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Gregor Kijanka

*Royal College of Surgeons in Ireland*

Derek Murphy

*Royal College of Surgeons in Ireland*

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# Protein arrays as tools for serum autoantibody marker discovery in cancer

Gregor Kijanka, Derek Murphy\*

Centre for Human Proteomics, Royal College of Surgeons in Ireland

## ARTICLE DATA

## ABSTRACT

Protein array technology has begun to play a significant role in the study of protein-protein interactions and in the identification of antigenic targets of serum autoantibodies in a variety of autoimmune disorders. More recently, this technology has been applied to the identification of autoantibody signatures in cancer.

The identification of tumour-associated antigens (TAAs) recognised by the patient's immune response represents an exciting approach to identify novel diagnostic cancer biomarkers and may contribute towards a better understanding of the molecular mechanisms involved. Circulating autoantibodies have not only been used to identify TAAs as diagnostic/prognostic markers and potential therapeutic targets, they also represent excellent biomarkers for the early detection of tumours and potential markers for monitoring the efficacy of treatment. Protein array technology offers the ability to screen the humoral immune response in cancer against thousands of proteins in a high throughput technique, thus readily identifying new panels of TAAs. Such an approach should not only aid in improved diagnostics, but has already contributed to the identification of complex autoantibody signatures that may represent disease subgroups, early diagnostics and facilitated the analysis of vaccine trials.

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## 1. Introduction

When studying disease processes, the knowledge of alterations on protein level, including protein structure, post-translational modifications or protein-protein interactions, is crucial to elucidate complex biological phenomena. Such insights are difficult to attain due to the enormous complexity of the protein world. Traditional methods for the analysis of proteomes include two dimensional gel electrophoresis and mass spectrometry, however, to compliment the functional analysis of proteins on a large scale, proteins can be studied in array formats. Preferably such arrays would contain functionally active proteins, in all their modified states, immobilised on a surface at high density or in solution in nanowells. Importantly, protein activity is dependant on a wide range of

factors (post-translational modifications, cellular localisation, pH, presence/absence of co-factors, etc.) which makes the production of a protein chip containing the whole proteome a daunting task.

Since their introduction over a decade ago [1], protein arrays have been successfully used in a wide range of studies. Large collections of proteins in an array format enabled identification of peptide-protein interactions [2], protein-protein interactions [3] and antibody-antigen interactions [4]. The specificity of these binding interactions was subsequently confirmed by pull-down, yeast two-hybrid, immunoblot and co-immunoprecipitation experiments. In another approach protein arrays were successfully used for identification of novel substrates for arginine N-methyltransferases [5] and for protein kinases [6]. More recent studies identified 59

\* Corresponding author. Centre for Human Proteomics, Royal College of Surgeons in Ireland, 123 St. Stephens Green, Dublin 2, Ireland. Tel.: +353 1 402 8518; fax: +353 1 402 8551.

E-mail address: [dmurphy2@rcsi.ie](mailto:dmurphy2@rcsi.ie) (D. Murphy).

novel substrates for Ubiquitin-protein ligases (E3s) in yeast [7] and novel substrates for Cyclin-dependent kinase (Cdk5) using commercially available protein microarrays [8,9].

In the field of oncology, protein arrays offer huge potential as tools for the identification of cancer biomarkers, in particular auto-antibody/antigen markers. The identification of tumour-associated antigens (TAAs) recognised by the patient's immune system represents an exciting approach to identify novel diagnostic cancer biomarkers and to contribute towards a better understanding of the molecular mechanisms involved. Circulating autoantibodies have not only been used to identify TAAs as diagnostic/prognostic markers and potential therapeutic targets [10,11], they also represent excellent biomarkers for the early detection of tumours and permit monitoring of the efficacy of treatment [12,13].

Protein array technology has already played a significant role in analyzing autoantibodies in a variety of autoimmune disorders [14-16] and cancers [17,18]. The ability to screen an immune response to a large number of proteins in a high throughput technique cannot only aid in improved diagnosis, but has identified autoantibody signatures that may represent disease subgroups, early diagnostics [19] and facilitate the analysis of vaccine trials [20].

This review investigates the current developments in protein array technology and their applications with particular emphasis on the field of tumour-associated antigen (TAA) discovery.

