Microarray BASICA: Background Adjustment, Segmentation, Image Compression and Analysis of Microarray Images

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This paper presents microarray BASICA: an integrated image processing tool for background adjustment, segmentation, image compression, and analysis of cDNA microarray images. BASICA uses a fast Mann-Whitney test-based algorithm to segment cDNA microarray images and performs postprocessing to eliminate the segmentation irregularities. The segmentation results, along with the foreground and background intensities obtained with the background adjustment, are then used for independent compression of the foreground and background. We introduce a new distortion measurement for cDNA microarray image compression and devise a coding scheme by modifying the embedded block coding with optimized truncation (EBCOT) algorithm (Taubman, 2000) to achieve optimal rate-distortion performance in lossy coding while still maintaining outstanding lossless compression performance. Experimental results show that the bit rate required to ensure sufficiently accurate gene expression measurement varies and depends on the quality of cDNA microarray images. For homogeneously hybridized cDNA microarray images, BASICA is able to provide from a bit rate as low as 5 bpp the gene expression data that are 99% in agreement with those of the original 32 bpp images.

Keywords and phrases: microarray BASICA, segmentation, Mann-Whitney test, lossy-to-lossless compression, EBCOT.

1. INTRODUCTION

The cDNA microarray technology is a hybridization-based process that can quantitatively characterize the relative abundance of gene transcripts [1, 2]. Contrary to conventional methods, microarray technology promises to monitor the transcript production of thousands of genes or even the whole genome simultaneously. It thus provides a new and powerful enabling tool for genetic research and drug discovery. To produce cDNA microarrays, the mRNA of the control and test samples are first reverse-transcribed into cDNA and fluorescently labeled with different dyes (typically red and green). Then the fluorescent targets are mixed and allowed to hybridize with gene-specific cDNA clones printed in an array format on a glass microslide. Finally, by scanning the microslide with a laser and capturing the photons emitted from different dyes into different channels with a confocal fluorescence microscope, a two-channel 16-bit microarray
image is obtained in which the pixel intensities reflect the level of mRNA expression. Usually a microarray image is shown RGB composite format, where the red and green channels correspond to the two channels of the microarray image obtained while the blue channel is set to zero. With the help of signal processing and data analysis operations such as ratio statistics, classification, and genetic regulatory network design, microarray images can shed light on the possible regulation rules of transcription production sought by biologists and clinicians.

Microarray images cannot be used for genetic data analysis directly. Appropriate image processing procedures are to be performed in order to extract information from the images for downstream analysis. Thousands of cDNA target sites must first be identified as the foreground by an image segmentation algorithm. Then the intensity pair \((R,G)\) that represents gene expression levels of both channels is extracted from every foreground target site with appropriate background adjustment. Subsequent data analysis is normally conducted based on the log ratio \(\log R/G\) of the intensity pair. As the very first step of CMA microarray signal processing, the accuracy of image processing is critical to the reliability of subsequent data analysis. Many image processing schemes have been developed for this purpose in recent years and can be found in various commercial and non-commercial software packages [3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19]. Generally, because each channel of the microarray image is typically more than 15 MB in size, highly efficient compression is necessary for data backup and communication purposes. In order to save storage space and alleviate the transmission burden for data sharing, the search for good progressive compression schemes that provide sufficiently accurate genetic information for data analysis at low bit rates while still ensuring good lossless compression performance has become the focus of CMA microarray image compression research recently [3, 4, 20].

This paper introduces a new integrated system called microarray BASICA. BASICA brings together the image processing procedures required to accomplish the aforementioned information extraction and data analysis, including background adjustment, segmentation, and compression. A fast Mann-Whitney test-based algorithm is presented for the initial segmentation of CMA microarray images. This new algorithm can save up to 50 times the number of repetitions required from the original algorithm [5]. The resulting images are then postprocessed to remove the segmentation irregularities. The segmentation results, along with the foreground and background intensities, are saved into a header file for both data analysis and compression. A novel distortion measure is introduced to evaluate the accuracy of extracted information. Based on this measure and the information provided by the header file, a new image compression scheme is designed by modifying the embedded block coding with optimized truncation (EBCOT) algorithm [3], which is now incorporated in the JPEG2000 standard. Our experiments show that there appears to be no common bit rate that ensures sufficiently accurate gene expression data for different CMA microarray images. On CMA microarray images of good quality, BASICA is able to provide from a bit rate as low as 5 bpp (bit per pixel) the gene expression data that are 99% in agreement with those of the original 32 bpp images.

2. DETAILS OF MICROARRAY BASICA

Microarray BASICA provides solutions to both processing and compression of CMA microarray images. The major components of BASICA and their relationship with the elements of a microarray experiment are shown in Figure 1. Each two-channel microarray image acquired through the laser scanner is first sent to the segmentation component, where the target sites are identified. With the result of segmentation, the background adjustment component estimates each spot’s foreground and background intensities and calculates the log ratio values based on the background-subtracted intensities. After this, the calculated log ratio values along with the segmentation information and other necessary data related to each spot are output for downstream data analysis. In the mean time, BASICA compiles the segmentation result and extracted intensities into a header file. With this header file, the compression component encodes the foreground and background of both channels of the original image into progressive bitstreams separately. The generated bitstreams, plus the header file, are saved into a data archive for future access or are transmitted as shared data. On the other hand, to utilize the archived or transmitted data, BASICA can either quickly retrieve the necessary genetic information saved in the header file or reconstruct the microarray image with available bitstreams through the reconstruction component and redo the segmentation and background adjustment.

2.1. Segmentation with postprocessing

Segmentation is performed to identify the target sites in each spot where the hybridization occurs. In [8], various existing segmentation schemes are summarized and categorized into four groups: (1) fixed circle segmentation, (2) adaptive circle segmentation, (3) adaptive shape segmentation, and (4) histogram segmentation.

