Improving Algorithm for the Alignment of Consecutive, Whole-Slide, Immunohistochemical Section Images

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Submitted: 03-Dec-2020 Revised: 29-Apr-2021 Accepted: 01-Jun-2021 Published: 03-Aug-2021

Abstract

Background: Accurate and precise alignment of histopathology tissue sections is a key step for the interpretation of the proteome topology and cell level three-dimensional (3D) reconstruction of diseased tissues. However, the realization of an automated and robust method for aligning nonglobally stained immunohistochemical (IHC) sections is still challenging. In this study, we aim to assess the feasibility of multidimensional graph-based image registration on aligning serial-section and whole-slide IHC section images. Materials and Methods: An automated, patch graph-based registration method was established and applied to align serial, whole-slide IHC sections at ×10 magnification (average 32,947 × 27,054 pixels). The alignment began with the initial alignment of high-resolution reference and translated images (object segmentation and rigid registration) and nonlinear registration of low-resolution reference and translated images, followed by the multidimensional graph-based image registration of the segmented patches, and finally, the fusion of deformed patches for inspection. The performance of the proposed method was formulated and evaluated by the Hausdorff distance between continuous image slices. Results: Sets of average 315 patches from five serial whole slide, IHC section images were tested using 21 different IHC antibodies across five different tissue types (skin, breast, stomach, prostate, and soft tissue). The proposed method was successfully automated to align most of the images. The average Hausdorff distance was 48.93 μm with a standard deviation of 14.94 μm, showing a significant improvement from the previously published patch-based nonlinear image registration method (average Hausdorff distance of 93.89 μm with 50.85 μm standard deviation). Conclusions: Our method was effective in aligning whole-slide tissue sections at the cell-level resolution. Further advancements in the screening of the proteome topology and 3D tissue reconstruction could be expected.

Keywords: Cell-level resolution, Image registration, Immunohistochemistry, Multidimensional graph-based registration, Proteome topology, Three-dimensional tissue reconstruction, Whole-slide images

Introduction

Automatic and precise alignment of consecutive whole-slide histopathology section images is a key process leading to the future cell-level three-dimensional (3D) tissue reconstruction and topographical proteomics. However, only few studies focused on this topic.1-4 The bottlenecks were attributed to the following four factors: (1) a ×10 magnified, whole-slide histopathology image was at average 32,947 × 27,054 pixels in size, and at its maximal resolution may reach 100,000 × 100,000 pixels. Dealing with data of this size was extremely challenging using the common computing hardware;5 (2) currently, most histopathology slides are manually made, thereby introducing various distortions to the tissue section, such as cuts, folds, tearing, and local tissue stretch or compression;1-4 (3) each whole-slide image was unique and slightly different than each other in spatial coordinates, and they were cut in a continuous manner with

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How to cite this article: Liang CW, Chang RF, Fang PW, Chen CM. Improving algorithm for the alignment of consecutive, whole-slide, immunohistochemical section Images. J Pathol Inform 2021;12:29. Available FREE in open access from: http://www.jspathinformatics.org/text.asp?2021/12/1/29/323081
4-μm spacing, thereby making the assessment of alignment adequacy difficult, as no ground-truth image was available in each level of the whole-slide image series; and (4) for nonglobal stained slides, such as immunohistochemical (IHC) slides, the images produced from different antibodies were usually viewed as independent objects by the current established registration methods and were, thus, difficult to be aligned, particularly in sections wherein the stained proteins are highly expressed in contrast to the low-counterstained background. Several image-registration models and algorithms were proposed in few previously conducted studies. They mainly addressed the first three above-mentioned issues and significantly improved the accuracy and precision of aligning serial neighborhood, globally-stained, histopathological section images.\cite{1-4} However, in daily laboratory usage, efficacy issues were still common, and the alignment and concatenation of multiple, differently and strongly stained IHC whole-slide images usually failed. In this study, we improve the current method to align consecutive, whole-slide gigapixel histopathology images, thereby enhancing the program efficiency and addressing the issue regarding the alignment of multiple, nonglobally and differently stained IHC section images. The Hausdorff distance was used to measure the effectiveness of image alignment.\cite{1} We prove that our method can successfully achieve image alignment at a near cell-level resolution, thereby not only leading to the future cell-level 3D tissue reconstruction but also displaying the ability to incorporate the topographical protein information into the reconstruction.