## 2. General applications of protein arrays

Protein arrays provide a powerful tool to examine interactions between proteins (including antibodies), peptides, DNA/RNA or chemical compounds on a large scale. In one proof-of-principle experiment, a small number of well-defined protein-protein interactions, including an interaction dependent on a small molecule, were demonstrated in a microarray format [21]. As mentioned above, calmodulin- and phospholipids-interacting proteins have been identified by screening almost 6000 yeast proteins, generated by expressing previously annotated open reading frames [22]. Another approach for the identification of protein-protein interactions on protein arrays is the use of short peptides as bait, which can permit studies of proteins that are otherwise difficult to work with in solution. This approach has been successfully applied to identify cytoplasmic interactors of the platelet integrin  $\alpha_{IIb}\beta_3$  [2,23]. Protein arrays also make an ideal platform to characterise antibodies. A human protein expression library, involving in situ expression of tens of thousands of recombinant proteins on large membranes, has been successfully screened to identify antibody-protein interactions [4], permitting identification of cross-reacting proteins and potential epitope definition.

Protein arrays can also be used to determine the target of antibodies previously identified as potentially interesting markers in disease. Our group is presently working to identify the antigens of antibodies, initially identified by immunohistochemistry as interesting markers in certain types of cancer. Each antibody can be screened against an appropriate protein expression library. Unlike previous approaches, which could at best identify potential epitopes of a particular antibody, this

approach identifies the actual target protein. In addition, potential cross-reacting proteins can also be identified using this method, which is important information for assessing an antibody in disease screening procedures. Similarly, protein arrays could provide a useful screening technique to be introduced into any antibody production process.

Just as it is possible to characterize the binding of a single antibody using protein arrays, it is also possible to characterize many antibodies present in a single sample, such as to profile the antibody repertoire in serum or plasma. One immediate application is the use of "allergen arrays" to screen for the presence of particular IgE molecules in a patient sample [24-27]. In a similar fashion, protein arrays can be used to profile antibodies present in the blood of patients with autoimmune diseases. Initial auto-antigen arrays consisted of almost 200 proteins, peptides and other biomolecules (including several forms of dsDNA and ssDNA) which were known auto-antigens to several well characterized auto-immune diseases, including rheumatoid arthritis and systemic lupus erythematosus [28]. Another approach is to use large arrayed libraries of recombinant proteins as a potential auto-antigen array. In a proof-of-principle experiment, a protein array chip containing almost 2500 purified recombinant human proteins was used to profile the auto-antibodies present in small volume samples from patients with alopecia and rheumatoid arthritis [29]. This approach permitted the identification of previously known auto-antigens and also previously uncharacterized protein auto-antigens. Initial results from screening a large recombinant mouse protein array also indicate the usefulness of this approach to characterize auto-immune disease in a mouse model system for systemic lupus erythematosus [14].

## 3. Autoantibody responses in cancer

Antibody responses to tumour associated antigens in cancer patients have been identified in many cancer types and offer new biomarkers with potentially high levels of specificity and sensitivity (see Table 1). The processes by which such self-proteins become immunogenic are not entirely understood [30]. However, the molecular changes in structure or expression of proteins during tumorigenesis suggest mechanisms by which such proteins could be perceived by the immune system as foreign and initiate antibody production.

The most extensively researched mechanisms of cancer immunity are genetic mutations leading to expression of defective tumour suppressor p53. These frequent mutations in many cancer types, lead to conformational changes in the protein. This in turn results in increased stability of the protein and extended half life time of several hours compared with 20 min for wild-type p53 [31,32]. It is generally accepted that p53 mutations are not directly responsible for priming the antibody production, but rather that it is caused by the high antigenic load resulting from the accumulation of the p53 protein [33]. The disparity between the mutation site and the actual epitope was also observed for Ras p21 in cancer patients [34]. Interestingly, in both cases the accumulation of the protein is required for the antibody response, however, the accumulation of the protein does not always result in antibody

