Exploiting noise in array CGH data to improve detection of DNA copy number change

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ABSTRACT

Developing effective methods for analyzing array-CGH data to detect chromosomal aberrations is very important for the diagnosis of pathogenesis of cancer and other diseases. Current analysis methods, being largely based on smoothing and/or segmentation, are not quite capable of detecting both the aberration regions and the boundary break points very accurately. Furthermore, when evaluating the accuracy of an algorithm for analyzing array-CGH data, it is commonly assumed that noise in the data follows normal distribution. A fundamental question is whether noise in array-CGH is indeed Gaussian, and if not, can one exploit the characteristics of noise to develop novel analysis methods that are capable of detecting accurately the aberration regions as well as the boundary break points simultaneously? By analyzing bacterial artificial chromosomes (BACs) arrays with an average 1 Mb resolution, 19 k oligo arrays with the average probe spacing <100 kb and 385 k oligo arrays with the average probe spacing of about 6 kb, we show that when there are aberrations, noise in all three types of arrays is highly non-Gaussian and possesses long-range spatial correlations, and that such noise leads to worse performance of existing methods for detecting aberrations in array-CGH than the Gaussian noise case. We further develop a novel method, which has optimally exploited the character of the noise, and is capable of identifying both aberration regions as well as the boundary break points very accurately. Finally, we propose a new concept, posteriori signal-to-noise ratio (p–SNR), to assign certain confidence level to an aberration region and boundaries detected.

INTRODUCTION

Amplification or deletion of chromosomal segments can lead to abnormal mRNA transcript levels and result in the malfunctioning of cellular processes. Locating chromosomal aberrations in genomic DNA samples is an important step in understanding the pathogenesis of many diseases, most notably cancers. Microarray-based comparative genomic hybridization (array CGH) is a powerful technique for measuring such changes (1–5). To realize the promise of the array CGH technique, it is very important to develop effective methods to identify aberration regions from array CGH data. Existing analysis methods (6–23) can be roughly classified into two categories: smoothing-based (6–9) and segmentation-based (10–19). The latter approaches explicitly model the observed array data as a series of segments, with unknown boundaries and unknown heights estimated from the data by employing certain optimization criterion. While the boundary points thus identified are reliable, the aberration regions identified may be less so, in the sense that some of them may be false positives. Smoothing-based approaches assume that true signals in a specific region, aberration or non-aberration, are smoother than any kind of noise superimposed on the signals. Therefore, they attempt to reduce noise by comparing individual data points to their adjacent ones and modifying them. While such methods can reduce the number of false aberration regions identified, the boundary points detected are usually less accurate than segmentation-based methods. It would be very desirable to develop new methods for analyzing array CGH data, with both the merits of smoothing and segmentation-based approaches. Such a goal may not be fully accomplished by just incorporating mean or median smoothing to a segmentation-based method.

The term ‘noise’ is often used in biology to describe experimental measurement imprecision. In particular, when evaluating the accuracy of an algorithm for detecting aberrations, it is commonly assumed that noise in the data follows normal distribution. However, this important assumption has not been verified/falsified based on the analysis of
CHARACTERIZATION OF NOISE IN ARRAY CGH

In this section, we carry out distributional analysis as well as spatial correlation analysis of array-CGH noise, and assess the effect of such noise on the performance of published algorithms for detecting aberrations from array-CGH data. Below, we first describe data.

Data

In this paper, we analyze data of three resolutions, BAC array (2), 19 k oligo array (24) and 385 k oligo array (http://www.nimblegen.com/products/cgh/). The BAC array (2) has an average 1 Mb resolution. It is from Stanford University, which can be freely downloaded from http://www.nature.com/ng/journal/v29/n3/suppinfo/ng754_S1.html. It consisted of 15 human cell lines with known karyotypes (12 fibroblast cell lines, 2 chorionic villus cell lines and 1 lymphoblast cell line) from the NIGMS Human Genetics Cell Repository. Each cell line had been hybridized with an array CGH of 2276 BACs, spotted in triplicate. The variable used for analysis was the normalized average of the log base 2 test over reference ratio, as processed by the original authors. In each cell line, there were either one or two aberrations. Among the 15 cell lines, 6 had aberrations that covered an entire chromosome. Note that some of these datasets were recently used by Olshen and Vankatraman (12) to evaluate the effectiveness of their algorithm for detecting aberrations from the data. For convenience, the names of the 15 cell lines are listed in the first column of Table 1.

