Breast Cancer Diagnosis Using a Microfluidic Multiplexed Immunohistochemistry Platform

Minseok S. Kim¹, Taemin Kim², Sun-Young Kong³, Soim Kwon⁴, Chae Yun Bae¹, Jaekyu Choi¹, Chul Hwan Kim⁵, Eun Sook Lee⁴*, Je-Kyun Park¹,6*

¹Department of Bio and Brain Engineering, Korea Advanced Institute of Science and Technology (KAIST), Daejeon, Republic of Korea, ²Department of Electrical Engineering and Computer Science, Korea Advanced Institute of Science and Technology (KAIST), Daejeon, Republic of Korea, ³Department of Laboratory Medicine, Research Institute and Hospital, National Cancer Center, Goyang, Republic of Korea, ⁴Department of Breast and Endocrine Surgery, College of Medicine, Korea University, Seoul, Republic of Korea, ⁵Department of Pathology, College of Medicine, Korea University, Seoul, Republic of Korea, ⁶KAIST Institute of NanoCentury, Daejeon, Republic of Korea

Abstract

**Background:** Biomarkers play a key role in risk assessment, assessing treatment response, and detecting recurrence and the investigation of multiple biomarkers may also prove useful in accurate prediction and prognosis of cancers. Immunohistochemistry (IHC) has been a major diagnostic tool to identify therapeutic biomarkers and to subclassify breast cancer patients. However, there is no suitable IHC platform for multiplex assay toward personalized cancer therapy. Here, we report a microfluidics-based multiplexed IHC (MMIHC) platform that significantly improves IHC performance in reduction of time and tissue consumption, quantification, consistency, sensitivity, specificity and cost-effectiveness.

**Methodology/Principal Findings:** By creating a simple and robust interface between the device and human breast tissue samples, we not only applied conventional thin-section tissues into on-chip without any additional modification process, but also attained perfect fluid control for various solutions, without any leakage, bubble formation, or cross-contamination. Four biomarkers, estrogen receptor (ER), human epidermal growth factor receptor 2 (HER2), progesterone receptor (PR) and Ki-67, were examined simultaneously on breast cancer cells and human breast cancer tissues. The MMIHC method improved immunoreaction, reducing time and reagent consumption. Moreover, it showed the availability of semi-quantitative analysis by comparing Western blot. Concordance study proved strong consensus between conventional whole-section analysis and MMIHC (n = 105, lowest Kendall’s coefficient of concordance, 0.90). To demonstrate the suitability of MMIHC for scarce samples, it was also applied successfully to tissues from needle biopsies.

**Conclusions/Significance:** The microfluidic system, for the first time, was successfully applied to human clinical tissue samples and histopathological diagnosis was realized for breast cancers. Our results showing substantial agreement indicate that several cancer-related proteins can be simultaneously investigated on a single tumor section, giving clear advantages and technical advances over standard immunohistochemical method. This novel concept will enable histopathological diagnosis using numerous specific biomarkers at a time even for small-sized specimens, thus facilitating the individualization of cancer therapy.

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* E-mail: esilee@korea.ac.kr (ESL); jekyun@kaist.ac.kr (JKP)

Introduction

Accurate prediction and prognosis are the most critical and difficult issues in breast cancer treatment. Because breast cancer, a leading cause of cancer death in women, is a heterogeneous disease that has several biological subtypes, single biomarker test is insufficient to predict the clinical outcome of individual neoplasms [1,2]. Many potential biomarkers with clinical value have been identified through advances in genomics, proteomics, and molecular pathology [3], and they have facilitated various kinds of personalized therapy for cancer patients [4]. However, this transition toward personalized therapy will require novel analytical methodology for accurate prediction and prognosis, particularly multiplex analysis [5,6]. For example, genomic techniques such as DNA microarray analysis (examining 4968 significant genes) [7] and RT-PCR analysis of formalin-fixed, paraffin-embedded tissues (examining 21 prospectively selected genes) [8] have been used for chemotherapy treatment planning in breast cancer. In addition, novel genetic and molecular classifications of breast cancers have assisted in the individualization of adjuvant systemic endocrine chemotherapy, and have reduced the severity of side effects [1,9–11]. However, although these genomic
approaches provide valuable information for breast cancer prognosis, significant changes in gene expression may not be reflected in the level of protein expression or practical function [12]. Therefore, to obtain a more accurate and sophisticated understanding of patient status, the development of a novel analytical method to detect various biomarkers at the proteomic level is critical, in addition to analysis at the genomic level.

Immunohistochemistry (IHC) has been widely used for assessing therapeutic biomarkers and has become a major part of practical diagnosis for various malignancies in surgical pathology [13]. IHC allows the identification of proteins of interest and provides information on protein localization and tissue morphology [14]. In addition, many studies showing the relationship between immunohistochemical profiles and molecular classification of breast cancers support that IHC might play a significant role in subclassification of breast cancer patients [15,16]. Therefore, IHC-based assays can represent an ideal method to realize personalized-tailored therapies if efficient multiplexing method is created. However, conventional IHC has been faced with several practical limitations to examine tens of biomarkers in clinics: time, labor, diagnostic expense, and the amount of tissues required. For example, when various target proteins were examined by IHC for precise prediction and prognosis such as Oncotype Dx which examines over 20 genes involved in breast cancer [17], much time and labors are required. Although an automated IHC machine is able to overcome these issues, not only high costs from many biomarkers but much tissue consumption still remain [3], which might be significantly raised as practical problems for personalized medicine. Moreover, qualitative evaluation, subjective decision and variable result in IHC represent other hurdles toward a robust and reputable proteomic tool [13].