Although the shape of a target site is determined by the physical attributes of the DNA probes and the mechanism of the printing procedure, most target sites are round or donut-like in shape. The fixed circle segmentation, which sets a round region of constant diameter in the middle of each spot as the target site, appears to be the most straightforward method and is provided in most existing software packages [9, 11, 12, 13, 17, 18]. The radius of the foreground is set either by a default value as a parameter of the robot arrayer and laser scanner or empirically determined by the user. The fixed circle method runs fast and performs well when the microarray spots are perfectly hybridized and aligned. In practical cases, however, the spots are far from perfect due to unpredictable nonuniform hybridization across the spot or misalignment of the probe array. GenePix [13] uses the adaptive circle segmentation to accommodate the varying sizes of different target sites, and Dapple [11] finds the best matched
Neither the fixed nor the adaptive circle segmentation can accommodate the variances in shape of the target sites in the images. To tackle this problem, more accurate and sophisticated segmentation methods are needed. The segmentation technique introduced in [8] uses seeded region growing [21], while other methods [5, 6, 10, 15, 17, 19] rely on more conventional histogram-based segmentation algorithms. The histogram-based methods generally compute a histogram of pixel intensities for each spot. Methods in [10, 17, 19] adopt a percentile-based approach, which sets the pixels in a high percentile range of the histogram as the foreground and those in a low range as the background. Methods in [6, 15] use a threshold-based approach. To ensure correct segmentation, methods in [10, 15] employ repetitions to find the most stable segmentation. The histogram-based segmentation demonstrates good performance when a target site has a high hybridization rate, that is, a high intensity. However, the intensities of most target sites are actually very close to the local background intensities, and it is hard to segment correctly by finding a threshold based on the histogram only.

In an attempt to solve this problem, Chen et al. introduced a Mann-Whitney test-based segmentation method in [5]. So far, no single segmentation algorithm can meet the demands of all microarray images. Segmentation algorithms are normally designed to perform well on microarray images acquired by certain type of arrayers and scanners. It is therefore hard to compare them directly.

### 2.1.1. Mann-Whitney test-based segmentation

In BASICA, we use the Mann-Whitney test-based segmentation algorithm introduced by Chen et al. in [5]. The Mann-Whitney test is a distribution-free rank-based two-sample test, which can be applied to various intensity distributions caused by irregular hybridization processes that are difficult to handle by conventional thresholding methods. Here we first give a brief description of the Mann-Whitney test-based segmentation algorithm.

Consider two independent sample sets $X$ and $Y$. Samples $X_1, X_2, \ldots, X_m$ are randomly selected from set $X$, and $Y_1, Y_2, \ldots, Y_n$ are randomly selected from set $Y$. All $N = m + n$ samples are sorted and ranked. Denote $R_i$ as the rank of the $i$th sample, $R(X_i)$ as the rank of sample $X_i$, and $R(Y_i)$ as the rank of $Y_i$. These ranks are used to test the following hypotheses:

- (H$_0$) $P(X < Y) \geq 0.5$,
- (H$_1$) $P(X < Y) < 0.5$.

Define the rank sum of the $m$ samples from $X$ as

$$T = \sum_{i=1}^{m} R(X_i). \quad (1)$$

To avoid deviations caused by ties, $T$ is commonly normalized as

$$\overline{T} = \sqrt{\frac{m(m+1)/2}{nm/N(N-1)\sum_{i=1}^{N} R_i^2 - nm(N+1)^2/4(N-1)}}. \quad (2)$$

Hypothesis (H$_0$) will be rejected if $\overline{T}$ is greater than a certain quantile $w_{1-\alpha}$, where $\alpha$ is the significance level.

In microarray image segmentation, hypothesis (H$_1$) corresponds to the case that set $X$ is the high-intensity foreground and set $Y$ is the low-intensity background, and...
hypothesis \( (H_0) \) corresponds to the reverse case. To segment a target spot, a predefined target mask (obtained by selecting, unifying, and thresholding strong targets) is first applied to the spot. Pixels inside the mask correspond to set \( X \), and pixels outside correspond to set \( Y \). To start the test, \( n \) samples are randomly selected from set \( Y \), while \( m \) samples with lowest intensities are selected from set \( X \). If hypothesis \( (H_0) \) is accepted, the pixel with lowest intensity is removed from set \( X \), and \( m \) sample pixels are reselected. The test is repeated until hypothesis \( (H_0) \) is rejected. Then the pixels left in set \( X \) are considered as the foreground at significance level \( \alpha \). The foregrounds obtained from the two channels are united into one to produce the final segmentation result.

The repetitive nature of this algorithm makes it cumbersome for real-time implementation. So, in BASICA, we proposed a fast Mann-Whitney test-based algorithm \cite{3} which runs much faster while generating identical segmentation results.

2.1.2. Speeding up Mann-Whitney test-based segmentation algorithm

Assume that the predefined target mask is obtained according to the way described in \cite{5, 6}. Samples \( X_1, X_2, \ldots, X_m \) and \( Y_1, Y_2, \ldots, Y_n \) are picked from the foreground and background, respectively. Without loss of generality, it suffices to assume that \( X_1 \leq X_2 \leq \cdots \leq X_m \) and \( Y_1 \leq Y_2 \leq \cdots \leq Y_n \). Since \( X_1, X_2, \ldots, X_m \) are the smallest \( m \) samples in set \( X \), all other samples can be determined if \( X_1 \) is set. Then Mann-Whitney test-based segmentation is actually an optimization problem of minimizing \( X_1 \) subject to \( T \geq x_{1-a} \). Chen et al.'s approach takes a large number of repetitions to reach the final segmentation. However, it turns out that the number of repetitions can be significantly reduced by carefully choosing the starting point and search strategy.

BASICA first finds an upper bound of the optimal \( X_1 \), denoted by \( X_1^{\text{max}} \), which is related to \( Y_1, Y_2, \ldots, Y_n \). With (2), \( T \geq x_{1-a} \) can be written as

\[
\sum_{i=1}^{m} R(X_i) \geq x_{1-a} \sqrt{\frac{nm}{N(N-1)} \sum_{i=1}^{N} R_i^2 - \frac{nm(N+1)^2}{4(N-1)} + m - \frac{N+1}{2}}.
\]

(3)

In the right-hand side of (3), only \( \sum_{i=1}^{N} R_i^2 \) is associated with \( X_1 \). If no tie exists, the ranks are from 1 to \( N \) and the sum is \( \sum_{i=1}^{N} R_i^2 \). If there is a tie, the ranks of the tied samples are the average of those ranks if there would have been no tie, and induce a reduction on the sum. A property of this reduction is that it is only related to the number of samples tied at that value. If there are \( k \) samples having the same value, the reduction is \( (1/12)(k^3 - k) \). With this property, one can easily reduce the upper bound of \( \sum_{i=1}^{N} R_i^2 \). Assume that \( \Delta Y \) is the decrease in the sum caused by the ties in the sorted \( Y_1, Y_2, \ldots, Y_n \), then we have

\[
\sum_{i=1}^{N} R_i^2 \leq \sum_{i=1}^{N} R_i^2 - \Delta Y,
\]

(4)

where the equation holds when \( X_1, X_2, \ldots, X_m \) have no tie among themselves and share no tie with any sample in \( Y_1, Y_2, \ldots, Y_n \). In most cases, the difference is very small and the bound is quite tight.