**Table 1 – Examples of both protein (non-antibody) and autoantibody cancer markers**

Protein (non-antibody) biomarker	Swiss-prot ID	Cancer type	Specificity	Sensitivity	Reference
Prostate-specific antigen	PSA	Prostate cancer	92%*	12%*	[87]
Carcinoembryonic antigen	CEA	Colorectal cancer	90% **	34% **	[88]
Cancer antigen CA15-3	CA15-3	Breast cancer	69%	23%	[89]
Cancer antigen CA19-9	CA19-9	Gastrointestinal cancer	86%	35%	[90]
Cancer antigen CA125	CA125	Ovarian cancer	41.5%***	80%***	[91]
Autoantibody biomarker	Swiss-prot ID	Cancer type	Specificity	Sensitivity	Reference
cAMP-dependent protein kinase A	ECPKA	Several cancer types	90%	87%	[45]
Annexin XI-A	ANXA11	Breast cancer	88%	77%	[92]
Tumour suppressor p53	p53	Several cancer types	96%	30%	[93]
Huntingtin interacting protein 1	HIP-1	Prostate cancer	73%	46%	[94]

\*3.1–4.0 ng/ml PSA level \*\*5 ng ml<sup>-1</sup> CEA level \*\*\*330 U/mL CA125.

Autoantibody based biomarkers show promising results and are currently a focus of cancer diagnostic research.

production. Of all cancer patients with p53 mutations, approximately half have detectable serum antibodies specific to the p53 protein.

A recent study by Engelhorn and colleagues investigated the direct potential of mutations to induce antibody-mediated immune responses [35]. The group found in mice experiments, using combinatorial DNA libraries encoding large numbers of random mutations in tyrosinase-related proteins, that truncations of the tyrosinase protein are sufficient to elicit antibody production. Additional amino acid substitutions in the protein can further enhance these immune responses. Antibody responses to mutated gene products were also identified at advanced stages of cancer, including in a mouse model [36]. Since thousands of genes can mutate during the progression of cancer [37], these findings demonstrated the potential of stage-specific immune responses.

An alternative pathway leading to initiation of antibody production is the deregulation of gene expression. The expression of many genes in cancer is deregulated and many genes are overexpressed [38]. For example, the non-secreted surface antigen GA733-2 is over-expressed in over 90% of colorectal cancers and anti-GA733-2 antibodies were detected in 14.5% of the patients [39]. A further example is the HER-2/neu protein, which is overexpressed in approximately 20% of colorectal cancer patients. HER-2/neu antibodies were detected in 46% patients overexpressing the protein and in only 5% patients with no detectable HER-2/neu [40]. The overexpression of genes may lead to an increased antigenic load which in return can be sensed by the immune system and lead to antibody production.

Aberrant cellular localization of antigens caused by malignant transformation of mammalian cells might represent a further potential pathway triggering autoantibody responses in cancer patients. The cAMP-dependent protein kinase A (PKA) is an intracellular enzyme [41,42] which can be secreted by cancer cells into the conditioned medium [43,44]. This PKA, designated as extracellular protein kinase A (ECPKA), is often found to be up-regulated in the serum of cancer patients [43,44] which correlates with the presence of ECPKA specific autoantibodies in sera of corresponding patients [45].

A further potential mechanism by which self-proteins could trigger antibody production is expression of genes that are not usually expressed in a particular tissue. The mRNA binding

protein p62 is normally expressed in the foetus and its expression is silenced in adult tissue. In some cases of hepato-cellular carcinoma, p62 expression can be reinstated leading to recognition by the immune system and antibody production [46]. Another example is a group of proteins normally expressed in testicular tissue and not recognised by the immune system as foreign (e.g.: NY-ESO-1, SSX2). When, however, aberrantly expressed in tumour cells, these proteins can trigger antibody responses [47].

#### 4. Cancer autoantibody discovery in the proteomic age

The vast amount of potential molecular changes during tumorigenesis in different cancer types and the individual variances in immune responses in cancer patients potentially result in complex antibody repertoires. In order to identify a disease specific antibody profile it is required to analyse as many potential antigens as possible.

Classical laboratory methods, such as Western blotting or enzyme-linked immunosorbent assays (ELISA), have been used to identify a number of tumour antigens. Antibodies to p53, p21 ras or Her/2neu were successfully identified in cancer patients in the previous two decades [34,48,49]. Several novel approaches have been developed to detect new antigens that elicit antibody responses. With the advent of proteomics and recombinant technologies, it is now possible to characterise hundreds of proteins in parallel for potential antibodies in sera of cancer patients. Followed by a short description of methods based on 2D gel electrophoresis, mass spectrometry and phage display we will focus our review on protein arrays as a tool for detection of antibodies associated with cancer.

