Development of a protein array for autoantibody profiling of blood
—Comprehensive disease diagnosis using the body’s defense system—

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[Translation from Synthesiology, Vol.7, No.3, p.154-162 (2014)]

We have developed infrastructure of the technologies and resources for post-human genome research to perform functional proteomics (the analysis of protein functions, protein-protein interactions, and human protein structures) on a large scale. A method for profiling autoantibodies in serum is developed using human protein expression resources and protein expression techniques. The human biological defense system responds to abnormalities in the body with extraordinary sensitivity. Hence, this system is an effective tool for detecting human diseases at an early stage. Health safety and security can be achieved by establishing an early diagnostic method for diseases using autoantibody profiling of blood with a protein array.

Keywords: Autoantibody, protein array, human protein, body’s defense system, antigens, diagnosis, biomarker

1 Introduction

The ability to diagnose a disease before its development or at an early stage and obtaining a general diagnosis in which as much health information can be obtained in a single test is an extremely important issue in achieving a safe and secure society. In this research, we aim to achieve this objective by analyzing the types and quantities of autoantibodies in one drop of blood. Originally, the antibody is a biological defense system acquired by higher-order organisms through evolution, to defend themselves against bacteria and virus invasions. The antibody system is known to produce autoantibody against its own protein in response to the abnormal release of protein from cells or to the excessive production of proteins due to disease, as well as against external antigens. We think it is reasonable to utilize the biological defense mechanism that responds sensitively to the abnormalities of the body to detect diseases. Particularly, since the autoimmune diseases occur due to the production of antibodies that attack their own cells or tissues, the autoantibodies can be the cause of disease as well as disease markers. The detection of autoantibodies may enable presymptomatic testing for autoimmune diseases, thereby allowing early treatment. However, in practice, comprehensive tests for autoantibodies have not been established, and in most cases, one visits a hospital only after the symptoms of autoimmune diseases develop. There are also several intractable diseases in which the involvement of autoantibodies is suspected, and the development of a comprehensive detection system for autoantibodies is extremely important. There are many papers that reported the use of autoantibodies as disease markers for diseases including diabetes, cancer, Alzheimer’s disease, rheumatism, and dilated cardiomyopathy. We developed a comprehensive detection system of autoantibodies and have correlated the autoantibodies and diseases, by preparing antigen proteins using the world’s largest human protein expression resource and a human protein synthesis technology that we have been working on for a long time. We hope to further the technological development for a comprehensive profiling of blood autoantibodies that are individually different and are closely related to health.

2 Construction of a human protein expression resource and its use

Based on the human full-length cDNA sequencing project of the Ministry of International Trade and Industry (currently Ministry of Economy, Trade and Industry) that was started in 1998, we started preparations for the following in the Protein Function Analysis Project of the New Energy and Industrial Technology Development Organization (NEDO) in 2000: (1) Human Proteome Expression (HUPEX) resource, (2) high-throughput protein synthesis technology, and (3) Human Gene and Protein Database (HGPD). At the time, the Human Genome Project to decode the genome DNA sequence was being done internationally, and taking the lead for the coming age of proteomics, Japan decided to fortify the environment for human genome research, constructed the HUPEX resource, and built a database. The preparations of the technological foundation to carry out large-scale analyses of human protein functions, protein interactions, and protein structures were conducted as national projects. As a result, the HUPEX resource was utilized in various national research projects.
projects, joint research with corporations, and academic joint researches with research institutes and universities, and yielded rich results in the respective fields. One of the major results was a joint project “Special Project for Yamanaka iPS Cell” of the Japan Science and Technology Agency (JST) with Dr. Shinya Yamanaka, Director of the Center of iPS Cell Research and Application (CiRA), Kyoto University. This project led to the new discovery of Glis1, a gene for promoting the induction of iPS cells.[4] Moreover, in the joint research with the School of Medicine, Gifu University, a factor to highly efficiently induce iPS cells was discovered from the dental pulp cells in teeth. In the joint study with the School of Medicine, Keio University, a factor to promote direct reprogramming where the myocardial cells can be made from heart fibroblasts was discovered. Many results were obtained in the search for factors that may be useful in regenerative medicine. Also, major results were obtained in the development of in vitro visualization technology of protein interaction for a drug discovery screening system, and the development of production of standard proteins for quantitative proteomics by mass spectrometer. Such research results were expected in the initial conceptualization of the HUPEX resource uses. The HUPEX resource technology that we have constructed so far was basically to support the smooth progress of industrial proteome research. Sometimes, as we deepen our research in a discipline, new horizons that we did not initially consider begin to unfold. When we climbed the mountain of the proteome study and looked back, we realized that we could study the comprehensive antibody fields when the protein groups that comprise the proteome are considered as antigens. As shown in Fig. 1, we could study the immunome (whole immune system) from the proteome (while not all antigens are proteins, they comprise a major part). The idea of comprehensively analyzing the autoantibody in blood using the HUPEX resource and applying it in diagnosis (Fig. 2) was not considered initially. However, we were capable of using more human proteins as antigens than any other researcher in the world, and we possessed the ability to use the resource to find out whether the antibodies are present in the blood serum. Many researchers have previously reported the idea of using the autoantibodies in the blood serum.[1] However, since it was difficult to prepare the antigens to detect the antibodies, comprehensive analysis of autoantibody had not been done until now. Currently, we are able to conduct the world’s most accurate profiling of the blood serum autoantibody using the HUPEX resource. To realize autoantibody profiling, in addition to the HUPEX resource, it was necessary to establish a comprehensive protein expression technology, a technology for manufacturing protein arrays, and an antibody detection method. These technologies will be explained below.