Recently, multicolor-based IHC studies have been reported with molecular dyes and quantum dots (QDs) for multiplexing [18–21]. Although the multicolor IHC including direct and indirect sequential staining methods has a unique advantage of co-expressions for biomarkers, several drawbacks are accompanied depending on the multicolor staining method [22,23]. They include low stability of primary antibodies from probe conjugation process, alteration of binding properties, difficulty of probe conjugation to antibodies, high cost of reagents, increases of time and labors, and cross-over nonspecific binding of secondary probes. Therefore, a parallel multiplexing approach is gaining the interest as an alternative to overcome the limitations of multicolor-based IHC and to enhance throughput of biomarker multiplexing. Unfortunately, to date, few studies for parallel multiplexing approach have been reported.

Here, we report a novel microfluidic parallel-multiplexed immunohistochemistry (MMIHC) platform for the quantitative pathological diagnosis of breast cancers. Since microfluidics enables the formation of a well-confined microenvironment [24,25], with fast and easy fluidic control [26] and the precise manipulation of fluids [27–30], not only the variation of immunohistochemical staining, the amount of time and labor required can be reduced via automation, but also multiple biomarkers can be analyzed on a limited cancer core area. In addition, because microfluidics uses much smaller volumes of reagents and antibodies, it allows cost-effective diagnosis and reduces financial burden of patients [31]. However, most microfluidic devices have been fabricated by using an irreversible bond between a microchannel and a glass slide, and only a few studies have introduced the interface between tissue slide and a microfluidic device; this also proves that few studies applied to human clinical specimens have been reported in microfluidics. In order to apply conventional thin-section tissues into on-chip without any additional modification process, a tissue slide-compatible assembler was developed for perfect compatibility of conventional IHC method, which was robust and optimal in a microfluidic device.

The goal of this study was to demonstrate significant improvement of IHC performances in reduction of labor and tissue consumption, quantification, consistency, sensitivity, specificity, cost-effectiveness, precise diagnosis and massive multiplexing. By comparing the biomarker scores from MMIHC platform with those of conventional whole-section analysis of breast cancer tissues, the usefulness of MMIHC platform to predict patient prognosis as well as to select drugs for chemotherapy was also evaluated.

Results

Operation of the MMIHC Platform

The design of the MMIHC device took into consideration 1) the number of solutions required for IHC, 2) the number of representative biomarkers in breast cancer, and 3) the appropriate reaction channel dimensions. We selected the biomarkers with the most frequently used and the most significant indicators in therapeutic decisions of breast cancers [21], so that estrogen receptor (ER), human epidermal growth factor receptor 2 (HER2), progesterone receptor (PR), and Ki-67 were chosen and thereby four straight reaction channels were designed. Accordingly, the device was composed of six reservoirs for reagents (R1–R6), four biomarker reservoirs (BR1–BR4), individual microvalves for reservoirs of reagents and biomarkers (RV1–RV6 and biomarker valve), four reaction channels, and one outlet (Figure 1A). Because biomarkers are investigated within a localized area among whole tissue in this approach, as in tissue microarray (TMA), the dimensions of the reaction channels were considered with the human breast tumor sample size in diameter and the partial area representing a tissue result. Recent results provided by the National Cancer Center of Republic of Korea have shown that >93% of biopsy breast tissues had tumor sizes >4 mm in diameter. Therefore, each reaction channel was 800 μm in width and 5 mm in length to apply to most of tissue sections, giving an area that was 14-fold larger than that of TMA with a 600-μm diameter.

The MMIHC device was fabricated via two-step multilayer soft lithography, poly(dimethylsiloxane) (PDMS; Sylgard 184; Dow Corning, MA) replica molding and aligning processes. Creating an appropriate interface between the MMIHC device and the tissue slide was one of the most important works in realizing chip-based MMIHC and minimizing tissue damage. A weight was set on top of the device to provide constant pressure and to create a reversible seal and a robust interface; in addition, this apparatus was quick and easy to assemble as shown in Figure 1B. In the assembly process of MMIHC assay, a tissue slide was loaded onto the bottom plate. The tissue was soaked with washing buffer and the plasma-treated MMIHC device having four reaction channels (attached under the upper plate) was put on the tissue and then aligned (Figure 1C). The MMIHC microchannels were filled with the buffer and any creation of microparticles was not allowed by the process. To avoid any leakage from microchannels, a weight was mounted on the upper plate, which the tissue was pressed by the walls of microchannels and fluids were perfectly flowed along the microchannels (Figure 1D).

Initially, deformation of reaction channels was doubted for pressure because of elastic characteristic of PDMS, which might cause different flow velocity profile for a reaction channel width. However, z-stacked images obtained via confocal laser scanning
microscopy (LSM 510, Carl Zeiss, Germany) showed that the reaction channels retained their original rectangular shape reasonably well, and were completely separated between each channel at 8 kPa (Figure S1A).

Tissue sample intactness was verified by white light scanning interferometry (Pemtron Co., Korea). Although the tissue area attached to the MMIHC device was damaged, tissue within the reaction channels was intact; this intact region was the staining area for scoring (Figure S1B). After characterization of the MMIHC device, fluidic control for various solutions and biomarkers was conducted. Even with a reversible seal, fluid flow was perfectly controlled; at a flow rate of 300 $\mu$L h$^{-1}$, solutions were flushed completely from the reaction channel after only 5 s. No leakage or bubble formation was observed even when the device was located on a cell block. De-waxing in xylene, rehydration, and heat-induced epitope retrieval (HIER) were conducted off-chip, whereas most of antibody–antigen interaction steps were performed on-chip. To enclose the stained tissue sample, dehydration and mounting processes were conducted after separating the MMIHC device from the slide.