**Materials and Methods**

Whole-slide, 4 μm thick, serial IHC histopathology section images were prepared from formalin-fixed, paraffin-embedded tissues of the skin, breast, stomach, prostate, and soft tissue using 21 different IHC antibodies [Figure 1 for selected images of p63, CK5/6, EMA, CK7 and HER2, Supplemental Table 1 for staining conditions, clones and methods for all used antibodies, and Supplemental Figure 1 for set design], with an average image dimensions of 32,947 × 27,054 pixels. IHC staining was performed in a Leica-Bond Max automated immunostainer (Leica, Taipei, Taiwan), and whole-slide images (MIRAX Virtual Slide File, .mrxs, binary, Carl Zeiss MicroImaging) were generated using a 3DHISTECH Pannoramic 250 Flash III scanner (3DHISTECH Ltd., Budapest, Hungary) at ×400 resolution using a ×20 objective, with a 24-bit red, green, and blue (RGB) format and 0.19 μm/pixel resolution. The images were then viewed by CaseViewer (ver. 2.3 or above, https://www.3dhistech.com/research/software-downloads/) and exported as. TIF files with a 1:4 conversion (equivalent to a ×100 view or ×10 objective in practice, a resolution of 0.76 μm/pixel) for further computation. This study was approved by the Institutional Review Board of the National Taiwan University Hospital (201412155RIND) and Fu Jen Catholic University (C108102).

The general flowchart of the proposed method is shown in Figure 2. The input images contained a reference (Sequence 0) image and translated (Sequences 1–n with phase correlation, scale/rotation recovery on RGB transformation) high-resolution images (HR\(_n\) and HT\(_n\)) and 1× low-resolution images of 3295 × 2705 pixels (LR\(_n\) and LT\(_n\)) rescaled from HR\(_n\) and HT\(_n\).

The alignment began with the initial alignment (preprocessing) of HR\(_n\) and HT\(_n\) (object segmentation and rigid registration) and nonlinear registration of LR\(_n\) and LT\(_n\), followed by the multidimensional graph-based image registration of the segmented patches, and finally, the fusion of deformed patches for inspection. To assess the alignment accuracy, a graph-based method was adapted to check the distance shift from the local highest and lowest probability points converged from the neighborhood regions (Hausdorff distance).\cite{1}

**Initial alignment**

The initial alignment was used to reduce the object deviations between serial whole-slide IHC images. The staining intensities from different antibodies between each image were normalized by the Gamma correction.\cite{7} Foreground object preidentification was performed to detect the extent of staining in the tissue according to the reaction intensity generated by the Otsu thresholding algorithm,\cite{8} while the noise were removed. To correctly identify the surrounding nonstained region, a segmentation template was generated using an image matting and segmentation algorithm\cite{9,10} and applied to segment the tissue based on the result of the foreground object preidentification. The segmentation template differentiated an image into three regions, namely foreground (object), background, and unknown regions. The ring region collected from the outer 50 pixels from the Otsu foreground region comprised the unknown region, and it was then verified by an image matting algorithm defined as

\[
L(x-y) = \alpha(x-y) \times F(x-y) + (1-\alpha(x-y)) \times B(x-y),
\]

where

Figure 1: Selected serial immunohistochemical histopathological section images scanned at ×10 resolution: (a) p63, (b) CK5/6, (c) EMA, (d) CK7, and (e) HER2
\( L_{(x,y)} \) denotes the input image pixel value and \( x \) and \( y \) the image coordinates. The terms \( F_{(x,y)} \) and \( B_{(x,y)} \) denote the pixel values in the foreground and background, respectively, and \( \alpha_{(x,y)} \) denotes the average opacity value of the foreground pixels.

