Japan.



Figure 3. NIBB Genome Informatics Training Course

#### **Publication List on Cooperation:**

### [Original papers]

- Akashi, H.D., Cádiz Díaz, A., Shigenobu, S., Makino, T., and Kawata, M. (2016). Differentially expressed genes associated with adaptation to different thermal environments in three sympatric cuban anolis lizards. Molec. Ecol. 25, 2273–2285.
- Higo, A., Niwa, M., Yamato, K.T., Yamada, L., Sawada, H., Sakamoto, T., Kurata, T., Shirakawa, M., Endo, M., Shigenobu, S., *et al.* (2016). Transcriptional framework of male gametogenesis in the liverwort *Marchantia polymorpha* L. Plant Cell Physiol. 57, 325–338.
- Ide, T., Mochiji, S., Ueki, N., Yamaguchi, K., Shigenobu, S., Hirono, M., and Wakabayashi, K. (2016). Identification of the *agg1* mutation responsible for negative phototaxis in a "wild-type" strain of *Chlamydomonas reinhardtii*. Biochem. Biophys. Rep. 7, 379-385.
- Koga, H., Fujitani, H., Morino, Y., Miyamoto, N., Tsuchimoto, J., Shibata, T.F., Nozawa, M., Shigenobu, S., Ogura, A., Tachibana, K., *et al.* (2016). Experimental approach reveals the role of *alx1* in the evolution of the echinoderm larval skeleton. PLoS ONE *11*, e0149067.
- Matsui, H., Takahashi, T., Murayama, S.A., Uchiyama, I., Yamaguchi, K., Shigenobu, S., Suzuki, M., Rimbara, E., Shibayama, K., Øverby, A., and Nakamura, M. (2016). Draft genome sequence of *Helicobacter suis* strain SNTW101, isolated from a Japanese patient with nodular gastritis. Genome Announc. 4, e00934-16.
- Oda, K., Kamiya, T., Shikanai, Y., Shigenobu, S., Yamaguchi, K., and Fujiwara, T. (2016). The arabidopsis Mg transporter, MRS2-4, is essential for Mg homeostasis under both low and high Mg conditions. Plant Cell Physiol. 57, 754-763.
- Sato, K., Tanaka, T., Shigenobu, S., Motoi, Y., Wu, J., and Itoh, T. (2016). Improvement of barley genome annotations by deciphering the Haruna Nijo genome. DNA Res. 23, 21–28.
- Tong, W., Imai, A., Tabata, R., Shigenobu, S., Yamaguchi, K., Yamada, M., Hasebe, M., Sawa, S., Motose, H., and Takahashi, T. (2016). Polyamine resistance is increased by mutations in a nitrate transporter gene NRT1.3 (AtNPF6.4) in *Arabidopsis thaliana*. Front. Plant Sci. 7, 834.
- Ueki, N., Ide, T., Mochiji, S., Kobayashi, Y., Tokutsu, R., Ohnishi, N., Yamaguchi, K., Shigenobu, S., Tanaka, K., Minagawa, J., Hisabori, T., Hirono, M., and Wakabayashi, K. (2016). Eyespot-dependent determination of the phototactic sign in *Chlamydomonas reinhardtii*. Proc. Natl. Acad. Sci. USA *113*, 5299-5304.

 Yatsu, R., Miyagawa, S., Kohno, S., Parrott, B.B., Yamaguchi, K., Ogino, Y., Miyakawa, H., Lowers, R.H., Shigenobu, S., Guillette, L.J. Jr., and Iguchi, T. (2016). RNA-seq analysis of the gonadal transcriptome during *Alligator mississippiensis* temperature-dependent sex determination and differentiation. BMC Genomics 17, 77