To simplify the notation, we use \( \sigma_{\text{max}} \) to denote

\[
\frac{nm}{N(N-1)} \left( \sum_{i=1}^{N} i^2 - \Delta Y \right) - \frac{nm(N+1)^2}{4(N-1)}.
\]

(5)

Then \( X_1^{\text{max}} \) must satisfy the inequality

\[
\sum_{i=1}^{m} R(X_i) \geq w_{1-a} \sigma_{\text{max}} + \frac{N+1}{2}.
\]

(6)

no matter what \( X_2, X_3, \ldots, X_m \) can be for as long as the assumption \( X_1 \leq X_2 \leq \cdots \leq X_m \) holds. So, to find \( X_1^{\text{max}} \) is to find the smallest \( X_1 \) so that the smallest rank sum of \( X_1 \leq X_2 \leq \cdots \leq X_m \) still satisfies inequality (6). To associate \( X_1^{\text{max}} \) with known information \( Y_1, Y_2, \ldots, Y_n \), assume that \( Y_u < X_1 \). Then the minimum rank sum is \( \sum_{i=1}^{m} R(X_i) = \sum_{i=1}^{m} (u+i) \), when \( u < X_1 \leq X_2 \leq \cdots \leq X_m < Y_u+1 \). By solving the inequality with \( \sum_{i=1}^{m} R(X_i) = \sum_{i=1}^{m} (u+i) \), \( u \) can be obtained as

\[
u = \left[ \frac{w_{1-a} \sigma_{\text{max}} + n}{m} \right].
\]

(7)

Thus, the upper bound \( X_1^{\text{max}} \) is the smallest sample in \( X \) that is larger than \( Y_u \). For any sample set \( X_1, X_2, \ldots, X_m \) with \( X_1 \geq X_1^{\text{max}} \), hypothesis \( (H_0) \) can be rejected outright. Since \( X_1, X_2, \ldots, X_m \) normally have similar intensities which bring on consecutive ranks, \( X_1^{\text{max}} \) is very close to the actual threshold. Hence, the repetitions can be greatly reduced if backward repetitions based on \( X_1^{\text{max}} \) are applied.

Besides changing the starting point and repetition direction, a two-tier repetition strategy can be used to reduce the repetition in case the upper bound is not so tight as expected. In the first tier, one does not perform the repetition in a pixel-by-pixel manner, but in a leaping manner instead. Then a pixel-by-pixel repetition follows up and locates the exact segmentation in the second tier. Larger step size means fewer repetitions in the first tier but more in the second, while smaller step size has the opposite effect. A natural choice of the repetition steps is indicated by \( Y_1, Y_2, \ldots, Y_n \) when \( n \) is not very large. The whole algorithm is described as follows.

Step 1: Calculate \( u \) using (7).

Step 2: Find the smallest \( m \) samples from set \( X \) that are larger than \( Y_u \), and execute the Mann-Whitney test.

Step 3: If hypothesis \( (H_0) \) is rejected, then set \( u = u-1 \) and go to step 2, otherwise, go to step 4.

Step 4: \( u = u+1 \). Find the smallest \( m \) samples from set \( X \) that are larger than \( Y_u \), and begin the pixel-by-pixel repetition in backward manner.

It should be noted that this modified Mann-Whitney test-based segmentation algorithm may not always generate identical results with Chen et al.’s original algorithm. In order
Table 1: The comparisons of the number of repetitions between Chen et al.’s algorithm and our modified method used in BASICA at different significance levels.

<table>
<thead>
<tr>
<th>α</th>
<th>0.001</th>
<th>0.005</th>
<th>0.01</th>
<th>0.05</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chen et al.</td>
<td>328.7</td>
<td>269.1</td>
<td>270.9</td>
<td>226.3</td>
</tr>
<tr>
<td>BASICA</td>
<td>7.5</td>
<td>7.3</td>
<td>5.9</td>
<td>3.7</td>
</tr>
</tbody>
</table>

to obtain identical results, the backward-searching nature of the new algorithm requires the normalized rank sum in (2) to be strictly increasing during the repetition of the original algorithm. This is not guaranteed due to the occurrences of ties in the sorted samples. In one extreme case, when all $N$ samples have the same intensity, the devisor will become zero and the normalized rank sum will be infinity. Actually, Chen et al.’s original algorithm can be viewed as trying to find the largest foreground that rejects hypothesis $H_0$, while the modified algorithm in BASICA tries to find the smallest foreground that accepts the hypothesis $H_0$. Since in most cases the normalized rank sum will be strictly increasing, we expect the segmentation results of the modified algorithm to be identical to the original algorithm most of the time.

The comparisons of the number of required repetitions between Chen et al.’s algorithm and our modified algorithm are given in Table 1. Results are averaged over 504 spots in both channels from different test images. Both algorithms set $m = n = 8$ and use the same randomly selected samples from the predefined background for the Mann-Whitney test. We find that the segmentation results on all test spots of the sample images used in this study are identical between the original algorithm and the modified algorithm. From the table, we observe that the modified algorithm reduces the number of repetitions by up to 50 times from what is required of the original algorithm.

### 2.1.3. Postprocessing

Like common threshold-based segmentation algorithms, there are always many annoying shape irregularities in the segmentation results obtained by the Mann-Whitney test-based algorithms. These irregularities occur randomly and can severely reduce the compression efficiency. Thus, an appropriate postprocessing procedure is necessary to achieve efficient compression. Moreover, because most irregularities are pixels with a high probability of noise corruption, eliminating them is unlikely to compromise the accuracy of subsequent data extraction and analysis.