![Fig. 1 From proteome to immunome](image1)

![Fig. 2 Autoantibody and disease](image2)
3 Development of a protein array that allows autoantibody profiling

3.1 Comprehensive human proteome expression resource and protein expression technology

We constructed a protein expression resource by introducing the Gateway cloning technology, a general-use cloning system, and created the plasmid DNA (entry clone) with site-specific recombination sequence at both ends of the open reading frame (ORF) of cDNA. In the Gateway cloning technology, the expression clone can be created simply by mixing the entry clone and the destination vector, and by conducting site-specific DNA recombination using the recombination enzyme in a test tube. This is an optimal DNA recombination technology when conducting high-throughput protein expression. This resource is the world’s largest protein expression resource that covers about 80 % of the human genome, and is named the HUPEX resource. Two types of entry clones were created: the N-type entry clone where the same amino acid sequence can be synthesized as the native protein at the C terminal per cDNA; and the F-type where the stop codon is replaced with a sense codon so a tag can be attached to the C terminal. Various types of ORF type entry clones were created including the full-length ORF type with whole ORF for the gene, processing ORF type where the signal peptides were eliminated from the full-length ORF, and the domain ORF that could express the extracellular domain or intracellular domain with membrane protein with single membrane penetration domain. These resources can be freely selected according to the research objective. Also, since the protein synthesis can be conducted with all types of protein synthesis systems, the SD (Shine-Dalgarno) sequence for E. coli expression and Kozak sequence for eukaryotic cell expression was added to the 5’ upstream of ORF. For these protein expression resources, the N-type and F-type entry clones were prepared, and about 60,000 types were created including the known and unknown clones, as well as the splicing variant clones (Table 1). To use these entry clones in a comprehensive proteome research, a clone with the longest ORF is selected for each gene as a representative clone, and they are functionally categorized according to the functions of proteins (transcription factor group, GPCR group, kinase group, unknown gene group, etc.). About 20,000 clones representing human genes were used in our research.

To conduct the comprehensive proteome research using human proteins, the technology to comprehensively synthesize the protein is necessary, as well as building the HUPEX resource. Around the year 2000 when we started constructing the protein expression resource, at the same time, Professor Yaeta Endo et al. of the Ehime University developed a wheat germ cell-free protein synthesis system. We developed the technology for high-throughput protein synthesis using the wheat germ cell-free protein synthesis system. The whole reaction from DNA structuring to protein synthesis was done in an in vitro system (96 hole or 384 hole plate), and the reaction was optimized so only the dispensing procedure where the reacted solution was transferred to the next reaction solution was necessary. As a result, we developed a technology where the whole process of protein synthesis could be completed in