To minimize the inherent global angle deviations due to manual tissue sectioning, the longest horizontal and vertical diameters of each image were first determined; the intersection point was considered as the object center, which was then used to calculate the displacement between two objects. A rigid similarity transformation,\(^{[10]}\) appropriate for whole-slide image registration owing to its low computation complexity, was applied to rescale the inconsistent objects and calibrate the angle differences between each \( HR_{0} \) and \( HT_{n} \). The component matrix, which included rotation, translation, and scaling, is described as follows:

\[
\begin{bmatrix}
X' \\
Y'
\end{bmatrix} = S \begin{bmatrix}
\cos \theta & -\sin \theta \\
\sin \theta & \cos \theta
\end{bmatrix} \begin{bmatrix}
X \\
Y
\end{bmatrix} + \begin{bmatrix}
dx \\
dy
\end{bmatrix} \quad (1)
\]

Where \( dx \) and \( dy \) denote translation parameters, \( S \) a rescale parameter with rotation matrix \( \theta \), \( X \) and \( Y \) the original coordinates of \( HR_{0} \) and \( X' \) and \( Y' \) the transformed coordinates of \( HT_{n} \).

### Nonlinear registration with low resolution images

The global calibration corrected most angle deviations due to manual sectioning, and it was followed by nonlinear image registration to align the inconsistent areas between major tissue elements.\(^{[11]}\) A patch-wise image registration model\(^{[3]}\) was adapted to efficiently discretize the deformation and evaluate the similarity between \( LR_{0} \) and \( LT_{n} \). To compute the deformation matrix \( Dy \) from each aligned \( LR_{0}/LT_{n} \), an objective similarity function with respect to the discretized deformation \( J(\mathbf{LT}_{n},\mathbf{LR}_{0},\mathbf{Dy}) \) was defined as the computation of

\[
\min_{\mathbf{D}} J(\mathbf{LT}_{n},\mathbf{LR}_{0},\mathbf{Dy}) = D(\mathbf{LT}_{n},\mathbf{LR}_{0},\mathbf{Py}) + S[\mathbf{u}] \quad (2)
\]

where the \( D \) function measured the image similarity, and the \( S \) function measured the reasonability of the transformation.

To assess the similarity, which described the correspondence between neighbor images, the normalized gradient field\(^{[12]}\) formula was applied, taking advantage of its intensity-gradient-driven algorithm that could accurately address the registration of differently stained IHC images. The \( D \) function that measured the image similarity was defined as follows:

\[
D(\mathbf{LT}_{n},\mathbf{LR}_{0},\mathbf{Dy}) = NGF[\mathbf{LT}_{n},\mathbf{LR}_{0},\mathbf{Dy}] = \int_{\Omega} \left( \frac{\nabla LT_{n}(Py(\hat{x}))}{\nabla LT_{n}(Py(\hat{x}))} \cdot \frac{\nabla LR_{0}(\hat{x})}{\nabla LR_{0}(\hat{x})} \right)^{2} dx \quad (2)
\]

Where \( D \) denotes the prolongation operator that interpolates

### Table 1: Average baseline differences in serial sections of the same immunohistochemical staining with 1 section apart, 2 sections apart, and 3 sections apart by the proposed method

| Distances between sections (Number) segmentation objects | Average (µm) (range) | SD (µm) | \( P \) (t-test) |
|----------------------------------------------------------|----------------------|---------|-----------------|
| 1 section apart, 4 µm in distance (30)                   | 44.64 (41.31-50.01)  | 7.49    | -               |
| 2 sections apart, 8 µm in distance (30)                  | 42.56 (36.98-48.14)  | 13.20   | 0.46            |
| 3 sections apart, 12 µm in distance (30)                 | 44.88 (35.95-53.81)  | 6.15    | 0.89            |

SD: Standard deviation

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**Figure 2:** Flowchart of the proposed multidimensional, graph-based image-registration method. The alignment began with the initial alignment of high-resolution reference and translated images (object segmentation and rigid registration) and nonlinear registration of low-resolution reference and translated images, followed by the multidimensional graph-based image registration of the segmented patches, and finally, the fusion of deformed patches for inspection.
the low-resolution deformation onto the image grid, and
\[
\|x\|^2 - \|x\|^2 + \mathcal{E}^2 \] is to assess image similarity. The square brackets in the expression \(y(x)\) denote the bilinear interpolation of \(y\) based on the four neighboring pixels on the grid of \(y\).