In BASICA, we categorize possible irregularities into two types and employ different methods to eliminate them. The first type includes isolated noisy pixels or tiny regions, which can be observed from the lower half of the segmentation result in Figure 2a. These irregularities are caused usually by nonspecific hybridization or undesired binding of fluorescent dyes to the glass surface. The second type includes the small branches attached to the large consolidated foreground regions, which are visible in the segmentation results of Figure 2. Since these irregularities are located between the foreground and background, their intensities are also inbetween, making them vulnerable to noise corruption. The irregularities in most segmentation results are usually made up of both these two types. For the first type, BASICA will detect and remove them directly from the foreground. As for the second type, BASICA applies an operation similar to the standard morphological pruning [22]. By removing and pruning repetitively, BASICA can successfully eliminate most irregularities in three to five repetitions. The right column of Figure 2 shows the postprocessing results on the original segmentation which are to be used for the compression of the images. Figure 3 shows a portion of a microarray image and its segmentation results.

### 2.2. Background adjustment

It is commonly believed that the pixel intensity of the foreground reflects the joint effects of the fluorescence and the glass surface. To obtain the expression level accurately, the intensity bias caused by the glass surface should be estimated and subtracted from the foreground intensity, and this process is known as background adjustment. Since there is no hybridization in the background area, the background intensity is normally measured and treated as an intensity bias. Although mean pixel intensity has been adopted in almost all existing schemes as the foreground intensity, several methods have been developed for background intensity estimation. The major differences of various methods lie in two aspects: (1) on which pixels the estimation is based and (2) how to calculate the estimation. Regarding the first aspect, the regions chosen for background estimation vary from a global background to a local background. For the global background, the background regions in all spots are considered, and a global background intensity is estimated and subtracted from every foreground intensity [9, 16]. The global background ignores possible variance between subarrays and spots. So, in [9], partial global background estimation is performed based on the background of one subarray or on several manually selected spots. The more common approach is to estimate the background intensity based on the local background for each target site separately. The local background can be the entire background region in one spot [18], or, to avoid interference from the foreground, it can be the region with a certain distance from the foreground target site [7, 11, 13, 15, 17]. In the extreme case, the algorithm in [14] uses the pixels on the border of each spot as the local background. However, using too few pixels increases the possibility of a large variance in background estimation. As to the second aspect, almost all existing systems adopt mean or median to measure the expression level. Besides these, mode and minimum are also used in some softwares [6, 16]. Unlike all the methods mentioned above, a morphological opening operation is performed in [8] to smooth the whole background and then estimate the background by sampling at the center of the spot.

Some commercial software packages [9, 16] offer more than one choice for background adjustment. ArrayMetrix [9] provides up to nine methods, while ArrayVision [10]
Figure 2: Segmentation and postprocessing of two typical spots. The left column shows the original microarray spots in RGB composite format. Some intensity adjustments are applied in order to show them clearly. The middle column shows the corresponding segmentation results using the Mann-Whitney test with significance level $\alpha = 0.001$. The right column shows the final segmentation results after postprocessing.

Figure 3: (a) Part of a typical cDNA microarray image in RGB composite format. Some intensity adjustments were applied in order to show the image clearly. (b) The segmentation results of (a).

provides seven ways of background region determination and six choices of averaging method. Experiments in [8] show that different background adjustment methods have significant impact on the log ratio values subsequently obtained. However, there is no known criterion to measure which approach is more accurate than the others.

BASICA chooses the average of pixel intensities in the local background as the estimate of background intensity. To prevent possible biases caused by either the higher intensity values of the pixels adjacent to the foreground target sites or the lower intensity values of the dark hole regions in the middle of the spots, the local background used in BASICA is the background defined by the predefined target mask obtained through the segmentation.

2.3. Data analysis

Because so many elements impact the pixel intensities of the microarray image, genetic researchers do not use the absolute intensities of the two channels, but the ratio between them to measure the relative abundance of gene transcription. Not all genetic information extracted are reliable enough for data analysis. If the spot has so poor quality that no reliable information can be extracted, it is qualified as a false spot; otherwise, it is a valid spot. For a valid spot $k$, the expression ratio is denoted by

$$T_k = \frac{R_k}{G_k} = \frac{\mu_{FR_k} - \mu_{BR_k}}{\mu_{FG_k} - \mu_{BG_k}},$$

where $R_k$ and $G_k$ are the background-subtracted mean intensities of the red and green channels, respectively, $\mu_{FR_k}$ and $\mu_{FG_k}$ are the respective foreground mean intensities, and $\mu_{BR_k}$ and $\mu_{BG_k}$ are the respective background mean intensities.
and $\mu_{BG}$ are the respective estimated background mean intensities. Because expression ratio has an unsymmetric distribution, which contradicts the basic assumptions of most statistic tests, the log ratio $\log T_k = \log R_k/G_k$ is commonly used instead in most applications. In addition to the log ratio, an auxiliary measure which is often helpful in data analysis is the log product $\log R_k G_k$. However, since the log transform does not have constant variance at different expression levels, some alternative transforms like $g \log$ [23] have recently been introduced. In gene expression studies, such transformed ratios are ordinarily normalized and quantized into three classes: down-regulated, up-regulated, and invariant. Expression level extraction and quantization provide the starting point for subsequent high-level data analysis, and their accuracy is crucially important. Therefore, compression schemes should be designed to minimize the distortion in the image, and their performance should be assessed by agreement/disagreement in gene expression level measurement caused by the compression. These topics will be discussed in detail in Sections 2.4 and 3.3.

### 2.4. Image compression

Since microarray images contain huge amounts of data and are usually stored at the resolution of 16 bpp, a two-channel microarray image is typically between 32 and 64 MB in size. Efficient compression methods are highly desired to accommodate the rapid growth of microarray images and to reduce the storage and transmission costs. Currently, the common method to archive microarray images is to store them losslessly in TIFF format with LZW compression [24]. However, such an approach does not exploit 2D correlation of data between pixels and does not support lossy compression. Due to the huge data size, microarray images require efficient compression algorithms which support not only lossless compression but also lossy compression with graceful degradation of image quality for downstream data analysis at low bit rates.