Proteomics is an alternative approach for identification of novel tumour antigens. Proteomics applies two dimensional gel electrophoresis (2DE), a process that separates large mixtures of proteins from natural sources such as tissue or cell lines. Once the proteins are separated using 2DE and transferred onto protein binding membranes using Western blotting, the autoantibodies can be identified by incubation with serum of cancer patients [50,51]. Although the natural protein source allows analysis of proteins in post-translationally modified states, a disadvantage of 2DE is that very large or

260 very small proteins and certain kinds of proteins such as  
261 membrane proteins, are difficult or impossible to visualize  
262 using 2DE and require further characterisation using mass  
263 spectrometry [52].

264 An alternative proteomics approach combines liquid phase  
265 protein separations with microarray technology. Proteins in  
266 cell and tissue lysates can be separated using chromatography  
267 and hundreds of lysate fractions can be arrayed onto  
268 nitrocellulose-coated slides [53]. Incubation of serum samples  
269 with lysate arrays may reveal fractions immunoreactive with  
270 serum antibodies. A study by Nam and colleagues used a  
271 fractionated colon adenocarcinoma cell line and identified  
272 ubiquitin C-terminal hydrolase isozyme 3 as an antigen  
273 eliciting antibody responses in a group of colon cancer  
274 patients [54]. The merit of post-translationally modified  
275 antigen detection is reduced by difficulties in characterising  
276 the identity of antigens, since each fraction may still contain a  
277 complex mixture of proteins.

278 These proteomics approaches represent a large step  
279 towards the identification of novel antigens eliciting antibody  
280 responses in cancer patients. However, identification of  
281 immune responses to a broad set of antigens in cancer is  
282 limited to individual and often abundant proteins. Without  
283 doubt, the technical advances in separation methods, image  
284 analysis and mass spectrometry will retain proteomics as a  
285 vital tool for antibody identification.

286 In recent years various protein array techniques combined  
287 with more classical research methods have been successfully  
288 applied to identify cancer associated autoantibody markers.  
289 Novel autoantibodies were identified in sera of hepatocellular-  
290 carcinoma patients using 2DE proteomic method. The validity  
291 of these findings was confirmed using a protein array  
292 consisting of the corresponding antigens [55]. Generation of  
293 protein arrays based on protein sources from malignant tissue  
294 samples is another potentially important step due to advances  
295 in protein fractionation methods and mass spectrometry  
296 supported by supervised learning analysis of gathered data.  
297 Two recent studies focusing on lung and prostate cancer  
298 utilized 2D gel electrophoresis and liquid phase chromato-  
299 graphy methods respectively as means for generation of  
300 highly specialised protein arrays for autoantibody discovery  
301 [56,57]. A growing number of current studies strongly indicate  
302 that protein arrays are emerging as a tool of choice for  
303 research towards deciphering complex protein-protein inter-  
304 actions, in particular for the discovery of novel autoantibody/  
305 autoantigen biomarkers. Such work has enormous potential in  
306 the discovery and development of panels of such biomarkers,  
307 which appear to offer superior accuracy in disease diagnosis  
308 compared to individual markers [58].

309 Phage display offers a powerful platform for identification of  
310 tumour associated antibodies [59]. Serological analysis of phage  
311 libraries was first introduced in 1995 by Sahin and colleges [60].  
312 Phage display technology allows presentation of peptides and  
313 proteins derived from human cDNA libraries on the surface of  
314 bacteriophage and testing these for reactivity with serum  
315 antibodies. Individual immuno-reactive clones can be enriched  
316 in a multi-step biopanning procedure from the random phage  
317 libraries and characterized by sequencing. A great advantage of  
318 phage display is the very large number of potential antigens  
319 encoded by the cDNA libraries. This enables detection of a wide

range of cancer-specific antibodies/antigens [61]. However, 320  
phage display technology is labour-intensive and peptides 321  
expressed by phages are often generated from untranslated 322  
DNA sequences and do not correspond to native antigens, thus 323  
limiting identification of molecular targets in cancer [11,62]. 324

Recent technical advances enable generation of phage-based 325  
protein/peptide arrays containing several hundred up to several 326  
thousand phages [11]. Screening the initial phage library with 327  
pooled sera from several patients and pooled sera from non- 328  
cancer controls identified in a multi-step biopanning procedure 329  
approximately 2000 immunoreactive clones. These clones were 330  
then arrayed on glass-slides and re-screened for immuno- 331  
reactive clones with non-pooled serum samples. The phage 332  
array technology allowed the identification of two panels of 333  
antigens specific for prostate and lung cancer [11,63]. Both 334  
panels were able to predict cancer with over 80% sensitivity and 335  
specificity. 336