**Multiplexed IHC on a Cell Block**

After aligning with a cell block (Figure 2A) and injecting various solutions and biomarkers into the MMIHC device, IHC was conducted on the chip. Four biomarkers, including ER, HER2, PR, and Ki-67, were examined simultaneously on a MCF-7 cell block. Each biomarker showed a different expression level, and the pattern of immunohistochemical staining was equivalent to the geometry of the reaction channels (Figure 2B). Because the reaction channels were completely separated for each biomarker, we were able to use the same label solution (containing 3,3′-diaminobenzidine tetrachloride [DAB]) for visualization of immunological reactions for all biomarkers. Thus, parallel multiplexing allowed rapid immunohistochemical staining and a direct comparison of staining between biomarkers at one site. In addition, this approach eliminated potential variation that may occur as a result of multiple IHC steps. After counterstaining on the chip, the slide was separated from the device without any damage to the sample, and then stored using a conventional slide storage procedure (Figure 2C).

Four breast cancer cell lines (AU-565, SK-BR-3, HCC70, and MCF-7) were examined for ER, HER2, PR, and Ki-67 expressions. All breast cancer cell lines showed staining for the indicated biomarkers at the appropriate cellular locations, providing comparable results to those obtained via conventional IHC (Figure 2D). In addition, biomarker staining was compared quantitatively for each cell line using image analysis. Microscopic images were analyzed based on the expectation-maximization (EM) algorithm and the Gaussian mixture model (GMM) to distinguish staining cells. Staining was presented as the expression
level, the value of which was determined by multiplying the ratio of the stained area and the average staining intensity. ER and PR were expressed only in the MCF-7 cell line and HER2 was expressed in AU-565 and SK-BR-3 cell lines. In contrast, Ki-67 was expressed in all cell lines (order of decreasing intensity: MCF-7, AU-565, SK-BR-3, and HCC70; Figure 3A). IHC is generally regarded as a qualitative method [13,32]; therefore, to validate the quantitative ability of the MMIHC platform, the results obtained above were compared to Western blotting results (Figure 3B). Western blotting showed that ER and PR were expressed only in the MCF-7 cell line, whereas HER2 was expressed in AU-565 and SK-BR-3 cell lines and both expression levels were similar as shown in MMIHC result. Ki-67 was expressed in all cell lines, similarly to the result that came from using the MMIHC platform; regression analysis showed that the correlation coefficient between Western blotting and MMIHC

Figure 2. MMIHC using cell blocks. (A) Image of an MCF-7 cell block aligned with the MMIHC device. Scale bar, 500 μm. (B) Image processing DAB reaction. The cell block was stained at discrete sites with ER, HER2, PR, and Ki-67 antibodies. Scale bar, 500 μm. (C) Completed cell block slide after detachment from the platform. Samples were not damaged by the detachment process. Inset shows a magnified view. Scale bars, 3 mm. (D) Comparison of immunohistochemical staining for ER, HER2, PR, and Ki-67 using conventional IHC versus the MMIHC platform (1000×). The staining quality of biomarkers using the MMIHC platform was comparable to that of the conventional method.

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Figure 3. Quantification of the MMIHC method. (A) Quantitative comparison of biomarker expression levels in AU-565, SK-BR-3, HCC70 and MCF-7 breast cancer cell lines; mean ± SD of four replicate assays per sample. (B) Western blotting of AU-565, SK-BR-3, HCC70 and MCF-7 cell lines for four biomarkers.

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MMIHC had better repeatability than IHC for both automatic and manual staining. To compensate systemic environment and condition for MMIHC, manual IHC and automatic machine, the expression levels were normalized by their mean value of each biomarker (data not shown). The Levene’s test, one of the tests for equal variance, was employed to compare the variances of normalized expression levels for above three methods (with significance level 0.05). The MMIHC had smaller variance than IHC for automatic staining (Levene’s test: \( p \)-value = 0.044). Although the IHC for automatic staining had smaller variances than for manual method, it was not statistically significant (Levene’s test: \( p \)-value = 0.395). This enhancement was likely the result of automated staining within the confined environment of a microfluidic system.

The MMIHC platform not only saved time and reagents, but also improved efficiency of antibody–antigen reaction. The semi-automated microfluidic platform completed IHC for the four biomarkers within 90 min, which was a 10-fold decrease in time required compared to conventional methods. In addition to time and volumetric effects, the optimal antibody concentrations were also approximately 10-fold lower. Although the time required compared to conventional methods. In addition to time and volumetric effects, the optimal antibody concentrations were also approximately 10-fold lower. Although equivalent samples were used, staining intensity remained similar to that obtained via conventional methods even when the biomarkers were diluted 10-fold. To better understand this phenomenon, we conducted computational fluid dynamics (CFD) studies on the kinetics of receptor–ligand binding. Under conventional IHC conditions, although the original concentration was maintained on the tissue in the initial state, the concentration of analyte exposed to the tissue was significantly lower because the analyte in the vicinity of tissue was bound to tissue receptors. In contrast, the concentration of analyte showed very little change at the tissue surface when the analyte was allowed to flow through the MMIHC platform. Because the reaction rate of receptor–ligand binding is decided by the absolute concentration of the analyte, and because mass transport to an area with very low fluid velocity (i.e., such as that near to the tissue sample) is determined by the concentration gradient, a high Reynolds number (Re) likely resulted in the continuous exposure of the initial analyte concentration and the formation of a steep concentration gradient (Figure S2).

