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Rapid identification of molecular subtyping of breast cancer cell lines using a multi-channel sensor array†

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In this paper, we present a multi-channel sensor array composed of three fluorescent probes that can be used to identify and differentiate breast cancer molecular subtypes, including luminal A, luminal B, HER2-enriched and triple-negative. The multi-channel sensor array allows the three probes to mix and interact with each cell sample in one well simultaneously, resulting in high efficiency, lower sample costs, and 100% accuracy in cell molecular subtype differentiation.

The use of sensor arrays to mimic mammalian taste and olfactory systems has gained considerable attention in research fields of analytical chemistry. The concept of “chemical tongues” or “electronic noses” was proposed as an alternative to traditional highly specific “lock-and-key” recognition systems.^{1–3} Array-based sensing relies on non-specific or semi-selective interactions between multiple sensor units and analytes that may lead to pattern recognition of complex mixture samples.^{2,4} Array-based sensing strategies have been used in discrimination of volatile organic compounds, metal ions, amino acids, proteins, food additives, beverages and so on.² In recent years, sensor arrays have shown potential applications in analyzing complex biological systems. For example, a three-unit fluorescence sensor array has been used to discriminate proteins in urine samples for urinary disease diagnosis in our previous work;⁵

Rotello's group has reported using different designs of sensor arrays, including fluorescence nanoparticle–polymers,⁶ ratiometric conjugated polymer–fluorescent proteins,⁷ and quantum dot-gold nanoparticle sensor arrays,⁸ to differentiate cell types and phenotypes, and macrophage phenotypes;^{9,10} other reports include stem-cell differentiation,¹¹ bacteria identification,^{12–17} microbial pathogen detection,^{18–20} recognition of gut microbiota,²¹ etc.^{22–24}

Lipids, carbohydrates, and several membrane proteins make up the membrane of cells.²⁵ In the course of carcinogenesis, the lipid composition of the membrane varies,²⁶ membrane receptors are overexpressed,²⁷ and surface proteins are glycosylated,²⁸ resulting in varying molecular patterns. Malignancy types and molecular subtypes also differ in the composition of cancer cell membranes. Breast tumours can be classified into four molecular subtypes based on gene expression patterns, namely luminal A (high expression of estrogen receptor), luminal B, HER2-enriched (human epidermal growth factor receptor 2-enriched), and basal-like (also known as triple-negative).²⁹ Different subtypes result in different clinical behaviours, which are crucial to the development of targeted therapies and to determining the prognosis of breast cancer (BC). In clinical settings, determining the intrinsic molecular subtype of BC requires immunohistochemistry or gene expression profiling, both of which are costly, time-consuming, and not widely available.

Recently, we reported the development of a 12-unit sensor array consisting of 12 fluorescent probes, including conjugated polyelectrolytes, fluorophore-labelled peptides, and monosaccharides/glycans for the differentiation of BC exosomes.³⁰ In contrast to a natural scenario, mammalian taste and olfactory systems actually “collect and analyze” responses delivered by different receptors within one breath of air or bite of food. This sensor array works on the principle of 12 different probes responding separately to each sample, and the response pattern is then analyzed through linear discriminant analysis (LDA), hierarchical cluster analysis (HCA) and a deep learning algorithm.

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Scheme 1 (a) Schematic illustration of identification of molecular subtyping of BC cells using a three-channel sensor array. (b) Chemical structures of the three sensor units used in this study. (c) The illustrative comparison of multi-well vs. one-well analysis.

Large numbers of measurements have limited the use of sensor arrays composed of many sensor units in biological analysis. A “one-well analysis” method was developed to overcome this limitation, which combines multiple probes with the sample in one well, instead of measuring with each probe separately. Herein, we report a simple and efficient one-well three-channel fluorescent sensor array for rapid identification of molecular subtyping of BC cell lines (Scheme 1).

To minimize the number of probes and ensure identification of subtle differences between diverse BC subtypes, three recognition units have been selected to form the multi-channel sensor array: RGD-FITC, UNO-FITC and Gly-DCM (glycan-fluorophore conjugate). RGD-FITC contains an arginine-glycine-aspartic acid (RGD) tripeptide sequence. RGD can preferentially bind to integrin $\alpha\beta_3$, which is overexpressed on tumor endothelial cells as well as on some tumor cells such as BC, liver cancer and so on. RGD-based probes have been used for molecular imaging of BC.³¹ The recognition unit of UNO-FITC is a tumor homing peptide codenamed “UNO” that targets CD206 (a specific mannose binding receptor) on cells. Previously, it has been reported that UNO can target 4T1 and MCF7 breast carcinoma.³² Gly-DCM has a sialylated glycoconjugate, which binds to immune regulatory receptors (sialic acid-binding immunoglobulin-type lectins) predominantly found on cells.³³ UNO and RGD have relatively high recognition specificity, while the glycan probe can cross-respond to a wider range of analytes. The combination of different interaction patterns is beneficial to maximize the recognition effect while reducing the number of probes.