The 19 k oligo array data (24) are from Harvard Medical School. It has an average probe spacing of <100 kb. The complete OligoLibrary consists of 18 861 oligos representing 18 664 LEADSTM clusters and 197 positive controls. There are four datasets for lymphoma tumors that developed in ATM deficient mice (24).

The 385 k oligo array data http://www.nimblegen.com/products/cgh/ has a median probe spacing of 6 000 bp through genic and intergenic regions. There are two datasets for the 385 k oligo array data. One is the normal female versus male case, where polymorphisms are observed by our method (to be described later) in chromosomes 1, 4 and 5. Another is the tumor case, where chromosome 8 has the longest aberration length (~2000), while chromosomes 10 and 19–22 do not have detectable aberrations at all (see Figure 1). Note that if we downsample the data to a resolution comparable to the BAC array data (2), then the longest aberration region in chromosome 8 only has less than 20 points. Therefore, the 385 k oligo array data http://www.nimblegen.com/products/cgh/ has the smallest aberration regions.

**Table 1.** \( p - SNR \) and Hurst parameter for noise of the 15 BAC array data (2)

| Cell line/chromosome(s) | \( p - SNR \) | \( H \) |
|-------------------------|--------------|---|
| GM01750/8/14            | 3.555/5.905  | 0.743 |
| GM01524/6               | 5.308        | 0.739 |
| GM01535/5/12            | 4.032/12.170 | 0.688 |
| GM03134/8               | 2.829        | 0.619 |
| GM03563/3/9             | 3.910/11.019 | 0.715 |
| GM05296/10/11           | 5.793/3.741  | 0.666 |
| GM07081/7               | 3.193        | 0.664 |
| GM13031/17              | 5.172        | 0.667 |
| GM13330/1/4             | 3.982/7.947  | 0.663 |
| GM00143/18              | 4.429        | 0.741 |
| GM02948/13              | 3.960        | 0.718 |
| GM03576/2/21            | 5.080/5.473  | 0.770 |
| GM04435/16/21           | 4.337/4.266  | 0.707 |
| GM07408/20              | 5.774        | 0.712 |
| GM10315/22              | 5.277        | 0.728 |

When there are two aberration regions, \( p - SNR \) is defined for both regions.

**Distributional analysis of array CGH noise**

When carrying out distributional analysis, an important issue to consider is the size of the sample points. For the 385 k oligo array data http://www.nimblegen.com/products/cgh/, we have considered noise for each chromosome in two scenarios, corresponding to that aberration and non-aberration regions (i) are considered together and (ii) are considered separately. The results are similar for both scenarios. For the BAC array data (2) and the 19 k oligo array data (24), we have also considered two cases, (i) each chromosome is considered separately and (ii) all the chromosomes are combined together. While the distribution for all the chromosomes combined is not the same as that for individual chromosomes, the qualitative feature of deviation from Gaussian distributions is similar for both cases. Comparisons of these different scenarios suggest that the non-Gaussian distributions discussed below may not be due to summation of multiple Gaussian distributions with different variance, corresponding to the euploid and copy-number-variant parts of the chromosome. In the main text here, we present results for the first scenario for all three types of data. Readers interested in knowing more details about the second scenario are referred to Supplementary Figures 1–3.
To simplify analysis, we have simply formed histograms. This is justified by noticing that the number of sample points is large in all the datasets. The distributions for the noise in two of the 15 BAC array data (cell line GM05296 and GM04435) are shown in Figure 2a and b. We observe that the distribution deviates from the Gaussian distribution considerably. In fact, these are typical results for the BAC array data (2). The distribution of noise in the 19 k oligo array data (24) is even more non-Gaussian, as shown in Figure 2c and d, which are typical among the four datasets. Since the 19 k oligo array data (24) has wider aberration regions, we suspect that the deviation from Gaussian distribution is positively correlated with the length of aberration regions. This hypothesis seems to be supported by analysis of the 385 k oligo array data http://www.nimblegen.com/products/cgh/. In Figure 2e–h we have shown the distributions for noise in chromosomes 8 and 9 of both the normal and tumor cases, where Figure 2e and g are for the normal case, while Figure 2f and h are for the tumor case. Note that distributions for noise in other chromosomes are very similar to those shown in Figure 2e–h. We observe that the distribution in Figure 2e and g is very close to Gaussian, while the distribution in Figure 2f and h is non-Gaussian, but the deviation from Gaussian is less severe than that shown in Figure 2a–d. Note that there are no aberrations in the normal case, while the aberration regions in the tumor case of the 385 k oligo array data (http://www.nimblegen.com/products/cgh/) are smaller than those in the BAC array data (2).