Table 1. Number of manufactured Gateway entry clones

<table>
<thead>
<tr>
<th>Type</th>
<th>Determined number of entry clones</th>
<th>C terminal stop</th>
<th>C terminal fusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full-length ORF</td>
<td>18,744</td>
<td>28,386</td>
<td></td>
</tr>
<tr>
<td>Processing ORF</td>
<td>4,068</td>
<td>2,863</td>
<td></td>
</tr>
<tr>
<td>Domain ORF</td>
<td>2,719</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>25,531</td>
<td>31,249</td>
<td>56,780</td>
</tr>
</tbody>
</table>

Fig. 3 Wheat germ cell-free protein synthesis system using the Gateway entry clone

Note) Modification of Fig. 6 in N. Goshima et al.: Constructing the foundation for comprehensive expression of human proteins, Jpn. J. Exp. Med. (Extra no.), 23 (4), Chap. 3, Sec. 3, Yodosha (2005) (in Japanese).
one week. The LR product, PCR product, and mRNA that are created in the process of protein synthesis can be stored for a long period at -80 °C. In case it becomes necessary to resynthesize the same protein, it can be resynthesized in 18 hours using the stored mRNA. By combining the protein synthesis technology and a dispenser, it became possible to synthesize about 20,000 protein types at one time, and all proteins can be used in the simultaneous assay system through the array technology.

The results of the genetic information of the entry clones, the status of clones, and the results of protein expression of the wheat germ cell-free system or E. coli system are stored in the Human Gene and Protein Database (HGPD: http://www.HGPD.jp/), and they can be searched freely. The created entry clones are available from the National Biological Resource Center (NBRC), National Institute of Technology and Evaluation (NITE) (http://www.nbrc.nite.go.jp/hgentry.html).

3.2 Development of the protein array technology

The protein array technology is useful in comprehensively analyzing the interactions of protein-protein, protein-low molecule, protein-nucleic acid, etc., as well as for analyzing the enzyme-substrate protein. In a conventional protein array, the protein is fixed on the nitrocellulose membrane sheet, or on the surface of the slide glass coated or specially treated with nitrocellulose. Due to the nature of the fixing method, the protein is fixed in a dehydrated condition on the substrate surface, and the three-dimensional structure is not maintained. Therefore, with the conventional array, the function of the fixed protein cannot be analyzed. We aimed to recreate and analyze the biological reaction on the array using the HUPEX resource and the high-throughput protein synthesis technology that we developed. For this purpose, it was necessary to find a way to fix proteins on the array substrate in a condition where the 3D structures were maintained and the proteins could express their functions on the array.

We developed a protein array where 3D structures and functions of the proteins were maintained. First, we focused on the protein refining technology using magnetic beads. By using our protein synthesis technology, the proteins can be synthesized with various tags attached. The target protein synthesized with tags can be easily refined using the magnetic beads with ligands. First, the target protein is bonded to the magnetic beads while maintaining the 3D structure at the time of synthesis. Normally, the protein bonded to the magnetic beads are eluted, recovered, and used, but we considered a way to create an array while maintaining the bond between the magnetic beads and the protein. A well plate for bonding with the magnetic beads was developed, and by combining the high-throughput protein synthesis technology and the protein refining and array technologies using the magnetic beads, we developed the technology for creating an array while maintaining the 3D structure of the protein (Fig. 4A). The protein array, in

Fig. 4A Manufacture of protein active array using magnetic beads and antibody detection

The GST-fused target protein synthesized by the wheat germ cell-free synthesis system is attached to the GSH magnetic beads surface. This suspension is dispensed into the magnetic plate (originally made) equipped with magnets at the bottom of the plate, and the non-adhering fraction is removed by cleansing. The protein active array with protein fixed using the magnetic beads in the solution is manufactured. The antibody (Y) or blood serum is added to the protein active array (PAA), the protein is bonded onto the array, and the bonded antibodies are detected by chemiluminescence using the HRP-fused secondary antibody.

Note) Modification of Fig. 3 in N. Goshima: Autoantibody analysis using the array, Handbook of Therapeutic and Diagnostic Antibodies, Chap. 1, Sec. 5, NTS (2012) (in Japanese).
which the panoramic analysis of protein function for human proteome can be conducted, is called the “protein active array (PAA)” (Fig. 4B).