Initial studies examined the rationality of probe selection. Through flow cytometry (FCM), we first validated the discriminatory efficiency of three probes against a variety of

cell lines: MCF10A (non-tumorigenic epithelial cell line), T47D (human BC cell line), HeLa (cervical cancer cell line), and HepG2 (liver carcinoma cell line). FCM provides a means of analyzing cell populations using different fluorescent stains. Using the probes of RGD-FITC, UNO-FITC, and Gly-DCM, a data matrix containing three probes, four cell types, and six replicates was generated. The results of the FCM analysis (Fig. 1a) provide different cell staining percentages with patterns. A number of machine learning methods have been developed to identify BC cell lines.^{34,35} Both LDA and principal component analysis (PCA) were used to analyze the experimental data in this study. As shown in Fig. 1b, LDA transformed raw fluorescence staining data into discriminant plots. Using leave-one-out cross-validation, four data groups clustered with 100% accuracy. PCA is an unsupervised learning algorithm (see Fig. S4†). In our study, LDA maximizes the signal-to-noise ratio of the data, allowing better differentiation between categories.

The same four cell lines and three probes were used to conduct the sensing experiment using a microplate reader to acquire fluorescence spectral data. Fig. 1c illustrates the fluorescence spectra, demonstrating details on individual intensities at different wavelengths. This provides a more advantageous approach than FCM in that FCM provides limited information on cell staining percentages and is relatively time-consuming to prepare samples. Meanwhile, fluorescence spectrometers and plate readers are more accessible than flow cytometers in most laboratories. Therefore, it is possible to collect high-dimensional fluorescence spectral data for the following data analysis. According to the LDA score plot (Fig. 1d), the four clusters were distributed in two-dimensional space without overlap and the accuracy is 100% with leave-one-out cross-validation.



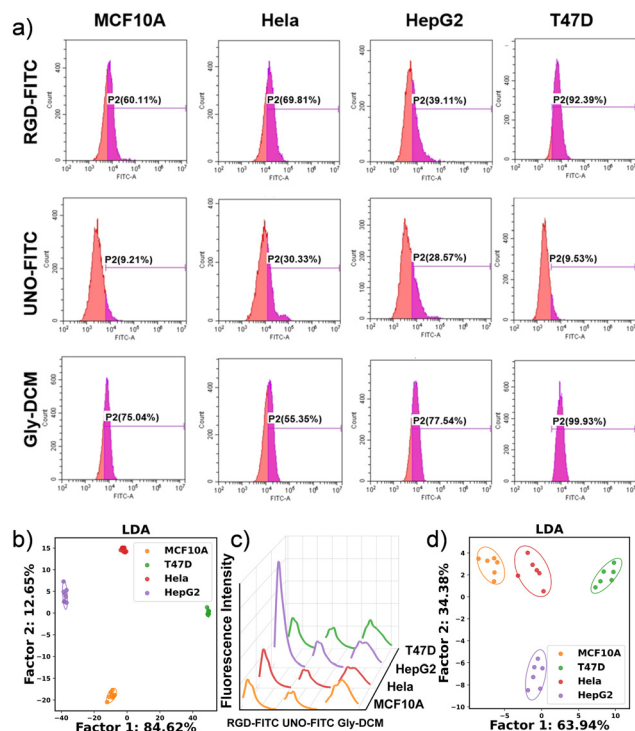


Fig. 1 Differentiation of normal cells and cancer cells using a multi-well sensor array. (a) FCM results and (b) LDA plot of the sensor array with MCF10A, HeLa, HepG2, and T47D, respectively. (c) Fluorescence spectra and (d) LDA plot of the sensor array with MCF10A, HeLa, HepG2, and T47D, respectively.

FCM and fluorescence spectral analysis were used to differentiate multiple cell lines, including one normal cell line MCF10A, and six BC cell lines, MDAMB231, MDAMB436, BT474, SKBR3, T47D, and MCF7 (Fig. 2). Both the FCM and fluorescence spectral analysis were successful in accurately classifying the samples, indicating the potential of the three-unit array for future applications.

In most sensor arrays reported in the literature, each sensor unit tests the sample individually, and all the separate signals are then collected and analyzed, whereas in mammalian olfactory and taste systems, all olfactory cells or taste buds respond to odors or tastes in concert and transmit a collective signal to the neural system for analysis. As a consequence, arrays with a large number of sensors require large quantities of samples and are labour-intensive and time-consuming. In order to observe the fluorescence from the three probes in one well, we modified the probes' recognition units with different coloured fluorophores. The chemical structures of the three modified probes are illustrated in Scheme 1b. We use Cy5 and TAMRA to modify RGD-peptide and UNO-peptide, to form RGD-Cy and UNO-TAMRA, respectively. Cy5 and TAMRA have strong light absorption and high quantum yields, which make them commonly used fluorophores. TAMRA fluoresces at 582 nm, whereas Cy5 fluoresces at 670 nm. Two glycine spacers (GG) were added between the 5-TAMRA unit and the circular UNO



Fig. 2 Differentiation of one normal cell line and six BC cell lines using a multi-well sensor array. (a) FCM results and (b) LDA plot of the sensor array with a normal cell line MCF10A and six BC cell lines. (c) Fluorescence spectra and (d) LDA plot of the sensor array with MCF10A and six BC cell lines.

peptide sequence. Therefore, the three-channel sensor array is equipped with a yellow probe Gly-DCM, a red probe UNO-TAMRA, and a green probe RGD-Cy.