### [Original paper (E-publication ahead of print)]

- Kondo, S., Wakae, K., Wakisaka, N., Nakanishi, Y., Ishikawa, K., Komori, T., Moriyama-Kita, M., Endo, K., Murono, S., Wang, Z., Kitamura, K., Nishiyama, T., Yamaguchi, K., Shigenobu, S., Muramatsu, M., and Yoshizaki, T. APOBEC3A associates with human papillomavirus genome integration in oropharyngeal cancers. Oncogene 2016 Oct 3.
- Murase, K., Shigenobu, S., Fujii, S., Ueda, K., Murata, T., Sakamoto, A., Wada, Y., Yamaguchi, K., Osakabe, Y., Osakabe, K., *et al.* MYB transcription factor gene involved in sex determination in *Asparagus* officinalis. Genes Cells 2016 Nov 21.

#### Research activity by S. Shigenobu

Specially Appointed Associate Professor: SHIGENOBU, Shuji NIBB Research Fellow: OGAWA, Kota Visiting Graduate Student: HSIAO, Yi-Min Technical Assistant: SUZUKI, Miyuzu

### **Symbiogenomics**

"Nothing, it seems, exists except as part of a network of interactions." (Gilbert & Epel, 2008)

Every creature on the earth exists among a network of various biological interactions. For example, many multicellular organisms, including humans, harbor symbiotic bacteria in their bodies: some of them provide their hosts with essential nutrients deficient in the host's diet and others digest foods indigestible by the host alone. In spite of numerous examples of symbioses and its intriguing outcomes, the genetic and molecular basis underlying these interactions remains elusive. The goal of our group is to establish a new interdisciplinary science "Symbiogenomics", where we aim to understand the network of biological interactions at the molecular and genetic level. To this end, we take advantage of state-of-the-art genomics such as next-generation sequencing technologies.

# I. Genomic revelations of a mutualism: the pea aphid and its obligate bacterial symbiont

Aphid species bear intracellular symbiotic bacteria in the cytoplasm of bacteriocytes, specialized cells for harboring the bacteria. The mutualism is so obligate that neither can reproduce independently. The 464 Mb draft genome sequence of the pea aphid, *Acyrthosiphon pisum*, in consort with that of bacterial symbiont *Buchnera aphidicola* illustrates the remarkable interdependency between the two organisms. Genetic capacities of the pea aphid and the symbiont for amino acid biosynthesis are complementary. The genome analysis revealed that the pea aphid has undergone characteristic gene losses and duplications. The IMB

antibacterial immune pathway is missing several critical genes, which might account for the evolutionary success of aphids to obtain beneficial symbionts. Lineage-specific gene duplications have occurred in genes in a broad range of functional categories, which include signaling pathways, miRNA machinery, chromatin modification and mitosis. The importance of these duplications for symbiosis remains to be determined. We found several instances of lateral gene transfer from bacteria to the pea aphid genome. Some of them are highly expressed in bacteriocytes.

We recently discovered a novel class of genes in the pea aphid genome that encode small cysteine-rich proteins with secretion signals that are expressed exclusively in bacteriocytes of the pea aphid, and named these bacteriocyte-specific cysteine-rich proteins (BCR). The BCR mRNAs are first expressed at a developmental time point coincident with the incorporation of symbionts strictly in the cells that contribute to the bacteriocyte, and this bacteriocyte-specific expression is maintained throughout the aphid's life. Some BCRs showed an antibiotic activity. These results suggest that BCRs act within bacteriocytes to mediate the symbiosis with bacterial symbionts, which is reminiscent of the cysteinerich secreted proteins of leguminous plants that also regulate endosymbionts. Employment of small cysteine-rich peptides may be a common tactic of host eukaryotes to manipulate bacterial symbionts.



Figure 1. Pea aphids and the bacterial symbiont, *Buchnera*. Adult aphids (Left). A developing viviparous embryo which symbionts are infecting (Right). Scale bar = 20um.