A diffusive regularizer\(^{[\text{13}]}\) was included to physically define the deformation model during image registration to ensure the robustness and computation speed. To avoid the local minimal value and optimize the joint objective function, the L-BFGS\(^{[\text{14}]}\) was applied as an optimizer using a discretized analytic derivative and was initialized with the analytic Hessian of the regularizer, which can be interpreted as a special case of linear elasticity as follows:

\[
S[u] = \frac{\alpha}{2} \sum_{\mathbf{u} \in \Omega} \langle \nabla \hat{u}, \nabla \hat{u} \rangle
\]  

(3)

**Multidimensional graph-based registration**

To address the minor local deformations of the detailed internal tissue structures generated between differently stained images, a multidimensional graph-based registration was proposed. The algorithm featured two stages: Registration by feature point aggregation and graph-based registration. The high-resolution slides \((H)\) were partitioned into patches first, each of size 4915 \(\times\) 4915 pixels, with overlapping areas up to 10\% of the patch size. For each patch, an individualized transformation was independently performed using a deformation matrix \(D_y\). High-dimensional vectors generated from NGF deformation were obtained, and a cubic B-spline transform was estimated using regularized least-squares error minimization. Finally, the step was refined by resolution progression from low \((LT_n)\) to high \((HT_n)\).

**Registration by feature point aggregation**

The pixels in the RGB color space were first translated into a gradient vector image, \(\mathbf{VG}_\alpha\), which was defined as

\[
\mathbf{VG}_\alpha(x, y) = (G_x^2 + G_y^2), i \in R, T, \]  

where \(G_x^2\) and \(G_y^2\) denote the gradient translations of the horizontal and vertical axes, respectively. The properties of the gradient magnitude contributed to the preservation of differently stained tissue structures, and they clearly demonstrated the differences and similarities in pixel intensity from different gradient vector directions, thereby facilitating the extraction of correct matching feature points. K-means clustering\(^{[\text{15}]}\) was applied to each translated patch for accurate feature point extraction in discretized-intensity categories.

Matching feature points were extracted using speeded up robust features,\(^{[\text{16}]}\) which contained steps including matching, feature point detection, and local neighborhood description. A Hessian matrix-based blob detector was used for detecting the feature points. The determinant of the Hessian matrix was applied as a measure of the local variety around the points, while the points were selected when the determinant was maximal. The detector of the feature points contained an indicator to assign invariability for every interest point for a local neighborhood description. These descriptors were adapted to search for correlations between the original image and transformed image when deformations occurred. By comparing the descriptors obtained from different images, matching pairs could be found.

Moreover, the process was refined by boundary control to compute the relative distance and orientation of the vector pointing from the centroid to boundary control points. The boundary control points were set along the object boundary, which was the result of foreground object segmentation, in a clockwise sequence 76 pixels apart. In each alignment after the feature point extraction, the distances of the matching feature points, >100 pixels and <15 pixels, were removed to enhance the significant feature points for the construction of the deformation graph model.

**Deformation graph model construction**

Nonlinear registration on the patched sections was performed using the deformation graph model construction to reduce the local deformation. The algorithm contained three stages: Deformation graph model construction, graph-based registration, and fusion of deformed patches. For the construction of the deformation graph model, Delaunay triangulation\(^{[\text{17}]}\) was used to define the deformation spatial relations by feature points. The entire patched image was covered with triangles, and any vertex of each triangle did not fall in the circumscribed circle of other triangles, thereby generating uniqueness. In each align, when the feature points were altered, only a part of the triangular structures needed to be changed, thereby enhancing the computation performance. Figure 3 shows the graphs of deformation spatial relations from a part of differently stained images.

After model construction, the one-to-one corresponding relationship between the pairs of triangle in two aligned images was defined for achieving smooth graph-based registration. Figure 4 shows an example of alignment; the triangle of moving 1s in the transformation patch was always mapped to the triangle of fixed 1s in the reference patch. All the pairs of the corresponding triangles were transformed using an affine transformation matrix\(^{[\text{18}]}\) for matching.