In selecting the magnetic beads used for the PAA, it is necessary to compare various magnetic beads from the following perspectives: 1) type of ligand of magnetic beads, 2) material and size of the beads, 3) amount of adsorbed target protein, 4) non-specific adsorption, and 5) method for dispensing the suspension. In general magnetic beads for His tag, GST tag, or Streptavidin tag adsorption are used. For the ligand of the magnetic beads, the GST tag adsorption magnetic beads was selected because the 5’ FLAG-GST tag that allows creation of active expressed protein for protein synthesis was used. From the perspectives of aforementioned 2) to 5), we selected the glutathione particles of the MagneGST Protein Purification System (Promega Corporation).

When creating the PAA, a special well plate is necessary to which the magnetic beads bonded with expressed proteins can be fixed, and which does not require dispensers or other equipment when cleansing or supplying the common reagent. We developed a well plate for the PAA, where the magnet is installed at the bottom of the well and the thickness of the well bottom is made as thin as possible to allow the magnetic beads to bind strongly to the well bottom. Normally, in assays such as ELISA, each well uses independent plates, but for the magnetic beads array, the wells are designed to be independent of each other but the reaction of the solution made of diluted serum can take place without division between the wells. We also developed a special cover plate that prevents biohazards when handling the serum samples, and that allows the reaction to be accomplished with a small amount of homogenous solution. By placing this cover plate, the target proteins on the surface of magnetic beads are covered with minute quantity of reaction liquid using a syringe, and the blood serum sample can be handled in a closed system rather than an open system.

By devising the equipment as described above, the manufacture process of the PAA can be accomplished very simply and in a short time. First, proteins synthesized by the wheat germ cell-free protein synthesis method using 96 well plates are added to the magnetic beads, and the proteins are bound to the surface of the magnetic beads. The suspension of the magnetic beads bound to proteins is dispensed to the well plate of the PAA, the proteins adhere to the well plate by magnetic force of the magnetic beads, and the array is created for the target protein. In case of long-term storage, the storage liquid is added and is stored at -80 ºC. As a result of storage tests, it was confirmed that the quality could be maintained for six months in the above storage condition.

3.3 Establishment of PAA detection method

The assay by PAA can be completed in about eight hours from the initial probe (serum, low molecular compound, proteins, etc. that will be investigated for the bond with human protein) reaction to detection. The number of samples that can be processed per day is four samples per person. For detection, secondary antibodies labeled with fluorescent pigment or secondary antibodies labeled with HRP for fluorescence detection are used. For detection, devices that can obtain the western blotting image can be used, such as the chemiluminescence image detector or fluorescence image detector that are commercially available. A liquid delivery pump is used for the cleansing process of the probe and the
reaction solution, and these can be automated in the future.

3.4 Establishment of the screening method for PAA

We are capable of synthesizing about 20,000 types of proteins in a short time by combining the protein synthesis technology and the dispenser. To screen proteins efficiently and economically using these characteristics, it is necessary to achieve higher density of PAAs or higher efficiency of screening. Cost and labor are needed to manufacture the protein arrays using the 20,000 proteins that were expressed individually and to identify the antigens, and problems may arise in the practical use and measurement of multiple samples. Therefore, we thought we could reduce the cost and labor to about one-tenth by conducting primary screening by creating an array using a compound of 10 types of proteins, and then conducting secondary screening using only the protein compounds containing the hits.

Several manufacture methods of the mixture of 10 proteins were considered. The method whereby simply synthesizing 10 proteins independently and then mixing the individual proteins would require cost and labor. Therefore, we investigated in which phase of the protein synthesis the 10 proteins could be combined for the co-expression of 10 types of proteins. In our protein synthesis system explained above (Fig. 3), the protein syntheses are conducted by mixing 10 types of entry clones, LR reaction products, PCR products, and mRNAs from each reaction step. The expressed proteins are separated by the SDS-PAGE method, and are detected by the western blotting method. The expressed proteins are separated by the SDS-PAGE method, and are detected by the western blotting method, anti-GST HRP-linked mouse mono Ab (NACALAI) is added to the antibody, diluted 5,000 times in 1.0 % skim milk in PBST, antibody reaction done for five minutes using ECL plus (GE Healthcare), and chemiluminescence detection is done using the Fluor-SMAX (Bio-Rad Laboratories, Inc.). As a result, for the co-expression of 10 proteins, 10 protein types were expressed efficiently in the mixture after the PCR product. Considering the cost and the complexity of the maneuver, it is concluded that the protein synthesis should be done after manufacturing the 10 mixtures using the PCR products from the uppermost stream (Fig. 5).