#### **Publication List:**

[Original paper]

• Hongo, Y., Ikuta, T., Takaki, Y., Shimamura, S., Shigenobu, S., Maruyama, T., and Yoshida, T. (2016). Expression of genes involved in the uptake of inorganic carbon in the gill of a deep-sea vesicomyid clam harboring intracellular thioautotrophic bacteria. Gene 585, 228-240.

## SPECTROGRAPHY AND BIOIMAGING FACILITY



Specially Appointed Associate Professor: KAMEI, Yasuhiro

Technical Staff:

Technical Assistant:

KONDO, Maki TANIGUCHI-SAIDA, Misako UCHIKAWA, Tamaki ICHIKAWA, Chiaki ISHIKAWA, Azusa

The Spectrography and Bioimaging Facility assists both collaborative and core research by managing and maintaining research tools that use "Light". The facility also provides technical support through management of technical staff assisting in the advancement of collaborative and core research projects, as well as academic support to researchers by Dr. Y. Kamei (refer to the Collaborative Research Group Research Enhancement Strategy Office section). Among its tools are advanced microscopes for biology and the Okazaki Large Spectrograph for photobiology. The Okazaki Large Spectrograph is the world's largest wide spectrum exposure mechanism, capable of producing a range of wavelengths from 250 nm (ultraviolet) to 1,000 nm (infrared) along its 10 meter focal curve; allowing exposure to strong monochromatic light. The facility's microscopes, which are cutting edge devices such as confocal and multi-photon excitation microscopes, are used by both internal and external researchers as vital equipment for core and collaborative research projects.

# **Representative Instruments: Okazaki Large Spectrograph (OLS)**

The spectrograph runs on a 30 kW Xenon arc lamp and projects a wavelength spectrum from 250 nm (ultraviolet) to 1,000 nm (infrared) onto its 10 m focal curve with an intensity of monochromatic light at each wavelength more than twice as much as that of the corresponding monochromatic component of tropical sunlight at noon (Watanabe *et al.*, Photochem. Photobiol. *36*, 491-498, 1982). The spectrograph is dedicated to action spectroscopical studies of various light-controlled biological processes.

The NIBB Collaborative Research Program for the Use of



Figure 1. An example of experiments using the Large Spectrograph. Various color rays (monochromatic light from right side and reflected by mirrors) were irradiated simultaneously to samples in cooling chambers.

the OLS supports about 10 projects every year conducted by both visiting scientists, including foreign researchers, as well as those in NIBB.

Action spectroscopical studies for various regulatory and damaging effects of light on living organisms, biological molecules, and artificial organic molecules have been conducted.

### **Microscopes**

This facility also has Bioimaging machines such as widefield microscopes (Olympus IX-81 and BX-63), confocal microscopes (Olympus FV1000, Nikon A1R, Nikon A1Rsi and Yokogawa CSU-X1 with EM-CCD camera), multiphoton microscopes (Olympus FV1000-MP, FV1200-MPs, Leica TCS-SP8 MPs) and other advanced custom-made laser microscopes with special aims (Digital Scanned Lightsheet Microscope: DSLM and Infrared Laser-Evoked Gene Operator microscope: IR-LEGO) for users in NIBB and collaborative guest researchers. We began two new types of Collaborative Research Program. One is a new category of the NIBB Collaborative Research for Integrative Bioimaging using machines and bioimage processing/analysis techniques, and the other is the Advanced Bioimaging Support Program (ABiS) of the Grant-in-aid for Scientific Research on Innovative Areas.

The DSLM was developed by Dr. Ernst Stelzer's group at the European Molecular Biology Laboratory (EMBL). This microscope can realize high-speed z-axis scanning in deeper tissue by illuminating a specimen from the side with a light sheet (more information is given in Dr. Nonaka's section: Lab. for Spatiotemporal Regulations). Dr. Shigenori Nonaka conducted and supported about 10 projects as Collaborative Research Programs for Integrative Bioimaging. The IR-LEGO was developed by Drs. Shunsuke Yuba and Yasuhiro Kamei at the National Institute of Advanced Industrial Science and Technology (AIST). This microscope can induce a target gene of interest by heating a single target cell in vivo with a high efficiency irradiating infrared laser. Details are described in the next section. The IR-LEGO was also used for about 10 Collaborative Research projects, including applications for animals and plants.