Figure 3: Example graphs of deformation spatial relations with (a) p63 and (b) EMA. Delaunay triangulation was used to define the deformation spatial relations by feature points. The entire patched image was covered with triangles, and any vertex of each triangle did not fall in the circumscribed circle of other triangles, thereby generating uniqueness.
The transformation matrix $AT$ was defined as follows:

$$AT = M_{\text{translation}} \times M_{\text{scaling}} \times M_{\text{rotation}} \times M_{\text{shear}},$$

which contains translation, scaling, rotation, and shear mapping, where

$$M_{\text{translation}} = \begin{bmatrix} 1 & 0 & x1 \\ 0 & 1 & x2 \\ 0 & 0 & 1 \end{bmatrix},$$

$$M_{\text{scaling}} = \begin{bmatrix} s1 & 0 & 0 \\ 0 & s2 & 0 \\ 0 & 0 & 1 \end{bmatrix},$$

$$M_{\text{rotation}} = \begin{bmatrix} \cos \theta & -\sin \theta & 0 \\ \sin \theta & \cos \theta & 0 \\ 0 & 0 & 1 \end{bmatrix},$$

and $M_{\text{shear}}$ was.

Finally, to obtain a global smooth deformation, bilinear interpolation was used to fuse the deformed patches into a complete graph. The global smoothness of the fused deformation was ensured by comparing the differences between the two deformation vectors in the overlapping areas.

All the experiments were conducted using MATLAB 2019b together with the C++ open source library. The hardware environments were Intel Core i7-8700K 4.60 GHz processor, 32 G of random access memory at 2400 MHz, and Microsoft Windows 10 operating system. For validating the effectiveness of the proposed method, the average (Avg) and standard deviations (SD) of the Hausdorff distance were calculated. The performance accuracy of the registration was evaluated by calculating the differences between the manual segmentations of the corresponding structures after registration. To assess the accuracy and precision of each alignment, thirty structure objects were manually segmented per slide pair by a pathology expert without the knowledge of the registration result to calculate the Hausdorff distance in a stack of every 5 different IHC stainings (HR$_a$ and HT$_{1-4}$, total 120 segmentation objects). To assess the inherent error (baseline difference) in serial sectioning, a 4-slide stack of the same CK5/6 staining was also created. The average Hausdorff distances were calculated between slides that are 1 section apart (4 μm in distance), 2 sections apart (8 μm in distance), and 3 sections apart (12 μm in distance).

**RESULTS**

Figure 5 shows an example of aligning paired patch images at different stages: Original offset, initial alignment using a published protocol (patch-based nonlinear image registration), and using our proposed method. The aligning precision and accuracy were gradually improved. In the original offset and initial alignment, macrostructures were aligned, although microstructures (e.g., a hair follicle, as marked in the figure) were not aligned. Using the proposed method, most microstructures were aligned. The results demonstrated that smooth microstructural correspondences were established.

Figure 6 shows the examples of feature point extraction in different IHC-stained patch images obtained after performing the proposed registration feature point aggregation at the multidimensional graph-based registration stage. The results demonstrate that the coordinates of most of the important microstructures (determined by a pathologist) were featured during the registration with correspondence.

To demonstrate the effectiveness of the proposed method in the practical screening of the proteome topology, a pseudocolor system was applied for visualization. Each color represented a different protein highlighted by the corresponding IHC antibodies. The proposed method showed the best alignment through which the protein–protein coexpression or mutually exclusive expression could clearly be elucidated.

The computation time was in average 300 s per patch of the proposed method, as compared with 320 s per patch for the published patch-based nonlinear registration method. It took about 5 h to align two whole-slide images with 60 patches and about 8.3 h to align two images with 100 patches.

The average baseline differences (inherent error caused by serial sectioning) are listed in Table 1. The results estimated the minimum Hausdorff distance that can be achieved. The different distances between the sections did not generate a
statistically significant ($P > 0.05$) deviation from the $HR_0$ slide in Hausdorff distance calculation.

Overall the proposed multidimensional graph-based registration method improved the aligning efficiency and precision [Table 2]. The average Hausdorff distances across a stack of five different IHC stainings ($HR_0$ and $HT_{1-4}$, total 120 segmentation objects) were 361.79 μm (range: 139.28–688.702 μm), 93.89 μm (range: 59.59–153.85 μm), and 48.93 μm (range: 43.75–54.77 μm) for initial alignment, published patch-based nonlinear registration method, and proposed multidimensional graph-based registration method, respectively. The average standard deviations were 62.82 μm, 50.85 μm, and 14.94 μm, respectively. The best alignment occurred in skin tissue using the antibodies of p63, CK7, and