Recently, a new method known as the segmented LOCO (SLOCO) was introduced in [20]. This method exploits the possibility of lossy-to-lossless compression for microarray images. SLOCO is based on the LOCO-I algorithm [25], which has been incorporated in the lossless/near-lossless compression standard of JPEG-LS. SLOCO employs a two-tier coding structure. It first encodes microarray images losslessly in near-lossless compression but also loss compression with graceful degradation of image quality for downstream data analysis at low bit rates.

In BASICA, we also incorporate lossy-to-lossless compression of microarray images. The aims of compression in BASICA are twofold: (1) to generate progressive bitstreams that can fulfill the requirements of signal processing and data analysis at low bit rates for data sharing and transmission applications and (2) to deliver competitive lossless compression performance to data archiving applications with a progressive bitstream. To achieve these objectives, the compression scheme in BASICA treats the foreground and background of microarray images separately. Obviously, the foreground and background usually have significant intensity differences and they are relatively homogeneous in their corresponding local regions. Hence, by compressing the foreground and background separately, the compression efficiency is expected to improve significantly. This is done by utilizing the outcomes of segmentation. Before encoding, BASICA saves all necessary segmentation information into a header file for subsequent compression.

SLOCO in [20] is based on spatial domain predictive coding. In contrast, BASICA employs bit-plane coding in the transform domain. Bit-plane coding enables BASICA to achieve truly progressive bitstream coding at any rate. To allow lossy compression, an appropriate distortion measurement is needed. Generally, medical image compression requires visually imperceptible differences between the lossily reconstructed image and the original. Traditional distortion measures, such as mean square error (MSE), are poor indicators for this purpose. However, unlike other types of medical images, the performance of microarray image compression does not depend on visual quality judgement, but instead on the accuracy of final data analysis. Therefore, it is reasonable to adopt a distortion measure adherent to the requirements of data analysis. Since almost all existing data analysis methods use the transformed expression values, we should seek to minimize the distortion under these measurements. In BASICA, we adopt distortion measures based on the log ratios and the log products because they are the most used transforms in common applications. However, as we will see later, the scheme employed in BASICA can be easily adapted for other transform measures.

The log ratios and the log products decouple the data of two channels into two separate log intensities, $\log R$ and $\log G$. This ensures that the compression can be done on each channel independently. Without loss of generality, we only refer to the $R$ channel in the rest of the paper.

BASICA currently employs the MSE of $\log R$ as the distortion measurement, which is defined as

$$MSE_{\log R} = \frac{1}{N} \sum_{i=1}^{N} (\log R_i - \log \hat{R}_i)^2,$$

where $N$ is the total number of pixels in the microarray image, and $R_i$ and $\hat{R}_i$ are background-subtracted mean intensities obtained from spot $i$ of the original and reconstructed image, respectively.

There is a direct relationship between the MSE of log intensity and the traditional MSE. For spot $k$, its log intensity $\log R_k$ can be further written as

$$\log R_k = \log (\mu_{FR_k} - \mu_{BR_k}) = \log \left( \frac{1}{M_k} \sum_{i=1}^{M_k} X_i - \mu_{BR_k} \right),$$

where $M_k$ is the total number of pixels in the foreground of spot $k$, and $X_i$ is the intensity of the $i$th pixel. So the unit error
\[ \Delta \log R_k \text{ is associated with the unit error } \Delta X_j \text{ of } j\text{th pixel by} \]
\[ \Delta \log R_k = \frac{\Delta X_j}{M_k(\mu_{FR_k} - \mu_{BR_k})}. \] (11)

For the pixels in the background, because most existing schemes do not compute the average intensity as \( \mu_{BR_k} \), but use nonlinear operations such as modulo or median filtering, the above derivation no longer holds. The foreground and background pixels have different impacts on the log intensity and should be considered separately.

Equation (11) indicates that the MSE of log intensity is actually a weighted version of traditional MSE. The weight \( 1/M_k(\mu_{FR_k} - \mu_{BR_k}) \) is a constant for pixels in the same spot and is inversely proportional to the spot’s intensity and foreground size. The higher a spot’s intensity or foreground size, the larger its allowable reconstruction error.

Quite similarly, one can easily derive other MSE distortion measurements for other transforms. For example, the \( g \log \) transform in [23] is
\[ g(R_k) = \log \left( \mu_{FR_k} - \alpha + \sqrt{(\mu_{FR_k} - \alpha)^2 + \epsilon} \right), \] (12)
where \( \alpha \) and \( \epsilon \) are parameters estimated from the microarray image. Then, with straightforward derivation, one can associate the unit error \( \Delta g(R_k) \) with the unit error \( \Delta X_j \) of \( j\text{th pixel by} \)
\[ \Delta g(R_k) = \frac{\Delta X_j}{M_k \sqrt{(\mu_{FR_k} - \alpha)^2 + \epsilon}}. \] (13)

Thus, the MSE of \( g \log \) is also a weighted version of traditional MSE, and like MSE of log ratio, the measurement allows larger distortions in spots of high intensities.

Although we can derive different distortion measurements for different transforms, the compression scheme in BASICA can only be designed based on one type of distortion measurement. As mentioned before, in BASICA we choose MSE of log ratio as the distortion measurement.

With the help of (11), we introduce a new lossy-to-lossless compression scheme in BASICA by modifying EBCOT [26] with several techniques specifically designed for the requirements of microarray technology. First, to encode the foreground and background separately, we modify the EBCOT context modeling according to different distortion measures for different transforms. For the pixels in the background, because most existing schemes do not compute the average intensity as \( \mu_{BR_k} \), but use nonlinear operations such as modulo or median filtering, the above derivation no longer holds. The foreground and background pixels have different impacts on the log intensity and should be considered separately.

Equation (11) indicates that the MSE of log intensity is actually a weighted version of traditional MSE. The weight \( 1/M_k(\mu_{FR_k} - \mu_{BR_k}) \) is a constant for pixels in the same spot and is inversely proportional to the spot’s intensity and foreground size. The higher a spot’s intensity or foreground size, the larger its allowable reconstruction error.