The success rates of phage based screening methods were 337  
critically improved with the introduction of protein arrays 338  
consisting of antigens identified in concluding steps of phage 339  
selection [11]. These recent developments permit screening to 340  
focus on several hundred to a few thousand identified leads 341  
and validate these in larger cohorts of patients using protein 342  
arrays [63]. 343

## 5. Generation of protein arrays for profiling autoantibody repertoires in cancer 346

The first requirement towards the generation of protein arrays 347  
is a source of large numbers of recombinant proteins. A number 348  
of strategies are currently employed to provide sources of 349  
thousands of proteins for the generation of protein arrays. One 350  
approach is high-throughput cloning or amplification using PCR 351  
of defined open reading frames coding for the proteins of 352  
interest [64,65]. The successful implementation of such an 353  
approach relies on the availability of sequence data and its 354  
correct annotation in the databases. In particular, the definition 355  
of the open reading frames of alternative splice variants of one 356  
protein remains difficult with such an approach. Also, pre- 357  
viously uncharacterised proteins will be absent, limiting this 358  
approach as a discovery tool. For these reasons, this approach 359  
has proved most valuable in the production of chips containing 360  
proteins from well characterized organisms, such as *Arabidopsis* 361  
[64] and *Caenorhabditis elegans* [66]. 362

Another approach is the use of protein expression libraries, 363  
which are used for a "shotgun" approach to generate 364  
recombinant proteins [1]. Such libraries are generated using 365  
mRNA isolated directly from the cell [67,68]. Here, mRNA is 366  
isolated from the tissue or organism of interest and direction- 367  
ally sub-cloned into a protein expression vector, suitable for 368  
heterologous expression in either a bacterial (e.g. *E. coli*) or 369  
eukaryotic organism (e.g. yeast). Several solutions exist for the 370  
rapid movement of coding region from one organism to 371  
another, such as the GATEWAY system (Life Technologies) 372  
[69] or dual expression vectors [67]. Not only does this 373  
approach circumvent the cloning of individual open reading 374  
frames, readily permitting the expression of tens of thousands 375  
of proteins, but also this strategy automatically includes splice 376  
variants and previously uncharacterized gene products. 377

378 The Gateway technology has been successfully applied for  
379 cloning the Ultimate™ ORF Clone collection resulting in  
380 generation of human protein expression clones for protein  
381 array applications. Recombinant bacmid DNA were isolated  
382 and transfected into Sf9 insect cells in order to generate a  
383 recombinant baculovirus that was used for expression experi-  
384 ments. A major advantage of this approach is the generation  
385 of protein arrays containing native proteins which are post-  
386 translationally modified [18].

387 Apart from just the question of expressing the recombinant  
388 proteins of choice in a particular host system, there is also the  
389 question of tagging these proteins. A large number of tags  
390 exist, the coding region of which can readily be introduced  
391 into vectors. Depending on the approach taken, many options  
392 are available, including N-terminal and/or C-terminal tagging,  
393 multiple different tags are also an option, in particular in  
394 conjunction with proteolytic sites for subsequent removal of  
395 the tag, if required. These tags can vary in size from the short  
396 His<sub>6</sub> tag to the 30 kDa GFP tag. Such tags are required for the  
397 detection of the expressed recombinant proteins and, in  
398 conjunction with an appropriate affinity separation method,  
399 for their purification. For a review of different expression  
400 systems see [70].

401 In one example, a human foetal brain cDNA library was  
402 subcloned into a bacterial expression vector, permitting  
403 controlled IPTG-inducible expression of His<sub>6</sub>-tagged recombi-  
404 nant proteins [1]. The *E. coli* clones of this library were then  
405 arrayed in high density onto PVDF membranes, grown over-  
406 night and recombinant protein expression induced for a  
407 controlled length of time. Those clones expressing a human  
408 protein are readily detected by means of an anti-His<sub>6</sub> antibody.  
409 In this fashion, a protein array containing 10–12,000 different  
410 human proteins has been generated. Approximately 66% of  
411 the proteins expressed in this library are, according to present  
412 annotation, full length. Also, such proteins have been shown  
413 to be readily expressed and purified in high-throughput using  
414 available automated platforms [29,71].