Table 2: Performance evaluation of image registration by Hausdorff distances on a stack of 5 serial, whole-slide, immunohistochemical tissue sections by different protocols

| Stainings ($HR_0$ and $HT_{1-4}$) (Number) segmentation objects | Initial alignment | Patch-based nonlinear registration | Proposed multi-dimensional graph-based registration |
|---------------------------------------------------------------|------------------|-----------------------------------|---------------------------------------------------|
|                                                              | Average (μm)     | SD (μm)                           | Average (μm)                                     | Average (μm)                                      |
| $HR_0-HT_{1}$ (30)                                           | 222.5687         | 58.1487                           | 82.9659                                          | 48.6919                                           |
| $HR_0-HT_{2}$ (30)                                           | 139.2823         | 59.6697                           | 59.5939                                          | 48.5248                                           |
| $HR_0-HT_{3}$ (30)                                           | 396.6143         | 56.0818                           | 153.8545                                         | 43.748                                            |
| $HR_0-HT_{4}$ (30)                                           | 688.702          | 77.3843                           | 79.1539                                          | 54.7710                                           |
| Average (120)                                                | 361.7918         | 62.8211                           | 93.8921                                          | 48.9339                                           |

SD: Standard deviation, HR: High-resolution reference image, HT: High-resolution translated images

Figure 5: Comparing different methods in terms of the alignment accuracy and precision (10×). (a-d) Results of the original offset, with significant macrostructural alignment deviations were noted (red asterisks). (e-h) Registration results of the initial alignment. (i-l) Registration results obtained by applying patch-based nonlinear region. (m-p) Registration results obtained by applying the proposed method. The alignment accuracy and precision were gradually improved. Within the red circles are the examples of a hair follicle microstructure, showing the best alignment by the proposed method.
CK5/6 (mean Hausdorff distance 46.99 μm), while the worst alignment occurred in prostatic tissue stained with MAOa, where the alignment failed [mean Hausdorff distance >1000 μm, or Hausdorff distance uncalculable, Supplemental Figure 2], and the data were excluded (see Discussion).

**Discussion**

The precise alignment and registration of consecutive whole-slide histopathological section images is a key step leading to the future topographical proteomics and 3D reconstruction of disease tissues at the cell-level resolution. However, only few models and algorithms, including unsupervised content classification-based nonrigid registration and patch-based nonlinear image registration, were established for this purpose.\(^{[3,4,6,20]}\) These methods could be programmed and automatized, providing significant improvements compared with the conventional outline-based or manual-based image overlaying systems. However, in daily laboratory practice, minor issues were still witnessed, one of which was in the application of topographical proteomics.

In addition, the alignment between the series of strongly and differently stained whole-slide IHC images was usually a problem. The conventional hematoxylin- and eosin stain or special stains (such as Periodic acid–Schiff stain and Giemsa stain) were global stains, i.e., the dyes stained all the tissue elements by chemical reactions, although certain stains would highlight certain tissue structures. Under these circumstances, pattern recognition can be used for image alignment and registration.\(^{[7,9]}\) IHC stains, however, were nonglobal stains.

The principle of IHC stains was the antigen–antibody reaction, and thus, only cells that contained certain proteins could be stained. Although there existed a hematoxylin counterstain in the background, when the stained protein was highly expressed, the background hematoxylin stain would be masked and pose a problem when aligning the neighborhood IHC images, because they were usually mistaken as different objects. In this study, a multidimensional graph-based registration algorithm improved the unsupervised content classification-based nonrigid registration method to address this issue. Our proposed algorithm offered the advantages of rigid and nonrigid registration methods. Initializing the registration of high-resolution slides using a rigid method preserves the main morphology information and providing a deformation matrix generated from nonlinear low-resolution slide registration for following multidimensional graph-based registration provides high-resolution patch images for the local microstructural alignment.

By calculating the Hausdorff distances [see Table 1], we proved that the proposed method could improve the overall alignment accuracy (represented by the average Hausdorff distances) and precision (represented by the standard deviation of the Hausdorff distances). The published patch-based nonlinear registration method,\(^{[3]}\) despite showing a significant improvement from the initial alignment, encountered problems in capturing the specific matching feature points in most local tissue microstructures, because the stained protein distribution and intensity were different in each IHC slide. The proposed graph-based deformation model bypassed this problem through intensity-independent feature point caption and construction, thereby performing effectively. The overall performance improvement was 52.1% and 29.3% for the average and SD values with the Hausdorff distance, respectively.