Quite similarly, one can easily derive other MSE distortion measurements for other transforms. For example, the \( g \log \) transform in [23] is
\[ g(R_k) = \log \left( \mu_{FR_k} - \alpha + \sqrt{(\mu_{FR_k} - \alpha)^2 + \epsilon} \right), \] (12)
where \( \alpha \) and \( \epsilon \) are parameters estimated from the microarray image. Then, with straightforward derivation, one can associate the unit error \( \Delta g(R_k) \) with the unit error \( \Delta X_j \) of \( j\text{th pixel by} \)
\[ \Delta g(R_k) = \frac{\Delta X_j}{M_k \sqrt{(\mu_{FR_k} - \alpha)^2 + \epsilon}}. \] (13)

Thus, the MSE of \( g \log \) is also a weighted version of traditional MSE, and like MSE of log ratio, the measurement allows larger distortions in spots of high intensities.

2.4.1. Modifying EBCOT for microarray image coding

Our major modifications to EBCOT are the following.

**Header file.** A header file is necessary for saving the information which will be used in the encoding and decoding procedures. To ensure that the encoder and decoder can correctly compress and reconstruct the foreground and background independently, the segmentation information must be saved in the header file. Besides, (11) indicates that the mean intensities of the foreground and background are also needed by the compression algorithm. To save storage memory, these data are coded with LZW compression. Although the segmentation information and spot intensities are enough for the compression component, other data, such as variances of pixel intensities in each spot, can also be saved in the header file for quick genetic information retrieval. In the practical implementation, the header file will be generated before encoding and must be transmitted and decoded first.

**Shape-adaptive integer wavelet transform.** Like other frequency-domain-based coding schemes, in BASICA the transform is performed before bit-plane coding during the encoding phase and after the bit-plane reconstruction during the decoding phase. To ensure lossless compression, integer wavelet transforms are required. The wavelet transforms are conducted on the foreground and background independently to prevent any interference between the coefficients from adjacent areas. Since the segmented foreground and background always have irregular shapes, critically sampled integer wavelet transforms for arbitrarily shaped objects are needed to ensure coding efficiency. Many approaches have been proposed for 2D shape-adaptive wavelet transforms. Our proposed coding scheme uses odd-symmetric extensions over object boundaries described in [27].

**Object-based EBCOT.** After shape-adaptive integer wavelet transform, we modify the EBCOT context modeling for arbitrarily shaped regions. The extension of EBCOT algorithm to shape-adaptive coding is rather straightforward. Because the shape-adaptive integer wavelet transform is critically sampled, the number of wavelet coefficients is the same as those in the original regions. Using the wavelet-domain shape mask, one can easily tell whether a coefficient belongs to a region to be coded. If any neighbor of that coefficient falls outside the region, we just set that neighboring coefficient’s values to zero, thus making it insignificant in context modeling. We call the resulting coder object-based EBCOT.

**Intensity shifts.** To minimize the initial MSE, the average intensity of the image is subtracted from each pixel before encoding and added back after decoding. Unlike eight-bit natural images, the foreground of a microarray image normally has an exponential intensity distribution. The exponential distribution property of the foreground makes the global average intensity subtraction less effective. However, the pixels...
in the foreground of any spot \( k \) normally have similar intensities and roughly have a symmetric distribution around \( \mu_{FR_k} \). So, for the encoding of the foreground, each pixel in spot \( k \) is subtracted \( \mu_{FR_k} \) instead of the global average intensity. Since \( \mu_{FR_k} \) is already saved in the header file, intensity shifts do not cost any overhead. With intensity shifts, the distribution of foreground intensities are transformed into a symmetric shape with a high peak around zero. As for the background compression, through our experiments, we find that the pixels in the background actually have a roughly symmetric intensity distribution, suggesting that the global average intensity subtraction will be appropriate.

**Bit shifts.** EBCOT uses block-based bit-plane coding. In order to minimize the distortions at different rates, one must code the bit planes of different spots according to their impacts on the MSE of log intensity. One straightforward solution is to scale the coefficients of each spot with the spot’s weight, so bits at the same bit plane of all spots have the same impacts on the MSE of log intensity. However, because the weights are noninteger fractions, lossless compression cannot be ensured under such a scaling. Furthermore, although one can round them to the closest integer as an approximation, any scaler \( w \) will increase a coefficient’s information up to \( \lceil \log_2(w) \rceil \) bits, which can lead to a very poor lossless compression performance. In BASICA, we apply the scaling by bit shifts, which is a good approximation and meanwhile does not compromise the performance of lossless compression. For spot \( k \), BASICA obtains

\[
S_k = \left\lfloor \log_2 \left( \frac{M_k(\mu_{FR_k} - \mu_{BR_k})}{w} \right) \right\rfloor + 0.5.
\]

Let \( S_{\text{max}} = \max(S_1, S_2, \ldots, S_N) \). Then it scales the coefficients of spot \( k \) by upshifting them \( S_{\text{max}} - S_k \) bits.

**Background compression.** With careful consideration, bit shifts have not been applied in the background compression in BASICA for several reasons. First, since there exist different approaches to compute the background intensity, and the values obtained by these methods also vary a lot, it is unclear how to find a unique weight for each pixel like what BASICA has for foreground compression. Second, unlike isolated target sites in the foreground, the local background is normally connected to each other. Thus, bit shifts will bring abrupt intensity changes along the borders of spots, which will in turn lower the compression efficiency significantly in lossless coding performance. Even though one can figure out the weights through a formula similar to (11) based on certain background extraction methods, there will be a significant trade-off on lossless compression, which is about 0.8 bpp according to our experiments. So, in BASICA, we apply a global average intensity subtraction and no bit shifts on the background compression, that is, the traditional MSEE measure is used for rate-distortion optimization. Normally, the pixel intensities in the background are located in a very small range, which means that the background is pretty homogenous. Thus, compression with traditional MSEE measure should be able to represent the background with fairly small bit rates.

To this end, the final code of a two-channel microarray image is composed of five different parts: a header file and two bitstreams representing the foreground and background, respectively, from each channel.

### 3. EXPERIMENTAL RESULTS AND DISCUSSION

Experiments have been conducted to test the performance of BASICA with eight microarray images from two different sources. We used three test images from the National Institutes of Health (NIH). Each of these images contains eight subarrays arranged in \( 2 \times 4 \) format. In each subarray, the spots are arranged in a \( 29 \times 29 \) format. There are a total of 20184 spots in all the three NIH images. In addition to these, we also tested on another set of five test images obtained from Spectral Genomics Inc. (SGI). Each of the SGI images contains eight subarrays arranged in \( 12 \times 2 \) format, and in each subarray, the spots are arranged in a \( 16 \times 6 \) format. These five SGI images contain a total of 9960 spots. The target sites in the NIH images exhibit noticeable irregular hybridization effect and have irregular brightness patterns across the spots. The intensities of these target sites span over a large range and vary considerably. The target sites in the SGI images appear to be hybridized more homogeneously, and many of them have nearly perfect circular shape.