### Workshop and Symposium

In 2016 we held courses during both the 9<sup>th</sup> NIBB International Practical Course which focused on basic techniques for medaka and zebrafish research including imaging, and the 4<sup>th</sup> biological image processing training course. We also have been holding a "Bioimaging Forum" every year which discusses Bioimaging from various directions such as microscopy, new photo-technology, and computer science. This year we held the 10<sup>th</sup> NIBB Bioimaging Forum focused on adaptive optics for microscopy. In addition, we held three symposiums focused on new emerging model animals, next generation research using amphibians, and heat and temperature in biology.

**Publication List on Cooperation** 

[Original papers (Selected)]

- Okada, K., Inohaya, K., Mise, T., Kudo, A., Takada, S., and Wada, H. (2016). Reiterative expression of pax1 directs pharyngeal pouch segmentation in medaka. Development 143, 1800-1810.
- Petroutsos, D., Tokutsu, R., Maruyama, S., Flori, S., Greiner, A., Magneschi, L., Cusant, L., Kottke, T., Mittag, M., Hegemann, P., Finazzi, G., and Minagawa, J. (2016). A blue-light photoreceptor mediates the feedback regulation of photosynthesis. Nature 537, 563-566.
- Shikata, T., Matsunaga, S., Kuwahara, Y., Iwahori, S., and Nishiyama, Y. (2016). Light spectrum regulates cell accumulation during daytime in the raphidophyte Chattonella antiqua causing noxious red tides. J. Photochem. Photobiol. B: Biology *160*, 128-133.
- Suthaparan, A., Solhaug, K. A., Stensvand, A., and Gislerod, H.R. (2016). Determination of UV action spectra affecting the infection process of *Oidium neolycopersici*, the cause of tomato powdery mildew. J. Photochem. Photobiol. B: Biology *156*, 41-49.
- Utagawa, U., Higashi, S., Kamei, Y., and Fukamachi, S. (2016). Characterization of assortative mating in medaka: Mate discrimination cues and factors that bias sexual preference. Horm. Behav. 84, 9-17.
- Yokoyama, R., Yamamoto, H., Kondo, M., Takeda, S., Ifuku, K., Fukao, Y., Kamei, Y., Nishimura, M., and Shikanai, T. (2016). Grana-localized proteins, RIQ1 and RIQ2, affect the organization of light-harvesting complex II and grana. Plant Cell 28, 2261–2275.
- Zeng, C.W., Kamei, Y., Wang, C.T., and Tsai, H.J. (2016). Subtypes of hypoxia-responsive cells differentiate into neurons in spinal cord of zebrafish embryos after hypoxic stress. Biol. Cell 108, 1–21.

#### Research activity by Y. Kamei

Specially Appointed Associate Professor:

NIBB Research Fellow: Postdoctoral Fellow: Technical Assistant: KAMEI, Yasuhiro HATTORI, Masayuki ANDO, Yoriko CHISADA, Eriko ATSUMI, Urumi

Our research group promotes two cutting-edge microscope projects; "observation" and "manipulation" using optical and biological technologies. The aim of our "observation project" is deep-seeing in living organisms using adaptive optics (AO) which were well-developed in the field of astronomy as a key technology of large telescopes such as the Subaru telescope in Hawaii. Although observation using telescopes on the earth may be disturbed by fluctuations in the atmosphere, AO technology can cancel this disturbance. On the other hand, living materials have particular refractive indexes, therefore, some organelles act as disturbances of the ideal optical path for microscope observation just like the atmosphere does for telescopes. AO technology can also compensate for this disturbance by sensing and correcting wave fronts using a wave front sensor and deformable mirror. Hence, we developed a custom-made wide-field microscope equipped with an AO system for observation of living organisms in collaboration with Dr. Tamada in NIBB and Dr. Hayano in the National Astronomical Observatory of Japan (NAOJ) and got high-resolution bright field and fluorescent images of living cells. Our results indicated that improvement of optical resolution was restricted to a small area which is called the "isoplanatic patch" (Figure 1).