However, the main limitation of the proposed method is the inherent tissue deviation due to manual serial sectioning. The neighboring two sections, in nature, are significantly alike but...
not the same. The average human cell size is approximately in
the range of 15–20 μm, and using the average and SD deviations
of the Housdorff distances (49 and 15 μm, respectively), a
deviation of three cells in accuracy and of one cell in precision
can be expected. Fortunately, in real human biology, cells with
similar or related functions usually work together as microunits,
such as in a microvessel, nerve, or folliculosebaceous unit,
which has a size >50 μm; additionally, with the microstructure
aligned, most functional topology proteomics can be correctly
interpreted [see Figures 5 and 7]. Most of the human biology
works through functional microunits rather than individual
cells. Some other modern methods, including next-generation
immunohistochemistry using mass cytometry,[21,22] multichannel
or multiplex immunofluorescence,[23‑25] or thick-section
confocal microscopy,[26] although could detect multiple protein
statives on a single slide in identical cells giving a highly
accurate evaluation on protein interactions, these methods are
usually expensive, labor intensive, limited in the interpreted
section size, require cell labeling, and demand high techniques,
thereby making them unlikely to be daily screening tools.
Interestingly, these methods were developed because in those
days “the conventional serial-section approach may have
instead approximated the cellular co-localization poorly owing
to heterogeneity effects.”[25] Now with our proposed method,
serial whole-slide IHC sections can be accurately and precisely
aligned. The “old-day” approach offers several clear advantages,
especially for screening purposes. First, whole-slide images can
be reviewed in a bright room (without requiring a dark room
for immunofluorescence), providing an overall view of the
proteome topology and facilitating pattern recognition. Second,
for each protein, the staining condition can be optimized
because the stain works on independent slides. By contrast,
there is usually a compromise in the staining condition when
multiple labels or stains work on the same slide. Third, most
pathologists worldwide are acquainted with the conventional
IHC images, thereby making the application of this method
highly practical from the daily usage viewpoint. Last, because
IHC is an essential part in daily pathology practice, even a local
hospital can provide all the needed instruments and slides for
the proposed method. No new instrument, reagent, or technique
is required.

The alignment quality, however, still heavily depends on the
quality of the antibody used. In this project, the best alignment
came from skin tissue sections stained with p63, CK7, and
CK5/6. These three antibodies gave highly specific and clear
staining signals with clean background and minimal cross
reaction, thus preserving the skin internal microstructures. By
contrast, the alignment failed in prostatic tissue sections stained
with MAOa. The antibody had intrinsic poor quality, giving a
blurred and nonspecific staining pattern with high background,
thus masking the underlying tissue microstructures.

Another limitation of the proposed method is that its ability to
treat extreme tissue torsion is still limited, as with other aligning
methods.[4] Tissue information based on consistent spatial
correlations may be incorrect because of the artifacts introduced
by cutting, folding, stretching, or tearing during tissue slicing.
Excessive extraction of registration feature points is possible
when artifacts are introduced so that mutually exclusive proteins
are brought together with variable overlaps. Additionally, the
proposed method, although provides accurate and precise
image alignment, show no apparent advance in terms of the
computation time (average 300 s per patch, as compared
with 320 s per patch for the published patch-based nonlinear
registration method). For screening purposes, the scanning
resolution could be decreased to reduce the computation time.

**Conclusions**

This study presented a three-step image-registration method to
efficiently align serial, whole-slide IHC tissue section images.

**Figure 7:** Pseudocolor system demonstrating the immunohistochemical protein expression for practical proteome topology: p63 (red), CK5/6 (green), EMA (blue), and CK7 (yellow) and HER2 (magenta) in protein staining (a–d) and background staining (e–h) areas. The stained areas in each whole-slide immunohistochemical image were extracted and transformed to a pseudocolor. The protein–protein coexpression or mutually exclusive expression could clearly be observed.
First, rigid registration was applied for the initial global alignment. Second, the patch-based nonlinear registration method was employed to perform low-resolution registration to achieve alignment between major tissue structures. Third, the proposed multidimensional graph-based registration, combined with a deformation matrix from patch-based nonlinear registration, was used to align the global and local deformation microstructures, improving the registration performance and robustness. Owing to its low cost and minimal requirements, our proposed method offers great advantages in screening the proteome topology in the daily laboratory practice.