Spatial correlations in array CGH noise

Denote array CGH noise by \(x_1, x_2, \ldots, x_t\) and the spatial resolution by \(\Delta x\). We have found that array CGH noise can be characterized as a type of fractal noise characterized by an algebraically decaying spatial autocorrelation function,

\[\gamma(m) = \frac{E(x_i x_{i+m})}{E(x_i^2)} \sim m^{2H-2},\]

where \(m\) corresponds to physical spacing \(m\Delta x\), \(0 < H < 1\) is the Hurst parameter (25). In particular, when \(1/2 < H < 1\), the summation of the autocorrelation becomes unbounded if \(m\) can go to infinity. Therefore, such noise process is often said to have long-range persistent correlations. We shall discuss its implications to DNA copy number change momentarily. When \(H = 1/2\), the noise is like the white Gaussian noise.
There exist many different ways to estimate the Hurst parameter \( H \) (25,26,27). For ease of interpretation, we choose variance-spacing relation. To use this method, one can analyze non-overlapping running means of the original array data by constructing a new time series \( X^{(m)} = \{X_t^{(m)}\}, t = 1, 2, 3, \ldots, m = 1, 2, 3, \ldots \)

\[
X_t^{(m)} = (x_{t-m} + \cdots + x_t)/m, \quad t \geq 1.
\]

For a noise process with the property described by Equation (1), the variance of the running means, \( \sigma^2 \), declines in a power-law manner as the size of the sample, \( m \), increases:

\[
\text{var}(X^{(m)}) = \sigma^2 m^{2H-2},
\]

where \( \sigma^2 \) is the variance of the original time series \( x_t \). Based on Equation (2), one can readily understand the meaning of \( H \) in terms of how effective smoothing can reduce variations in the noise. For example, when \( H = 0.5 \), \( \text{var}(X^{(m)}) \) drops to \( \sigma^2/10 \) when \( m = 10 \). However, if \( H \) becomes 0.75, then for the variance to drop to \( \sigma^2/10 \), one needs to take \( m = 100 \). This is an order of magnitude larger than if \( H = 0.5 \). In other words, smoothing is less effective in reducing variations in the data. For notational convenience, we shall re-write Equation (2) as

\[
F(m) = m [\text{var}(X^{(m)})]^{1/2} = \sigma m^H.
\]

We examine the long-range spatial correlations in the 385 k oligo array data (http://www.nimblegen.com/products/cgh/) first. Since the datasets have very high-spatial resolution, in each chromosome, we have plenty of data points. We carry out variance-spacing relation analysis of the data in each chromosome separately. When aberration regions exist in the data, we pre-process the data using two methods. One is to discard the part of data corresponding to the aberration regions. Another is to retain the part of data corresponding to the aberration regions, with the mean of that part removed. It is clear that after processing by either method, the remaining data is the fluctuations or noise in the array probes. The characteristics of fluctuations by both methods are similar. Below, we present results for the first method. Figure 3 shows \( \log_2 F(m) \) versus \( \log_2 m \) for the data of the six chromosomes of the 385 k oligo array data, where the red diamond is for the tumor case, while the black circle is for the normal case. The solid black and dashed red lines are straight lines fitted by linear least squares regression. There are no aberrations in chromosomes 19 and 20. The Hurst parameters are obtained as the slopes of the straight lines, which are indicated in the figure.

Figure 3. \( \log_2 F(m) \) versus \( \log_2 m \) for the data of the 6 chromosomes of the 385 k oligo array datasets. The red diamond is for the tumor case, while the black circle is for the normal case. The solid black and dashed red lines are straight lines fitted by linear least squares regression. There are no aberrations in chromosomes 19 and 20. The Hurst parameters are obtained as the slopes of the straight lines, which are indicated in the figure.
(24) are in the range of 0.6–0.77 (H for the BAC array data (2) are listed in the 3rd column of Table 1). While they are all larger than 0.5, suggesting long-range spatial correlations, we have to emphasize that such long-range spatial correlations are the correlations across chromosomes, and therefore, are different than the correlations in the 385 k oligo array data (http://www.nimblegen.com/products/cgh/), which are within chromosomes. We believe this is the primary reason that the Hurst parameters for the BAC array data (2) and the 19 k oligo array data (24) are smaller than those of the 385 k oligo array data (http://www.nimblegen.com/products/cgh/) with tumors. While the long-range spatial correlations in the BAC array data (2) and the 19 k oligo array data (24) might not have much biological relevance, they are important features to consider when one designs methods to detect DNA copy number changes from them.