In the experiments, for each two-channel image, the summed bit rate of all the bitstreams from both channels, plus the shape information, were reported in bpp format, which represents either the compression bit rate or the reconstruction bit rate, depending on the type of test performed. And the corresponding bit rate of the uncompressed original image is 32 bpp. BASICA first segmented the image and generated the header file. The average overhead of the header file was 0.5 bpp for the NIH images and 0.24 bpp for the SGI images, based on the postprocessed segmentation results. The header file overheads were smaller on the SGI images because of different settings of the microarray arrays used to acquire the images: there were much fewer spots in each SGI image than those in each NIH image. After generating the header file, the foreground and background of each channel were compressed independently.

#### 3.1. Comparisons of wavelet filters and decomposition levels

The framework of the proposed compression scheme in BASICA does not specify which wavelet filters and how many wavelet decomposition levels used. In order to find the optimal choice for microarray image compression, we compare the results generated with different wavelet filters and decomposition levels. All the results presented in this section are based on the NIH images unless stated otherwise.

Table 2 lists the lossless coding results by BASICA using nine different wavelet filters with one-level wavelet decomposition. From these we found that the compression results vary only in a small range of about 0.07 bpp. Among all the nine sets of filters, the 5/3 wavelet filters achieved the best result. This is probably because the 5/3 wavelet filters have more advantage in representing details. As for the background compression, normal,
discrepancies in the results were small, the choice of the wavelet filters appeared to be not critical to the system performance.

Table 3 lists the lossless coding results by BASICA with different wavelet decomposition levels. Only the best-performing 5/3 wavelet filters were evaluated in these tests. The performance appeared to get worse when the decomposition level increased and compression with only one-level decomposition achieved the best result. This is partly due to the fact that although with more decompositions more data energy is compacted into smaller subbands, it also introduces a higher model-adaptation cost to arithmetic coding in the newly generated subbands, which cancels out the gains. Similar to the comparison among the wavelet filters, the discrepancies of lossless compression performance using different decomposition levels are very small. To confirm this observation, lossy compression tests were also performed to compare the performances based on the choices of the wavelet decomposition level.

To evaluate the effect of lossy compression on data analysis, the test images were first reconstructed at a target rate. Then the reconstructed images were processed and genetic information (i.e., log ratio) was extracted and compared with the same information extracted from the original images. To ensure credibility of the comparisons, the Mann-Whitney test-based segmentation started with the same selection of random pixels in the predefined background in both the reconstructed image and the original image. The segmentation was conducted under three different significance levels $\alpha = 0.001, 0.01,$ and $0.05$. At each significance level, log ratios were extracted and distortions were computed. The distortions shown are the average distortions at the three significance levels over the three test images. Both the $l_1$ distortion and $l_2$ distortion (i.e., MSE) of log intensity were used as the error measures. Figure 4 shows the average reconstruction errors using BASICA at different bit rates with three different decomposition levels of the 5/3 wavelet transform. From this figure, we can see that one-level decomposition yielded a significantly better performance than the others. Based on the above lossless and lossy compression results, we decided to use the 5/3 wavelet filters with one-level wavelet decomposition as a default setting in BASICA.

### Table 2: Lossless compression results (in bpp) of BASICA using different integer wavelet filters with one-level wavelet decomposition. The results are averaged over the NIH images.

<table>
<thead>
<tr>
<th>Wavelet filters</th>
<th>9/7 – F</th>
<th>(2 + 2, 2)</th>
<th>5/3</th>
<th>$S + P$</th>
<th>(4,2)</th>
<th>(2,4)</th>
<th>(4,4)</th>
<th>(6,2)</th>
<th>2/6</th>
</tr>
</thead>
</table>

### Table 3: Lossless compression results (in bpp) of BASICA using the 5/3 wavelet filters with different wavelet decomposition levels. The results are averaged over the NIH images.

<table>
<thead>
<tr>
<th>Decomposition levels</th>
<th>1 level</th>
<th>2 levels</th>
<th>3 levels</th>
<th>4 levels</th>
<th>5 levels</th>
</tr>
</thead>
<tbody>
<tr>
<td>File size</td>
<td>13.97</td>
<td>14.00</td>
<td>14.01</td>
<td>14.02</td>
<td>14.02</td>
</tr>
</tbody>
</table>

#### 3.2. Comparisons of lossless compression

We first compared the lossless compression performance of BASICA with three current standard coding schemes: TIFF, JPEG-LS, and JPEG2000. In the comparisons, TIFF, JPEG-LS, and JPEG2000 all compress a microarray image as a single region and no header file is added. To evaluate the improvement brought by the postprocessing in segmentation, along with the intensity and bit shifts in compression, we also performed the tests of BASICA without the intensity and bit shifts and without postprocessing, respectively (denoted by BASICA w/o PP and BASICA w/o shifts, respectively, in Figures 5, 6, and 7, and Table 4).

The coding results are shown in Table 4. The TIFF format, which is commonly used in existing microarray image archiving systems, produced the poorest results, about 4 bpp worse than all the other methods compared. JPEG-LS achieved the best performance on the NIH images. But like TIFF, it does not support lossy compression. The proposed BASICA turned out to be about 0.27 bpp worse than JPEG-LS on the NIH images and 0.12 bpp better on the SGI images. Besides, BASICA was significantly better than JPEG2000 with the savings of 0.48 bpp and 0.56 bpp on the NIH and SGI images, respectively. BASICA without intensity and bit shifts yielded almost the same performance as BASICA in lossless compression. On the other hand, one can see clearly that the irregularities in segmentation reduced compression efficiency substantially. Without postprocessing, the average size of a header file was 0.33 bpp larger than that of BASICA on the NIH images and 0.09 bpp larger on the SGI images, respectively. Thus, BASICA with postprocessing was preferred on all the test images.