**Financial support and sponsorship**

This work was supported by the Ministry of Science and Technology of Taiwan [grant numbers 105-2633-B-002-006- to C.W.L]; National Taiwan University Hospital [grant number 105-M3314 to C.W.L]; Taipei Institute of Pathology [TIP-109-004 to C.W.L].

**Conflicts of interest**

None declared.

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### Table S1: Antibodies used in creating whole-slide immunohistochemical tissue section images

| Antibody | Isotype | Clone | Source | Dilution | Retrieval |
|----------|---------|-------|--------|----------|-----------|
| BCL2     | Mouse IgG1 | 124   | Dako   | 1:200    | Citrate PH 6.0 |
| CD10     | Mouse IgG1 | 56C6  | Leica  | 1:30     | EDTA PH 9.0  |
| CD31     | Mouse IgG1 | 9G11  | BioGenex | 1:75     | EDTA PH 9.0  |
| CD34     | Mouse IgG1 | QBE10  | Dako   | 1:50     | EDTA PH 9.0  |
| CK5/6    | Mouse IgG1 | D5/16B4 | Thermo | 1:75     | EDTA PH 9.0  |
| CK7      | Mouse IgG1 | OV-TL 12/30 | Dako   | 1:300    | EDTA PH 9.0  |
| CK (AE1/3)| Mouse IgG1 | AE1/AE3 | BioGenex | 1:300    | EDTA PH 9.0  |
| D2-40    | Mouse IgG1 | D2-40  | Dako   | Ready to use | EDTA PH 9.0  |
| Desmin   | Mouse IgG1 | 33    | BioGenex | 1:250    | EDTA PH 9.0  |
| E-cadherin | Mouse IgG1 | NCH-38 | Dako | 1:400     | EDTA PH 9.0  |
| EMA      | Mouse IgG2a | E29   | Dako   | 1:200    | EDTA PH 9.0  |
| ERG      | Rabbit     | EP111  | Bio SB | 1:50     | EDTA PH 9.0  |
| GATA3    | Mouse      | L50-823 | Leica | 1:100    | EDTA PH 9.0  |
| Her-2    | Rabbit IgG | 4B5   | Ventana | Ready to use | EDTA PH 9.0  |
| MAOa     | Rabbit IgG | EPR7101 | Epitomics | 1:50     | EDTA PH 9.0  |
| MYC      | Rabbit IgG | Y69   | Abcam  | 1:50     | EDTA PH 9.0  |
| p40      | Rabbit IgG | Polyclonal | Biocare | 1:200    | EDTA PH 9.0  |
| p63      | Mouse IgG1 | DAK-p63 | Dako | 1:100     | EDTA PH 9.0  |
| PR       | Rabbit     | 1E2   | Ventana | 1:2      | EDTA PH 9.0  |
| PSA      | Mouse IgG1 | ER-PR8 | Dako   | 1:100    | EDTA PH 9.0  |
| S100     | Rabbit     | Z0311 | Dako   | 1:1800   | EDTA PH 9.0  |

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**Supplemental Figure 1:** Set design for consecutive, whole-slide, immunohistochemical section images. (a) Every set contained 1 Hematoxylin-and-Eosin stained slide and 5 immunohistochemical slides. The Hematoxylin-and-Eosin slide was the conventionally and globally stained slide used for histopathology interpretation. A total of 21 antibodies were successfully applied on five different kinds of tissues. (b-d) Examples of consecutive, whole-slide, immunohistochemical sections: (b) EMA staining, (c) Desmin staining, and (d) CK (AE1/3) staining.
Supplemental Figure 2: Aligning sections stained with MAOa in prostatic tissue. (a) The MAOa antibody showed high background staining across the prostatic tissue, masking the underlying tissue microstructures. (b) The low-resolution image extracted for nonlinear registration after discretization. (c) A pseudocolor system showed the result of aligning other immunohistochemical sections with the section stained with MAOa. The results were of poor accuracy and precision using published and proposed methods.