Using the PAA manufactured from the 10 types of co-expressed proteins synthesized from the mixed PCR product, we conducted an investigation of antigen-antibody reaction for the antibody to determine the antigen. As a result, the antigen could be identified using the PAA manufactured using the 10 types of co-expressed proteins.

The two-step screening method, where the 10 types of proteins obtained in the primary screening or comprehensive PAA (C-PAA) using 10 types of co-expressed protein are individually used to conduct antigen identification in the secondary screening expanded PAA (E-PAA), was employed. The screening is simplified using this two-step screening method (Fig. 6).

Using this array technology, we manufactured PAAs for about 20,000 types of proteins that are categorized by function. The manufacturing technology of PAAs was transferred to Cell Free Sciences, Co. Ltd. that is working on the product realization of the PAAs.

3.5 Analysis of the autoantibody in the blood serum of a patient with ovarian carcinoma derived paraneoplastic cerebellar degeneration

Using the PAA, the analysis of the autoantibody in the serum of a paraneoplastic cerebellar degeneration (PCD) patient was conducted. The patient visited the Department of Neurology,
The University of Tokyo Hospital, with numbness in the hands as an initial symptom. The patient’s blood serum was diluted 1,000 times with PBS-T, and the autoantibody detection was conducted using the chemiluminescence detector using the HRP-labeled anti-human IgG antibody. From the detection results, about 37 types of autoantibodies were found (Fig. 7). The list of detected autoantibodies are shown as antigens in Table 2. LIMS1 and TRIM21 are autoantibodies that are detected highly frequently (80 % or higher) in healthy individuals. Although it has been reported that TRIM21 may be a marker antibody for lung cancer,\(^9\) in our research, it was detected at high percentage (65 % or higher) in healthy individuals. Recently, the relationships to cancer have been reported for the SSX family of SSX1, SSX2, SSX3, SSX4B, and SSX5,\(^10\) cancer antigens (CTA) CTA45A4 and CTA45A5, and lipase MGLL.\(^11\) The antigen proteins of the autoantibodies detected this time were also reported to be related to cancer. On the other hand, IRX2 is a factor related to the formation of cerebellum,\(^12\) and CTNNB2 is a protein related to the intercellular communication in the neurological connection.\(^13\) From these findings, it was shown that the autoantibodies in the serum could be comprehensively detected using the PAA, and the antibodies related to cancer and cerebellar degeneration were detected. Since the cancer antibody was detected by the serum antibody analysis, the patient underwent thorough tests for cancer, and an ovarian tumor was found. It is not uncommon to find cancer in patients who visit the neurology department after experiencing some neurological symptoms. First, the cancer develops and several autoantibodies are produced as a result, and some of the autoantibodies may lead to neurological diseases. Currently, we are investigating whether these detected antibodies may become new biomarkers, by employing the JST Advanced Measuring and Equipment Development Program for 2012~2015 to obtain and analyze the autoantibody data of several patients and healthy subjects. If the autoantibody profiling can be done inexpensively and quickly using the PAA, it may be an extremely effective method for increasing the precision of various diagnoses.

### 4 Future issues

It is said that the antibodies are produced in large amounts, almost as much as the amplification of PCR against the protein.
amount of antigens. The antibodies circulate throughout the body through blood, and therefore, it is thought that minute changes in the body can be found by studying the antibodies. In fact, it is becoming possible to profile the changes in autoantibodies of the body using PAAs, and the autoantibody detection system is entering the practical phase in terms of technology.

In the future, the comprehensive detection system for autoantibody using PAAs will allow the autoantibodies to be profiled comprehensively for various diseases such as cancer and autoimmune diseases, and we hope to accumulate the data on the relationships between several diseases and autoantibodies. By doing this, we hope it will become possible to conduct an integrated test through autoantibody profiling of the blood, to allow the evaluations of progress and treatment of disease, early detection, policy for diagnosis and treatment, and therapeutic effects. In the development of PAAs, we are working on achieving higher density of protein arrays, and on finding economic ways of conducting highly sensitive measurement using small quantities of blood. We also wish to create a system that can be used easily in general hospitals and research facilities.