We also showed that staining was more intense after the same incubation period when the flow rate was high (Figure 4A). Quantification clearly showed that higher flow rates produced higher expression levels for HER2 (Figure 4B). In addition, expression levels at flow rates of 60 and 180 \( \mu l \) h\(^{-1}\) when using a concentration of 0.1 \( \times \) HER2 were similar to those obtained using a conventional IHC method at a concentration of 1 \( \times \)HER2; this low concentration corresponded to the optimal antibody concentration determined by a pathologist. Based on these results and those of several other trials, we determined the optimal incubation conditions for primary antibodies (flow rate, 100 \( \mu l \) h\(^{-1}\); incubation period, 10 min), which translate into a >6-fold decrease in required time and 200-fold decrease in antibody consumption.

**MMIHC for Human Breast Cancer Tissues**

After verifying the utility of MMIHC on a chip and examining repeatability and the possibility of quantification using cell blocks, we applied the platform to patient tumor tissues which are heterogeneous in terms of morphology, genetic alterations and histopathological features. Over one hundred cases of human breast tumor tissues (113 cases) were randomly selected and the distributions of the investigated biomarkers were not biased. All experiments, including the MMIHC operation and clinical analysis, were blindly conducted. A pathologist judged the cases did not know not only the whole-section results of the cases but also any information of cases tested by the MMIHC platform.

Unlike cell blocks, tissue samples are not homogeneous; therefore, aligning the MMIHC device over the least heterogeneous area is critical for diagnostic outcome. The device was aligned at the area of highest cancer cell density, which was determined with the use of hematoxylin and eosin (H&E) staining slide (Figure 5A). After that, we examined the biomarker expression. Four biomarkers were examined simultaneously on the same tissue sample. In contrast to the cell blocks, which

![Figure 4](https://www.plosone.org/figure/10.1371/journal.pone.0010441.g004)

**Figure 4. Staining efficiency as a function of incubation conditions.** (A) Staining results for the SK-BR-3 cell line according to changes in flow rate and HER2 antibody concentration (incubation time, 30 min). Stronger expression was observed at higher flow rates. (B) Quantitative evaluation of HER2 expression under static and dynamic incubation conditions. As shown in the graph, the expression level of the 1 \( \times \) HER2 (conventional method) was not significantly different to that of 0.1 \( \times \) HER2 at a flow rate of 60 \( \mu l \) h\(^{-1}\); incubation period, 10 min), which translate into a >6-fold decrease in required time and 200-fold decrease in antibody consumption.

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showed homogeneous staining patterns, it was difficult to distinguish exact areas of biomarker staining because of tissue heterogeneity and some weak or negative staining results. This was resolved by injecting Mayer’s hematoxylin into the microchannels, which clearly demarcated the reaction channels (Figure 5B). Staining for the biomarkers was localized properly and score values for biomarkers were also equivalent to that obtained via conventional IHC (Figure 5C and 5D). Especially for PR, more non-specific staining was shown in conventional automatic IHC machine (Figure S3).

Although the biomarkers were examined in the areas in which cancer was most severe, a comparative study was essential to clarify whether the results from such a localized examination using the MMIHC platform could be considered representative of the whole tissue section. A blind experiment was performed and the 105 patient tissue slides investigated via MMIHC were scored by a pathologist (Table S1). The results revealed that Kendall’s coefficient of concordance (KCC, \( n = 105 \)) was 0.96 for ER, 0.90 for HER2, 0.95 for PR, and 0.98 for Ki-67; the agreement rates (\( \kappa \) coefficient, \( n = 105 \)) were 0.92, 0.65, 0.79 and 0.87 for ER, HER2, PR, and Ki-67, respectively (Table 1). HER2 showed the lowest KCC, although it should be noted that many cases for mismatches were slight (scores of 0 versus 1+). PR also showed a lower match rate than ER, which is consistent with previous studies showing that PR had lower sensitivity, specificity, and overall \( \kappa \) values than ER [34].

After confirmation of concordance between whole tissue section analysis and the MMIHC platform, we conducted the reproducibility study whether the platform also showed the same results within tissues originated from the same patient. Six cases were tested for reproducibility where four slides were made from the breast tumor of the same patient. To verify the concern of tissue

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\text{Table 1. Statistical concordance data for whole-section analysis versus the MMIHC platform (n = 105), including Kendall's coefficient of concordance (KCC), \( \chi^2 \) test, \( \kappa \) statistics, concordance and its 95% confidence interval (CI).}
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|                | Kendall’s coefficient of concordance (KCC) | \( \chi^2 \) | \( p \)-value | \( \kappa \)-value | \( p \)-value | % Concordance | 95% confidence interval (CI) |
|----------------|---------------------------------------------|-------------|--------------|-----------------|--------------|--------------|-------------------------------|
| ER             | 0.96                                        | 200         | <0.0001      | 0.92            | <0.0001      | 98.1         | 93.3 – 99.4                  |
| HER2           | 0.90                                        | 187         | <0.0001      | 0.65            | <0.0001      | 85.0         | 76.4 – 91.0                  |
| PR             | 0.95                                        | 198         | <0.0001      | 0.79            | <0.0001      | 90.5         | 83.2 – 95.3                  |
| Ki-67          | 0.98                                        | 204         | <0.0001      | 0.87            | <0.0001      | 91.4         | 84.4 – 96.0                  |

The KCCs were 0.96 for ER, 0.90 for HER2, 0.95 for PR, and 0.98 for Ki-67, and the agreement rates (\( \kappa \) coefficient) were 0.92, 0.65, 0.79, and 0.87 for ER, HER2, PR, and Ki-67, respectively. It is noted that KCC values >0.90 are regarded as almost perfect degree of agreement [45] and \( \kappa \) values >0.61 are considered as substantial agreement [34].