Effect of array noise on detection of aberrations

To illustrate the effect of array noise on detection of aberrations, we choose CGHseg algorithm (15), which is one of the best segmentation based algorithms, and consider simulated data of various aberration widths (5, 10, 20 and 40 probes) and noise levels (SNR of 1, 2, 3 and 4). SNR is defined as the mean magnitude of the aberration (i.e. signal) divided by the SD of the superimposed noise. Two types of noise are considered. One is simulated Gaussian noise. The other is the actual noise in the BAC array data (2). For each aberration width and SNR, we generate 100 artificial chromosomes, each consisting of 100 probes and with the square-wave signal profile added to the center of the chromosome. To generate receiver operating characteristic (ROC) curve corresponding to a particular aberration width and SNR, we calculate the true positive rates (TPR) and the false positive rates (FPR). TPR is defined as the number of probes inside the aberration whose fitted values are above the threshold level divided by the number of probes in the aberration. FPR is defined as the number of probes outside the aberration whose fitted values are above the threshold level divided by the total number of probes outside the aberration. We vary the threshold value for aberration from the minimum log-ratio value to the maximum. Each threshold value results in a TPR and a FPR, represented by a point on the ROC curve. Similar simulation procedure has been used by Lai et al. (23). Figure 4 shows the ROC curves corresponding to different aberration widths and SNRs, where the purple curves correspond to the simulation with real array noise, while the green curves correspond to the simulation with Gaussian noise. We observe that the green curves are generally above the purple ones. This is especially so when SNR = 1 and the aberration width is 20. Therefore, we can conclude that performance of CGHseg algorithm for detecting aberrations from array CGH data are worse when real array noise is used than when Gaussian noise is used. Interestingly, other existing methods behave similarly.

Summary

Summarizing the results discussed so far, we can conclude that when there are aberrations, noise in array CGH is highly non-Gaussian and possesses long-range spatial correlations. It appears that the non-Gaussian feature as well as the long-range spatial correlation feature become stronger when the aberration regions become larger. When SNR is low and the aberration width is large, performance of existing methods for detecting aberrations is worse with this type of noise than when noise is assumed to be Gaussian.
EXPLOITING NOISE TO IMPROVE DETECTION OF ABERRATIONS FROM ARRAY CGH DATA

We now present an algorithm for detecting aberrations from array CGH data that has considerably taken into account the character of array noise. Since the method has merits of both smoothing and segmentation based methods, we denote it by CGHss. It consists of five steps. They are detailed below.

1. To reduce noise, the original log_2 ratio data \( y(n) \), \( n = 1, 2, \ldots, N \) is filtered by a median filter. In order not to lose too much information about the boundary, the length of the filter is 3-point. Let us denote the resulting data by \( y_1(n) \).
2. Construct a random walk process from \( y_1(n) \) by simply forming partial summation of \( y_1(n) \) based on the following formula,
   \[
   y_0^1(n) = \sum_{i=1}^{n} [y_1(i) - \mu],
   \]
   where \( \mu \) is the mean of \( y_1(n) \); then \( y_0^1(n) \) is partitioned into overlapping segments of length 3 and overlap 2, and the local trend in each segment is calculated to be the ordinate of a linear least-squares fit for the random walk in that segment. Denote the difference between the original walk and the local trend by \( y_2(n) \).
   Note this step plays the role of smoothing. In particular, as will be shown in Figure 5c, the pattern of \( y_0^1(n) \) can be well-associated with the aberration regions. We emphasize that unlike conventional lossy smoothing, here, information on original data can be recovered. Hence, it is a lossless smoothing. Furthermore, \( y_2(n) \) can be used to estimate the Hurst parameter through the method called detrended fluctuation analysis (28). Therefore, this is the step that has fully taken into account the spatial correlation feature of the data.
3. Let \( \text{var}(a,b) \) denote the variance of two variables \( a \) and \( b \), which is simply \( (a - b)^2/2 \). Now we modify \( y_1(n) \)