Second, the aim of our "manipulation project" is to control gene expression *in vivo*. Gene function analysis must be evaluated at the cell level *in vivo*. To achieve spatiotemporalcontrolled gene expression we employed one of the stress responses, the heat shock response. The heat shock promoter



Figure 1. Effects of adaptive optics (AO) to wide-field microscope images (bright field and fluorescence of plant cells).

is the transcription regulation region of heat shock proteins and all organisms have this mechanism. Positioning the target gene downstream of the promoter, we can induce the target gene expression by local heating.

Infrared (IR) beams can heat water molecules, which are the main constituent of cells, hence, we can heat a single cell by irradiating IR to a target cell using a microscope. We have developed a microscope, IR laser evoked gene operator (IR-LEGO), specialized for this purpose (Figure 2). The IR-LEGO microscope can irradiate an IR laser to a single cell *in vivo* such as in *C. elegans*, *Drosophila*, medaka, zebrafish, *Xenopus* and *Arabidopsis*, to induce the heat shock response at a desired timing. In 2015, additionally, we confirmed the system was effective in the moss *Marchantia polymorpha* and in the newt *Pleurodeles waltl*.



Figure 2. Schematic illustration of heat shock response of cells (left) and an infrared laser-evoked gene operator (IR-LEGO) microscope system.

Optimal heating induces the heat shock response and subsequent gene expression, while an excess results in cell death. Hence, we must precisely control laser heating. We evaluated time course and spatial heating profiles, and the results presented that temperature of the target area rose rapidly and kept a constant level dependant on IR laser power, additionally, the heated area was adequately as small as a typical cell size.

With this in mind, we tried to induce gene expression in various species. At first, we reported an IR-LEGO experiment in living C. elegans. Target gene expression in a target cell could be induced with only 1 s-IR irradiation. Whereas the optimal power range which can induce gene induction without cell damage was limited. Excess laser power resulted in cell death or cessation of cell division. We confirmed that an optimal irradiation, e.g. 11 mW for 1 s, induced physiological gene expression in the target cell and subsequent cell division or morphogenesis underwent normal development. Next, we tried the experiment in other animals, such as, medaka, zebrafish and Xenopus, and the higher plant, Arabidopsis, since all organisms have a heat shock response system. We succeeded in local gene induction in all the species as expected. Moreover, this system can be combined to the cre/loxP recombination technique for long-term gene



Figure 3. Long-term gene expression system with cre/loxP recombination system and its example of practical experiment in medaka embryo to adult.

## expression (Figure 3).

As mentioned above, excess irradiation resulted in cell damage, so we utilized the system to ablate target cells with strong pulsed irradiation. In collaboration with National Taiwan University, we used the system for neuronal regeneration study in zebrafish and revealed that a kind of neuronal precursor cell played an important role within the neuron regeneration step the in zebrafish spinal cord (Zeng et al. Biol. Cell 2016). In addition, the IR-LEGO system can be utilized for thermo-biology because spatiotemporal micrometer order local heating is difficult without this system. Now we are trying to estimate thermal properties of cells and biomaterials *in vivo* using the IR-LEGO system.