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Figure 5. Application of the MMIHC platform to human breast cancer tissues and comparison with conventional IHC. (A) The alignment of MMIHC device with an H&E-stained tissue sample. The device was aligned at a site showing numerous stained nuclei, indicating an area containing proliferating cancerous cells. Scale bar, 3 mm. (B) MMIHC image for a human breast cancer tissue sample. Compared to conventional IHC, the image shows the expression of the four biomarkers (ER, HER2, PR, and Ki-67) simultaneously on a tissue slide. Scale bar, 500 \( \mu \)m. (C) Magnified images of panel b (\( \times 400 \)). (D) Images of ER, HER2, PR, and Ki-67 staining via conventional IHC. The expression of biomarkers and the assigned Allred scores were equivalent.

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heterogeneity with reproducibility, for each case, a simple Latin square of order 4 was used by cyclic permutation tests in the first slides for subsequent slides to eliminate the sequential effect of tests; ER, HER2, PR and Ki-67 for 1st slide, HER2, ER, Ki-67 and PR for 2nd slide, PR, Ki-67, ER and HER2 for 3rd slide, and Ki-67, PR, HER2 and ER for 4th slide (Figure 6A). The biomarkers and their score levels were expressed with colors and their intensity (Figure 6B). The KCCs within appraiser showed that all biomarkers had over 0.95 values by repetition of measurement as all p-values are sufficiently small (Table S2). This result indicates that the platform is also reproducible in tissue sample and the sequence of the biomarkers does not affect score results significantly.

Not all tissue samples were suitable for the MMIHC platform. Although most tissues were firmly attached to the slides, tissue detachment occasionally occurred after HIER. In total, about 9% of the slides showed tissue detachment (10 cases among 115 samples). This problem is likely resolvable through optimization of the sample preparation process, including fixation, sectioning, and drying.

**MMIHC on a Needle Biopsy Sample of Human Breast Cancers for Precise Diagnosis in Early Stage**

Preoperative chemotherapy has been used for large primary and inflammatory breast cancers, and the examination of specific biomarkers in needle biopsy samples greatly facilitates the early individualization of neoadjuvant therapy [35]. Therefore, we applied the platform on a tissue from needle biopsy of human breast cancers, which reduced consumption of the rare tissues and enabled investigation for more various biomarkers even in small-sized samples. Despite the narrow area of core biopsy samples, the MMIHC device was aligned easily and we were able to examine the expression of four biomarkers on a single slide (Figure 7A). We noted that fatty tissue (solid arrow in Figure 7A) should be avoided when selecting an inspection window; biomarker expression in such regions was inconsistent compared to other non-fatty areas (dotted arrow in Figure 7A). Similar to cell blocks and tissue samples, the four biomarkers were also expressed in needle biopsy tissues (Figure 7B). A concordance study showed that the agreement rate (Cohen’s κ coefficient, n=8) was 1 for ER ($p=0.0001$), 0.71 for HER2 ($p=0.0175$), and 0.73 for PR ($p=0.0044$). In the case of tissues from needle biopsy, the κ statistics showed substantial agreement in concordance and ER showed 100% concordance and HER2 and PR showed 87.5% concordance (data not shown).

**Discussion**

The MMIHC platform, which was realized IHC on a chip and was applied to human clinical tissues specimens for the first time, minimized the use of externally connected equipment and formed a simple interface with the tissue sample. The unique platform significantly reduced the probability of assay failure (under 1%), which is of critical importance in practical use when dealing with clinically rare samples. Because the use of microfluidic channels creates a confined microenvironment and staining was semi-automated, MMIHC showed better repeatability of immunohistochemical staining compared to conventional manual IHC method and automatic IHC machine. This benefit seems to give improvement for inconsistent problem of IHC. Furthermore, the MMIHC yielded a 200-fold reduction of antibody consumption, fast immunological reaction, and the ability to examine various biomarkers for cancer chemotherapy in rare tissue samples. The characteristics of MMIHC platform are expected to reduce costs required for examination of various biomarkers when it is fully developed with a full automated and high-throughput manner.

Quantitative scoring is one of the main topics to overcome scoring subjectivity immunohistochemical analysis. By comparing our results with those obtained via Western blotting, we showed that the MMIHC platform is suitable for the semi-quantitative analysis of cell blocks. In addition, since MMIHC enabled the direct comparison of biomarker staining at a single site and eliminated the unexpected variation that may arise from multiple IHC steps, more accurate relative quantification was expected. Because the inspection window for each biomarker is relatively small, it would be also beneficial to perform quantification with image analysis as same rationale with TMAs [36]. To adopt automated quantitative image analysis into tissue results, however, staining between ductal carcinoma in situ (DCIS) and invasive

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**Figure 6. Reproducibility study within tissues originated from the same patient.** (A) Four slides were made from the human breast tumor of the same patient and cyclic permutation tests were conducted; ER, HER2, PR and Ki-67 for 1st slide, HER2, ER, Ki-67 and PR for 2nd slide, PR, Ki-67, ER and HER2 for 3rd slide, and Ki-67, PR, HER2 and ER for 4th slide. (B) Results of cyclic permutation tests for six cases. When the intensity of each color is equivalent in the boxes, the case was perfectly matched. The black dotted square for Patient 4 denotes different intensity compared to others. doi:10.1371/journal.pone.0010441.g006
cancers should be distinguished. Therefore, additional advances should be made in extensive image analyses and improved algorithms to decide clear scores.