References


Yoshitaka KAWAKAMI
Withdrew from the doctorate course after obtaining the necessary credits from the Graduate School of Science, Osaka City University in 2008. Participated in the Regional Innovation Creation R&D Program “Development of Efficient Tumor Marker by Random Immunization” and “Development of Comprehensive Diagnostic System for Specific Cancers Using Autoantibodies,” as a technical staff at the Biomedical Information Research Center, AIST. Also worked in the “Multi-Institutional Pre-Phase II Clinical Trial for Post-Operative Esophageal Cancer Cases for CHP/ NY-ESO-1 Polypeptide Cancer Vaccine” of the Ministry of Health, Labour and Welfare. Joining the Fukushima Medical Industrial Translational Research Project as a researcher of the Regional Innovation Consortium in 2008~2010. Conducted the autoantibody profiling in blood sample to search for autoantibody biomarker. In this paper, was in charge of the creation of protein array and autoantibody measurement.

Naoki GOSHIMA
Completed the courses at the Department of Biochemistry, Graduate School of Agriculture and Biological Sciences, Osaka Prefecture University in 1987 (Doctor of Agricultural Science). Flexible Researcher, RIKEN; Assistant, Kyoto Pharmaceutical University; Assistant Professor, Graduate School of Science, Hiroshima University; and currently, Team Leader, Quantitative Proteomics Team, Molecular Profiling Research Center for Drug Discovery, AIST. Participated in the “Protein Function Analysis Project” of the New Energy and Industrial Technology Development Organization (NEDO) from 2000, and engaged in genome-wide protein functional analysis based on the Gateway creation of human full-length cDNA and comprehensive protein expression. Utilized the human protein expression resource and obtained results for the NEDO “Chemo-Bio Project,” NEDO “TR Product,” and “Special Project for Yamana iPS Cell” of Japan Science and Technology Agency in 2006. Engaged in the development of PAA in the Kyushu Regional Consortium in 2006 and the Kanto Regional Innovation in 2008~2010. Conducted the autoantibody profiling in blood sample to search for autoantibody biomarker. In this paper, was in charge of supervising the protein array research and collaboration with the medical practices.

Discussions with Reviewers

1 Overall comments

Question and comment (Yasushi Mitsuishi, AIST Tohoku)

I read this paper with interest, on the use of the results of the national project following from the human full-length cDNA project, and on the possibility of enabling early diagnosis of disease through blood serum autoantibody profiling that was not initially considered. I understood the idea that some kind of autoantibody is produced excessively in the development of a disease and that this can be used as a disease marker, but I felt the explanation of the technological result was too brief and I was somewhat left unsatisfied. The paper seems to finish with the conclusion that you created a PAA that allows the comprehensive analysis of the bonded protein in blood serum. While it is great that you can do comprehensive analysis of proteins, but in this paper, please describe in detail the types of serum autoantibodies, the amount, and the level at which the presence and degree of disease can be estimated.

Answer (Naoki Goshima)

The detailed results described in the initial manuscript will be submitted to a specialized journal, including the consideration of autoantibody as a marker. Therefore, I responded to your comment by including the analysis of the autoantibodies in the blood serum of a patient with ovarian carcinoma derived paraneoplastic disease through blood serum autoantibody profiling that was not initially considered. I understood the idea that some kind of autoantibody is produced excessively in the development of a disease and that this can be used as a disease marker, but I felt the explanation of the technological result was too brief and I was somewhat left unsatisfied. The paper seems to finish with the conclusion that you created a PAA that allows the comprehensive analysis of the bonded protein in blood serum. While it is great that you can do comprehensive analysis of proteins, but in this paper, please describe in detail the types of serum autoantibodies, the amount, and the level at which the presence and degree of disease can be estimated.

2 Concept of PAA

Question and comment (Noboru Yumoto, AIST)

The objective of this research is “the development of a comprehensive detection system for autoantibodies,” and centering on the major breakthrough of “the construction of the world’s largest protein expression resource,” the scenario, in which the elemental technologies including comprehensive protein expression technology, protein array technology, antibody detection technology, and screening technology were integrated, is understandable to the readers of the bio field. However, I think it is difficult for people outside the field to understand what actually is a proteome array. Therefore, can you include a conceptual diagram of the PAA that you developed?

Answer (Naoki Goshima)

I added the conceptual diagram of the PAA to Fig. 4B, and added explanations to the diagram, Fig. 4.