Although the modified probes on the fluorophore share the same recognition units, they differ from the original probes in reducing overlap among fluorophores and maximizing the amount of information collected across multiple detection channels while maintaining the cross-response capability of the original probes. Photophysical characterization of the probes is included in Fig. S1, ESI†. Gly-DCM exhibits a relatively wide spectrum of absorption and emission ($\lambda_{\text{ex}} = 456$ nm; $\lambda_{\text{em}} = 620$ nm). RGD-Cy ($\lambda_{\text{ex}} = 600$ nm; $\lambda_{\text{em}} = 673$ nm) and UNO-TAMRA ($\lambda_{\text{ex}} = 520$ nm; $\lambda_{\text{em}} = 582$ nm) both exhibit strong and narrow emission signals.

The three probes were then mixed together in one microwell and their spectra were measured. In the mixed system, Gly-DCM and UNO-TAMRA showed a slight change in spectral shape, which could be explained by the Förster resonance energy transfer (FRET) effect between the probes (Fig. S2, ESI†). FRET results in both a decrease in the fluorescence of the donor molecule and an increase in the fluorescence of the acceptor molecule. Some references have demonstrated that the use of FRET fluorophore pairs in a sensor array can provide more spectral information for pattern recognition.^{7,36} We then examined the fluorescence spectra of the sensor array with MCF10A and three tumor cell lines (Fig. 3a). As part of a one-well analysis, all probes were placed in one microwell in order to interact with the cell sample. Fluorescence spectra were then collected from each well at corresponding bands. A one-well analysis was able to discriminate four cell lines with 100% accuracy (Fig. 3b), demonstrating the feasibility of using multi-channel sensor arrays.



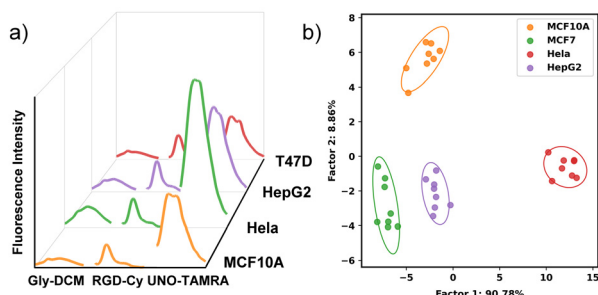


Fig. 3 Differentiation of a normal cell line and three cancer cell lines using the multi-channel sensor array. (a) Fluorescence spectra and (b) LDA plot of the sensor array with MCF10A, HeLa, HepG2, and T47D, respectively.

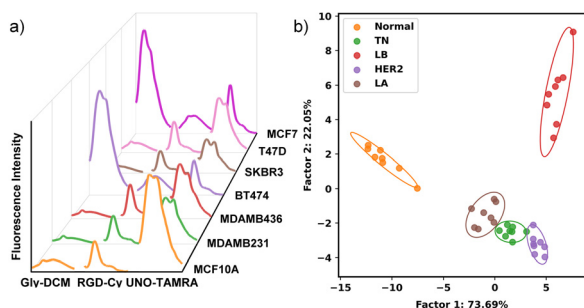


Fig. 4 Differentiation of molecular subtypes of BC cell lines using the multi-channel sensor array. (a) Fluorescence spectra and (b) LDA plot of the sensor array with one normal cell line and six BC cell lines.

We further investigated the discrimination of different BC subtypes using a multi-channel sensor array with six different BC cell lines and one normal cell line. Based on the LDA result, these 7 cell lines can be divided into five groups: two LA subtype cell lines, T47D and MCF7, are grouped together; two TN cell lines, MDAMB 231 and MDAMB436 are grouped together; while MCF10A, SKBR3, and BT474 are grouped separately (Fig. 4b and Table S1, ESI†). As a heterogeneous disease with multiple subtypes, BC poses many treatment challenges. In these results, we demonstrate the capability of our multi-channel sensor array to differentiate molecular subtypes of BC cell lines.

In conclusion, a three-unit fluorescence sensor array for accurate discrimination of molecular subtypes of BC cells has been designed and constructed. With the multi-channel sensor array, three probes can mix and interact with each sample simultaneously, resulting in high efficiency, lower sample costs, and 100% accuracy in cell subtype differentiation. Moreover, due to its high sensitivity and excellent ability to recognize biological systems, this sensing platform might be well suited to more complex biological systems.

Author contributions

Yuyao Jin: data analysis, validation, and investigation. Yuanjie Sun: data analysis, validation, investigation, writing –

original draft and review. Nan Du: validation, investigation, and data analysis. Wei-Tao Dou: resources. Ying Tan: formal analysis. Xiao-Peng He: resources. Naihan Xu: formal analysis and funding acquisition. Chunyan Tan: conceptualization, supervision, resources, writing – review & editing, methodology and funding acquisition.

Conflicts of interest

There are no conflicts to declare.

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