Although this platform has many advantages over the conventional and automated machine-based IHC methods, it was doubted whether the MMIHC results are consistent with those obtained via conventional whole-tissue section analysis. Actually, this was also a critical concern regarding the use of TMA, which uses small tissue cores that may not be representative of the whole tissue section. However, after publication of the initial TMA results, many subsequent studies have shown excellent correlation for various tumor types [37−39]. Because the reaction channel of the MMIHC device was 14-fold wider than the 600-μm TMA core, it was expected to correlate well with the results of conventional methods, although TMA correlations are advantageous in their own right for permitting the selection of several tissue cores and thus enhancing representativeness. As we expected, in statistical aspect, results showed all of biomarkers showed over 0.90 KCC values, reflecting the whole section IHC scores with almost a perfect degree of agreement. This tendency was also shown in tissue samples from needle biopsies.

In clinical aspects, the MMIHC method is likely to be no detriment to patient care in clinical settings. Although there are many reasons for discrepancy of scores between whole-section analysis and MMIHC such as inborn errors with IHC itself (interobserver and interlaboratory variations), specimen selection, processing and representativeness of MMIHC results, total cases showing discordance in this study were 34 cases. Among our 34 discordant results, there were only two cases (case #27 and 93) of disagreement between control and MMIHC for ER which did not result in change of treatment plans (2+ = >3+). For PR results, three cases (case #7, 86, and 103) would result in a difference when making treatment decisions. However, only one case (case #103) could actually be treated differently since the ER status is always considered simultaneously. The other seven cases (case #18, 20, 30, 39, 59, 60, and 68) did not result in any change of treatment plans (1+ = >2+ or 3+ = >2+). Most of the discordance occurred in HER2 assessment. Specifically, two cases (case #28 and 69) among 16 discordant cases had potential risk for receiving trastuzumab. In case #69, since HER2 2+ by automated conventional method should be subsequently tested by fluorescence in situ hybridization (FISH) and treatment is dependent on the FISH result. Three cases (case #26, 100, and 101) would require additional FISH test. In summary, the cases that cause clinically different treatment are 0 case for ER, 1 case for PR and 1 case for HER2, showing only 1.9% variation in clinical treatment. Therefore, on the basis of our results, the MMIHC platform showed sufficient possibility of adoption as a method for the presentation of clinical specimens.

Another concern was the different scores depending on the sequence of biomarkers owing to tissue heterogeneity. Cyclic permutation test, however, revealed that scores of the biomarkers were reproducibly repeated and the sequence of the biomarkers did not affect score results significantly. This result is also likely to imply that the aligning position is not such extremely critical; meaning that the slight different position at the area of highest cancer cell density was affordable in reproducibility. Although this study was conducted in a microfluidic device with 800-μm-wide reaction channels to satisfy so much as tissues having small cancer core, increasing the width and number of the channels in different directions is expected to enhance the representativeness of the results, in the same way that the concordance of results obtained via TMA was improved by increasing the number and size of the cores [39−41].

In improving the standard of patient care, many issues must still be resolved before the current format of the MMIHC platform can compete with existing methods. Although the MMIHC approach contains a sampling process within a whole section of tissue, it is obvious that an assay that could accurately quantify several cancer-related proteins simultaneously on single tumor section or small tumor specimens does offer clear advantages and technical advances over standard immunohistochemical method [21]. Therefore, a novel method showing substantial agreement for KCC statistics is expected to be useful as a decision supporting tool for pathologist and clinician.

In IHC, multiplex staining is growing need within limited quantity of clinical samples. Many studies for multiplex IHC have been based on the multicolor approach [18−21]. Sequential indirect multiplexing method, which conducts blocking, antibody reaction and then tagging with fluorescence in repeat, has an advantage to look at a wide range of co-expressions for biomarkers. However, it is a labor-intensive method that consumes a lot of time and reagents.
Materials and Methods

Ethics Statement

Human tissue samples from each tumor lesion were obtained from the National Cancer Center Hospital (Goyang, Korea) and the Korea University Anam Hospital (Seoul, Korea), with the corresponding written consents given by the patients or their relatives. This study was approved by the Institutional Review Board (IRB) at the National Cancer Center Hospital and Korea University Anam Hospital.

Fabrication of a MMIHC Device

The fluidic channel mold for a MMIHC device was fabricated via two-step multilayer soft lithography. To construct rectangular reaction channels, SU-8 2025 (Microchem Corp., MA) was spin-coated to form a 50-μm thick layer on a bare silicon wafer, patterned by UV exposure. After developing the wafer, a masking layer was patterned on the reaction channel area. To make a round-shaped remnant fluidic channel, AZ 9260 was spin-coated to form a 25-μm thick layer. After lifting off the masking layer, it was exposed to UV light, and developed using AZ photore sist developer. The fabricated mold was refilled by heating, and the fluidic channels were transformed into a round shape, except for the reaction channels. The control channel mold was fabricated by conventional SU-8 photolithography. After spin-coating the fluidic channel mold with PDMS and curing, the fluidic channel was aligned and bonded with the control channel using an O2 plasma ash (270 W for 30 s).