415 In the recent years several novel methods have been  
416 developed for cell free *in situ* synthesis of protein arrays [72–75].  
417 *In situ* protein array synthesis uses cell free expression systems  
418 to produce proteins directly from nucleic acid templates. The  
419 major advantage of these methods is the expression of many  
420 proteins in parallel on a chip, avoiding labour intensive and often  
421 costly processes of DNA cloning, expression and protein  
422 purification. The *in situ* cell free protein generation methods  
423 show the capability of high-yield and high-throughput synthesis  
424 of hundreds of proteins with minimal variation and good  
425 reproducibility [76]. In addition, these methods demonstrate  
426 intrinsic purity of synthesized proteins [74] which makes them a  
427 desirable tool for TAA detection from complex and polyclonal  
428 antibody mixtures present in blood of patients. The high purity  
429 of proteins arrayed on such self-assembling chips encourages  
430 applications in cancer immunology, as outlined in a recent  
431 breast cancer study. The findings show that the identified  
432 autoantibody profiles are less prone to false positive results due  
433 to highly pure proteins directly assembled on the chip [77]. An  
434 extensive review on cell free protein array synthesis methods  
435 was recently published by He et al [78].

436 Once a source of proteins has been established, considera-  
437 tion must be given to the format that a protein microarray can

438 have. Presently, there is a number of possible surfaces  
439 available for the manufacture of protein arrays. In general,  
440 chip surfaces can be divided into two major categories: flat  
441 (planar), and 3D (mostly gel-like) surfaces [79,80]. Planar  
442 surfaces have been primarily developed in the area of cDNA  
443 arrays. The glass surface of these chips is usually treated to  
444 produce a thin layer of a particular chemical group, e.g. an  
445 aldehyde group or poly- $\epsilon$ -Lysine [81]. The proteins are bound  
446 to the surface of these chips by either covalent bonds or simple  
447 electrical charge. Another type of planar chip surface available  
448 is plastic polymer coated slides, such as the MaxiSorb slides  
449 from Nunc – an approach used in ELISAs for many years. The  
450 simple adoption of technology from the area of cDNA arrays  
451 brings with it a number of major concerns when used with  
452 proteins. Firstly, unlike DNA, the surface charge of proteins is  
453 highly variable and the use of simple uniform electrostatic  
454 interaction for the immobilisation of different proteins results  
455 in large variation in the amount of protein bound. Secondly,  
456 the structural conformation of proteins deposited onto a  
457 planar surface cannot be expected to very closely mimic  
458 “native” proteins, but would be more similar to proteins  
459 present on membranes, such as in traditional Western  
460 blotting or dot blot experiments. Thirdly, there is no control  
461 over the orientation of the proteins on the surface, which may  
462 result in, for example, the inaccessibility of an active site.

463 The development of the gel-like 3D chip surface was partly  
464 driven in an attempt to minimise the denaturation of the  
465 immobilised proteins in the arrays. A number of such surfaces  
466 exist based around polyacrylamide and agarose coated on a  
467 glass surface, which provide a hydrophilic environment for  
468 the proteins. Such surfaces allow the user to adjust various  
469 conditions, such as pH and salt concentrations, by incubating  
470 the chips in the appropriate buffer. Using a “home made”  
471 polyacrylamide surface, a glass chip containing 2413 non-  
472 redundant purified human fusion proteins, arrayed at a  
473 density of up to 1600 proteins/cm<sup>2</sup>, has been successfully  
474 employed in antibody binding studies, including the screening  
475 of human serum samples [29], indicating the maintenance of a  
476 degree of structural conformation of the proteins involved.  
477 There are also non-gel-like 3D surfaces available, such as the  
478 FAST slides from Schleicher and Schuell which have a  
479 nitrocellulose surface. Like any 3D surface, these surfaces  
480 allow a much higher concentration of proteins per spot.