#### 3.3. Comparisons of lossy compression

During the experiments, we also compared the lossy compression results at different bit rates. Since TIFF and JPEG-LS do not support the lossy compression functionality, JPEG2000 was the only standard compression scheme compared in the experiments. Our comparisons were based on three different measurements.

##### 3.3.1. Comparisons based on $l_1$ and $l_2$ distortions

We first compared the rate-distortion curves based on the $l_1$ and $l_2$ distortions of log intensity. Figure 5 shows the average reconstruction errors of these methods at different bit rates. We observe that, due to the effect of relatively more homogeneous hybridization, the distortion on the SGI images was uniformly smaller than the distortion on the NIH images. JPEG2000 produced surprisingly small $l_1$ distortion values at
low bit rates, only inferior to BASICA on the NIH images and similar to the others on the SGI images. Nevertheless, it produced relatively large $l_2$ distortion values. Apparently, without adjusting the MSE for log intensity, JPEG2000 spent too many bit rates on high-intensity pixels/spots, which led to high $l_2$ distortion. Furthermore, the distortion of JPEG2000 decayed slowly in both $l_1$ and $l_2$ senses. For bit rates beyond 6 bpp, it degraded to produce the worst distortion among all the methods. Without the intensity and bit shifts, BASICA performed poorly at lower bit rates. Only when the bit rates went above 6 bpp did its performance become acceptable. BASICA without postprocessing produced different performances on images of different sources. On the NIH images, it obviously suffered from the irregularities of segmentation, yielding a performance between BASICA and BASICA without the intensity, and bit shifts at low bit rates. But it quickly became worse than both of these schemes when the bit rates increased. On the SGI images, in which target sites had more uniform hybridization, there was almost no difference between its performance and BASICA’s. Compared to the other schemes, BASICA yielded the best performance in both $l_1$ and $l_2$ distortions at all the bit rates on all test images.

3.3.2. Comparisons based on scatter plots

Besides $l_1$ and $l_2$ distortion measures, a more intuitively visual way to compare the distortion of different methods is by scatter plotting. Figure 6 shows the extracted log ratios and log products by different methods at a bit rate around 4 bpp for two test images. In each scatter plot, the blue diagonal line corresponds to the information extracted from the original images. From the plots, we can see that BASICA had a better performance than the other methods. BASICA without postprocessing had a worse performance on the NIH images and a good performance on the SGI images. JPEG2000 and BASICA without intensity and bit shifts yielded worse performances on both sets of test images. This observation is consistent with the results shown in Figure 5. Since a scatter plot cannot provide quantitative performance measurements and can only visually display the data for comparisons at one bit rate per plot, it does not provide a practical performance measurement.

3.3.3. Comparisons based on gene expression data

Rather than judging the performance based on the $l_1$ and $l_2$ distortion measures and the scatter plots, biologists and clinicians in gene expression studies are likely to care more
Figure 5: Rate-distortion curves of log ratio in terms of $l_1$ distortion (left column) and $l_2$ distortion (right column) under different reconstruction bit rates for different compression schemes: (a-b) results based on the NIH images; (c-d) results based on the SGI images. The segmentation was performed at the significance level $\alpha = 0.05$.

whether a gene is differently detected or identified due to a lossy compression. Hence, it is meaningful to look at the rate of disagreement on detection and identification between lossily reconstructed image and original image. The detection and identification disagreement are defined as follows.

1. The detection disagreement is defined to be the valid spots in the original image being detected as false spot, or vice versa, after a lossy reconstruction.
2. The identification disagreement is defined to be a different classification outcome among up-regulated, down-
regulated, and invariant gene expression levels after a lossy reconstruction, even though the detection outcome is the same.

We conducted experiments using a simple quantitative model of gene expression data analysis to compare different methods. We determined that a spot was false if its foreground intensity was less than its background intensity in either channel or no foreground target site was found by the segmentation. We also decided that if the log ratio was larger or smaller than a certain threshold range \([\theta, -\theta]\), then the spot was up- or down-regulated; otherwise, it was invariant. For these experiments, no normalization was performed to reduce the interimage data variations. The experiments were performed on the NIH images and the SGI images separately and the results are shown in Figure 7. From this figure we can see that the identification disagreement rate was about 10 times higher than the detection disagreement rate. These results were similar to what have been shown in Figure 5. The disagreement caused by the lossy compression of JPEG2000 was comparable to that of BASICA only at 2 bpp, and dropped slowly when the bit rate increased. On the other hand, the disagreement caused by BASICA without intensity and bit shifts became acceptable only after 6 bpp. BASICA without postprocessing yielded a performance similar to that of BASICA on the SGI images but did worse on the NIH images. One can also observe that the disagreement rates on the NIH images were much higher than on the SGI images at the same bit rate. This is probably because NIH images are much noisier than SGI images, and hence require more bit rates to compress. These results are consistent with Figure 5, where the NIH images have much larger \(l_1\) and \(l_2\) distortions than the SGI images.
at the same bit rate. For the NIH images, the identification disagreement rate was larger than 10% at 2 bpp and was around 1.5% at 10 bpp. For the SGI images, the identification disagreement rate was smaller than 2.5% even at 2 bpp, and was around 0.1% at 10 bpp. All these results consistently suggested that one could hardly find a common bit rate that led to similar disagreement/agreement rates for different microarray images. For images with homogeneous
hybridization, which are becoming more available with the advance of microarray production technology, lossy compression at low bit rates appears to be viable for highly accurate gene expression data analysis.

4. CONCLUSIONS AND FUTURE RESEARCH

We have introduced a new integrated tool, microarray BASICA, for cDNA microarray data analysis. It integrates background adjustment, and image segmentation and compression in a coherent software system. The cDNA microarray images are segmented by a fast Mann-Whitney test-based algorithm. Postprocessing is performed to remove the segmentation irregularities. A highly efficient image coding scheme based on a modified EBCOT algorithm is presented, along with a new distortion measurement specially chosen for cDNA microarray data analysis. Experimental results show that cDNA microarray images of different quality require different bit rates to ensure sufficiently accurate gene expression data analysis. For homogeneously hybridized cDNA microarray images, BASICA is able to provide from a bit rate as low as 5 bpp the gene expression data that are 99% in agreement with those of the original 32 bpp images. Future research includes finding the optimal rate allocation between the background and foreground, and between the two channels of a cDNA microarray image.

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