Preparation of Cell Blocks and Tissues

Four commercially available breast carcinoma cell lines, MCF-7, SK-BR-3, AU-565, and HCC70, were obtained from the American Type Culture Collection (ATCC; Manassas, VA). HCC70, MCF-7, and AU-565 were maintained in RPMI-1640 and SK-BR-3 cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 100 IU ml⁻¹ penicillin, and 100 mg ml⁻¹ streptomycin. All cell lines were cultured at 37°C and incubated in 5% CO2. Adherent cells were harvested by trypsinization before reaching confluence. For IHC analysis, the harvested cells were centrifuged, fixed in formalin, suspended in agar, and embedded in paraffin to produce a cell block. Paraffin-embedded cell blocks were sectioned at 4 μm thickness using a microtome (Leica, Germany). The sections were baked onto positively charged slides and allowed to dry for 1 h at room temperature, followed by 1 h in an incubator at 60°C.

Tissue samples from each tumor lesion were fixed for 24 h in 4% neutral-buffered formalin, Bouin’s fixative, acetic formalin alcohol (AFA), or 4% or 10% unbuffered formalin; 4 h in PreFer (Anatech, Battle Creek, MI) or Pen-Fix (Richard Allen Scientific; Kalamazoo, MI); or 48 h in 4% neutral-buffered formalin. After paraffin embedding, tumor specimens were cut into 4-μm-thick sections and allowed to dry for 1 h at room temperature, followed by 1 h in an incubator at 60°C.

Immunohistochemical Staining

Four biological markers were investigated. ER (SP1) antibody (Ventana, Tucson, AZ) and PR (1E2) antibody (Ventana) were used in conventional methods at 1× concentration and in MMIHC at a dilution of 1:10. HER2 oncoprotein antibody (Dako, Denmark) was used in conventional methods at a dilution of 1:500 and in MMIHC at a dilution of 1:5000. Ki-67 (clone MIB-1) antibody (Dako) was used in conventional methods at a dilution of 1:50 and in MMIHC at a dilution of 1:500. Cell blocks and tissues were de-waxed in xylene and rehydrated through a graded series of ethanol solutions. A microwave antigen-retrieval technique was used and the samples were treated in target retrieval solution, pH 9 (Dako), for 20 min at 750 W. The Cap-plus kit (Zymed, San Francisco, CA) was used for immunostaining and
Mayer's hematoxylin (Sigma, St Louis, MO and Labvision, Fremont, CA) was used for counterstaining.

**Western Blotting**

Protein was extracted from cells by the addition of lysis buffer followed by centrifugation at 16,000 g for 10 min at 4°C. The supernatant fractions were separated by polyacrylamide gradient gel (8–16%) containing sodium dodecyl sulfate. Following electrophoresis, the proteins were transferred to polyvinylidene fluoride membranes (Millipore, Bedford). The membranes were blocked in TBS-T (0.1% Tween 20) containing 5% non-fat milk (Bio-Rad, Richmond) for 1 h at room temperature. After blocking, the membranes were incubated for 2 h at room temperature with primary antibodies against ER, HER2, PR, and Ki-67 (clone MIB-1) antibodies. Then, the membranes were washed in TBS-T (0.1% Tween 20), for 15 min at a time, and incubated with diluted HRP-conjugated secondary antibody (Southern Biotech, Birmingham, AL) for 1 h at room temperature. This was followed by washing with TBS-T (3×15 min), incubation with WEST-ZOL® plus chemiluminescence reagent (ßNiRON Biotechnology, Korea) for 1 min, and exposure to film (Kodak Blue XB-1, Rochester, NY). The immunoblot of ß-actin (R&D Systems, Korea) was used as a loading control.

**Image Acquisition and Analysis for Quantification of MMIHC**

Tissue images were taken in optimal condition considering shading and glare. After completing MMIHC, the sample was placed on an inverted microscope (Carl Zeiss, Germany) and images were acquired with a microscope CCD camera (Olympus DP71, Japan) under 15,000 Lux light intensity. The microscope CCD camera has 12.5 megapixels, 12-bit digital color that displays the native CCD's full-resolution live image at 15 frames per second. The microscopic images were divided into three parts (Figure S4A): the staining part (SP) of the cell, the non-staining part (NSP) of the cell, and the background (Figure S4B). Then the expression level was defined by multiplying the staining ratio and intensity, where the staining ratio is the area ratio of the SP to the cell part (SP and NSP) and the staining intensity is the average intensity in the SP. At least five images were randomly taken along the individual reaction channel via 400× magnification and the average expression level for the images was presented as a representative value of immunohistochemical staining for a biomarker.