481 However, these surface solutions still leave the problem of  
482 controlling the orientation of the proteins on the surface of the  
483 slides, which is necessary to maximise the activity of these  
484 proteins. For example, in order to maximise the binding of  
485 antibodies arranged on a surface to their epitopes, it would  
486 clearly be an advantage if the heavy chain were attached to, or  
487 close to, the surface and the antigen binding site as far away  
488 from the surface as possible. One approach has been the use of  
489 affinity tags, for example a nickel-coated slide has a natural and  
490 specific affinity to His<sub>6</sub>-tagged recombinant proteins. This  
491 approach was used to array 5800 yeast proteins which were  
492 screened for their ability to interact with calmodulins and  
493 phospholipids [22]. Similarly, successful orientation of antibo-  
494 dies and antibody Fab fragments was achieved by biotinylating  
495 such antibodies/Fab fragments and arraying them on a strepta-  
496 vidin-coated surface [82]. A further development of surface  
497 chemistry involves the use of a polyethylene glycol layer (PEG)

[80] or dendrimers [83]. These approaches involve the coupling of the proteins to epoxy groups, which act as spacers preventing direct protein-surface contact and, thus, eliminate the need for blocking reagents to reduce background binding. One further development of this approach has been to link chelating iminodiacetic acid groups to PEG, which in turn can be bound by  $\text{Cu}^{2+}$  ions and so provide a highly specific binding site for His<sub>6</sub>-tagged proteins [72]. A number of studies have been carried out comparing the different surfaces available for protein array work, including antibody arrays which assess background noise, sensitivity/detection limits, reproducibility and storage for a variety of experimental designs [79,80].

While the development of 3D surfaces and spacers partially addresses the problems of protein-protein interactions on a chip, many experiments that involve interactions of proteins in a functional state will prove difficult to perform on these chips, which is obviously a major drawback of the current protein arrays. Ideally, we would like to look at protein interactions, where the proteins are in their native state and are functional, i.e. in conditions as close as possible to those in nature. This may be solved by developing a microfluidic chip, which is a series of enclosed micro-channels within a chip format, such as silicon, plastic or glass. The potential of microfluidic chips would include the ability to maintain proteins in their functional conformations and to therefore perform interactions such as protein-protein, protein-peptide, protein-compound, protein-DNA, protein-ligand and protein-antibody interactions in solution. Also, the area of enzyme studies would greatly profit from such a system. In fact the first steps have been taken to use a "lab-on-a-chip" to study some of the reactions in the glycolytic pathway of yeast. Using this system, enzymatic reactions in volumes as low as 6.3–8 nL could be studied [84]. The ability to use such small volumes in microfluidic chips is very important due to the high cost of proteins, antibodies, compound libraries, peptides and even the difficulties in obtaining enough patient samples to screen in high-throughput. Another exciting innovation is the first attempts to establish a lipid bilayer membrane chip that would allow the functional investigation of transmembrane proteins and their interactors [85]. These advances would also permit a more thorough exploitation of the libraries of proteins that currently exist.

The antibody repertoires in humans consist of a complex polyclonal mixture of antibodies with a wide range of specificities, resulting in immunoreactivity with a vast number of potential antigens. Consequently, minute impurities of a protein sample on a protein array may result in a non-specific false-positive signal. To assure the analytical quality of protein microarray experiments, the protein array production and antibody profiling experiments require highest standard and quality. Appropriate quality control and assurance measures are extensively described in an review article by Kricka and Master [86].

## 6. Conclusion

The understanding of molecular events leading to cancer is of the utmost importance to identify effective diagnostic and therapeutic targets. With the complexity of cells and organisms a genome-wide gene expression and protein analyses are

essential to elucidate human diseases in terms of dysfunctions of molecular systems. This is especially true for cancer, where multiple key molecular alterations may destabilize essential molecular pathways setting off an avalanche of associated modification on molecular and cellular levels leading to malignancy. As outlined in this review the humoral immune responses to tumour antigens offer an exciting opportunity to identify autoantibody based diagnostic biomarkers. The immune system may serve as a sensitive tool to detect minute amount of cancer specific antigens amplifying autoantibody responses to a measurable level. However, in addition to the complexity of cancer, the vast quantity of antibodies circulating in human blood and the diversity between individuals needs to be taken into account. In this context, individual patients produce an immense repertoire of antibodies of which many might be not cancer related. Therefore, to comprehensively study autoantibodies in patients and to identify cancer specific autoantibody signatures the research relies on large sets of proteins which can serve as potential antigens. The recent developments in protein array technology deliver an excellent tool for analysis of thousand of proteins for their antigenic ability. Supported by current findings, a major future application of high-content protein arrays will contribute to the identification of novel of tumour associated autoantibodies as diagnostic markers for cancer.

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