#### **Publication List:**

[Original papers]

- Nishihama, R., Ishida, S., Urawa, H., Kamei, Y., and Kohchi, T. (2016). Conditional gene expression/deletion systems for Marchantia polymorpha using its own heat-shock promoter and the cre/loxPmediated site-specific recombination. Plant Cell Physiol. 57, 271-280.
- Suzuki, M., Takagi, C., Miura, S., Sakane, Y., Suzuki, M., Sakuma, T., Sakamoto, N., Endo, T., Kamei, Y., Sato, Y., Kimura, H., Yamamoto, T., Ueno, N., and Suzuki, K.T. (2016). In vivo tracking of histone H3 lysine 9 acetylation in *Xenopus laevis* during tail regeneration. Genes Cells 21, 358-369.
- Utagawa, U., Higashi, S., Kamei, Y., and Fukamachi, S. (2016). Characterization of assortative mating in medaka: Mate discrimination cues and factors that bias sexual preference. Horm. Behav. 84, 9-17.
- Yokoi, S., Ansai, S., Kinoshita, M., Naruse, K., Kamei, Y., Young, L.J., and Takeuchi, H. (2016). Mate-guarding behavior enhances male reproductive success via familiarization with mating partners in medaka fish. Front. Zool. 13, 21.
- Yokoyama, R., Yamamoto, H., Kondo, M., Takeda, S., Ifuku, K., Fukao, Y., Kamei, Y., Nishimura, M. and Shikanai, T. (2016). Grana-localized proteins, RIQ1 and RIQ2, optimize the dynamics of light-harvesting complex II and grana stacking in Arabidopsis. Plant Cell 28, 2261-2275.
- Zeng, C.W., Kamei, Y., Wang, C.T., and Tsai, H.J. (2016). Subtypes of hypoxia-responsive cells differentiate into neurons in spinal cord of zebrafish embryos after hypoxic stress. Biol. Cell, 108, 357-377.

## Data Integration and Analysis Facility

Assistant Professor: Technical Staff:

Technical Assistant:

UCHIYAMA, Ikuo MIWA, Tomoki NISHIDE, Hiroyo NAKAMURA, Takanori OKA, Naomi

The Data Integration and Analysis Facility supports research activities based on large-scale biological data analysis, such as genomic sequence analysis, expression data analysis, and imaging data analysis. For this purpose, the facility maintains high-performance computers with large-capacity storage systems. On the basis of this system, the facility supports development of data analysis pipelines, database construction and setting up websites to distribute the data worldwide as well as providing users' basic technical support. In addition to computational analysis, the Data Integration and Analysis Facility supports NIBB's information infrastructure, the maintenance of the network systems in the institute and computer/network consultation for institute members.

### **Representative Instruments**

Our main computer system is the Biological Information Analysis System (BIAS) (Figure 1), which consists of a high-performance cluster system (SGI Rackable server C2112-4RP; 40 nodes/800 cores, 96GB memory/node), a shared memory parallel computer (HP ProLiant DL980 G7; 80 cores, 4TB memory), a high-throughput storage system (DDN SFA7700; 480TB), and a large capacity storage system (DELL PowerEdge R620; 720TB). All subsystems are connected via a high-speed InfiniBand network so that large amounts of data can be processed efficiently. Some personal computers and color printers are also available. On this system, we provide various biological databases and data retrieval/analysis programs, and support large-scale data analysis and database construction for institute members and collaborative researchers. Especially, we have supported the construction and maintenance of published databases of various model organisms including XDB (Xenopus laevis), PHYSCObase (Physcomitrella patens), DaphniaBASE (Daphnia magna), The Plant Organelles Database, and MBGD (microbial genomes).

The facility also provides network communication services. Most of the PCs in each laboratory, as well as all of the above-mentioned service machines, are connected by a local area network, which is linked to the high performance backbone network ORION connecting the three research institutes in Okazaki. Many local services, including sequence analysis services, file sharing services, and printer services, are provided through this network. We also maintain a public World Wide Web server that hosts the NIBB home page (http://www.nibb.ac.jp/en).



Figure 1. Biological Information Analysis System

# **Research activity by I. Uchiyama**

Assistant professor I. Uchiyama is the principal investigator of the Laboratory of Genome Informatics, which currently focuses on microbial comparative genomics studies. For details, please refer to the laboratory page (p. 71).