Bayesian classification was employed to segment a microscopic image into the three parts based on their colors. The color distribution of each part in RGB color space was represented by the Gaussian mixture model (GMM), and estimated by the expectation-maximization (EM) method. The probability density function (PDF) of GMM with C components is defined as a convex combination of Gaussian PDFs

\[
p(x; \theta) = \sum_{j=1}^{C} p_j N(x; \mu_j, \Sigma_j),
\]

where \(N(x; \mu, \Sigma)\) is the d-dimensional Gaussian PDF with mean \(\mu\) and the covariance \(\Sigma\), and \(p_j\) is the portion of the \(j\)-th component such that \(0 < p_j < 1\) for all components, and \(\sum_{j=1}^{C} p_j = 1\). The parameter list

\[
\theta = (p_1, \mu_1, \ldots, p_C, \mu_C, \Sigma)
\]
defines a particular PDF of the GMM. The parameters for the SP, NSP, and background were obtained via the EM method using data collected from IHC sample images. The numbers of clusters in the GMM for the SP, NSP, and background were set at 3, 10, and 2, respectively. Thirty independent IHC images were used to train the PDF. The optimized Gaussian PDF was automatically applied to the experimental images.

**Data and Statistical Analyses**

We compared the quantification of ER, HER2, PR, and Ki-67 expressions between MMIHC and the respective quantified Western blot bands by using Pearson’s correlation. For ER and PR, the Allred scores (0–8) assigned by a pathologist were translated into negative (−) for 0 score, weak (+) for 2 and 3 scores, intermediate (+++) for 4, 5, and 6 scores, and strong (+++) for 7 and 8 scores. For HER2, we followed the HER-2/new FDA-approved scoring system and translated into (−) for 0, (+) for 1+, (+++) for 2+, and (+++) for 3+. For Ki-67, stained cell number was counted and translated into (−) for ≤5%; (+) for 5% < x ≤20%; (++) for 20% < x ≤40%; (+++) for > 40%. Attribute agreement analysis for tissue data was performed using Minitab version 15. The relative agreement between conventional IHC and MMIHC data for ER, HER2, PR, and Ki-67 was assessed using KCC. Fleiss’ and Cohen’s κ statistics were calculated to evaluate the agreement between the two systems, for which the contingency tables between conventional IHC and MMIHC for ER, HER2, PR, and Ki-67 are presented. Hypothesis testing was conducted using the two sample t test to analyze HER2 and Ki-67 expression in the four breast cancer cell lines.

**Supporting Information**

Table S1 Comparison of scores obtained via whole-section analysis (control) versus the MMIHC platform in human breast cancer tissues (n = 105). For ER and PR: negative (−); weak (+); intermediate (++); strong (+++). For HER2: score 0 (0); score 1 (+); score 2 (++); score 3 (+++). For Ki-67: ≤5% (−); 5% < x ≤20% (+); 20% < x ≤40% (++); > 40% (+++). The unit of tumor size is centimeter. DCIS, ductal carcinoma in situ.

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Table S2 Statistical concordance table of cyclic permutation test, including KCC, \(\chi^2\) test and \(p\)-value. Six cases were tested for reproducibility where four slides were made from the breast tumor of the same patient (total number of tissues: 24). The KCCs were 1.00 for ER, 1.00 for HER2, 1.00 for PR, and 0.96 for Ki-67, respectively. The cyclic permutation tests indicate that the MMIIHC platform is reproducible in tissue sample and the reproducibility where four slides were made from the breast tumor of the same patient (total number of tissues: 24). The KCCs were 1.00 for ER, 1.00 for HER2, 1.00 for PR, and 0.96 for Ki-67, respectively. The cyclic permutation tests indicate that the MMIIHC platform is reproducible in tissue sample and the sequence of the biomarkers does not affect score results significantly.

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**Figure S1** Characterization of the MMIIHC platform. (A) Plane and z-stacked confocal laser microscopy images of the reaction channel area under 8 kPa. Reaction channels retained their original rectangular shape and each was separated completely. (B) Surface image of a cell block visualized using white light scanning interferometry. Cells in the reaction channels were intact under pressure, except for those in areas in direct contact with the MMIIHC device.

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**Figure S2** Computational fluid dynamics (CFD) study examining the kinetics of receptor-ligand binding. (A) Concentration
profile of the analyte using the conventional method (transient state after 80 s). The red dotted line indicates a tissue sample with antigens. The analyte concentration in the vicinity of the tissue decreased with time because the tissue functioned as a sink. (B) Concentration profile of the analyte using the MMIHC platform (RS = 4.5; transient state after 80 s). Fresh analyte flowed into and was maintained in the vicinity of the tissue; therefore, the concentration of the analyte showed little decrease at the tissue surface as time progressed. (C) Concentration distribution of the analyte according to incubation conditions. The concentration profiles between a non-flowing microchannel and the conventional method were similar, and the concentration of the analyte exposed to the tissue was higher when the flow velocity of the analyte increased. (D) Concentration gradient versus analyte incubation conditions. When the flow velocity was high, diffusion of the analyte was dominant.

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**Figure S3** Images of immunohistochemical staining for PR. (A) An image of PR staining from automatic machine (×400). (B) An image of PR staining from the MMIHC platform (×400). Blue solid arrows indicate non-specific staining. Normally, more non-specific staining was shown in conventional automatic IHC machine.

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**Author Contributions**

Conceived and designed the experiments: MSK ESL JKP. Performed the experiments: MSK SK CHK. Analyzed the data: MSK TK CHK ESL JKP. Contributed reagents/materials/analysis tools: TK SYK CYB JC. Wrote the paper: MSK JKP. Final approval of manuscript: JKP MSK ESL.

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**Figure S4** Image analysis of biomarker expression level. (A) A microscopic image acquired via MMIHC. (B) The image was divided into three parts: the staining part (SP), the non-staining part (NSP), and the background. Only the cell area (SP and NSP) was considered to minimize the variation of expression level according to cell density. (C) Image after analysis. Only the brown-colored areas remained.

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