Figure 5. A schematic figure illustrating the proposed method. (a) Shows the original array data, (b-g) Show the signals \( y_1(n), y'_1(n), y_2(n), y'_2(n), y_3(n) \) and \( y_4(n) \), respectively. The dashed red lines in (f) denote the threshold \( T_1 \).
according to the following rule:

\[
\begin{align*}
  y^*_1(n) &= y_1(n + 1), & \text{if } \var{y_1(n), y_1(n - 1)} \\
  y^*_1(n) &= y_1(n - 1), & \text{otherwise}
\end{align*}
\]

Note this step does both segmentation and smoothing.

(4) Let

\[
y_3(n) = y^*_1(n) \times y_2(n),
\]

and define

\[
y_4(n) = \begin{cases} y^*_1(n), & \text{if } |y_3(n)| > T_1, \\
  y_3(n), & \text{otherwise}
\end{cases}
\]

where \(T_1\) is a threshold value. This step yields a square wave-like signal. With this signal, we can make simple decisions, which is our step (5).

(5) Set thresholds \(T_2\) (a positive number) and \(T_3\) (a negative number), and identify the regions in \(y_4(n)\) data greater than \(T_2\) or less than \(T_3\) as potential amplifications or deletions in array CGH data. Sometimes in order to reduce false positives, we may discard small regions with only one or two probes. However this should be done with caution, since some microdeletions may only contain a single probe.

To make the above steps concrete, we have simulated an artificial chromosome data consisting of 100 probes, with the centering 10 probes having aberrations. The simulated data \(y(n)\) is shown in Figure 5a. Figure 5b–g show the signals \(y_1(n), y_1^*(n), y_2(n), y_3(n)\) and \(y_4(n)\), respectively. The red dashed lines in Figure 5f correspond to a more or less arbitrarily chosen threshold value \(T_1\).

Note all the three thresholds, \(T_1, T_2\) and \(T_3\), can be defined by users. Also note that the ROC curves presented below do not depend on \(T_1\) sensitively. After we describe the concept of \(p - \text{SNR}\), we shall provide some guidelines as how to choose \(T_2\) and \(T_3\).

We now compute the ROC curves for our method under the same setting as when we discussed the CGHseg algorithm (Figure 4, green and purple curves). They are shown in Figure 4 as red and black curves, for array noise and Gaussian noise, respectively. First we note that comparing with the results of a recent comparison paper (23), our method is comparable to the best smoothing based methods. Next, we make two interesting observations from Figure 4: (i) The ROC curves for our method are very similar for the array noise and the Gaussian noise. This is because our method has fully taken into account the character of the array noise and the Gaussian noise. This is because our method does not seem to be able to cope with such low SNR data. This is evident in Figure 6c and d (as well as Supplementary Figures 4–7): The CGHseg algorithm, however, does not seem to be able to cope with such low SNR data. 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com/products/cgh/ are around 0.9 to 1.5, and are even smaller for the 19 k oligo array data (24) (around 0.7–1.1). In particular, we have estimated the \( p/C_0 \) SNR for the 385 k and 19 k oligo array data based on the X chromosome using sex-mismatched samples. We have found that the \( p/C_0 \) SNR for the two array platforms are 1.26 and 1.03, respectively, both falling in the range for each type of data listed above. Although we do not have access to the sex-mismatched X chromosome data for the BAC array data (2), based on our analysis of the other two types of data, we have good reason to believe that the \( p/C_0 \) SNR for the BAC array data’s sex-mismatched X chromosome would be at least around 3, therefore, much larger than the \( p/C_0 \) SNR of the other two types of data.

Our concept of \( p/C_0 \) SNR suggests that a good starting point to choose the parameters \( T_2 \) and \( T_3 \) used in step (5) of our algorithm may correspond to signal intensity divided by \( p/C_0 \) SNR. This amounts to choosing one SD of the data. This rule suggests an iterative operation: starting from arbitrarily chosen \( T_2 \) and \( T_3 \), calculate the corresponding \( p/C_0 \) SNR, then use the criterion discussed above to obtain a new estimate of \( T_2 \) and \( T_3 \), finally calculate the new \( p/C_0 \) SNR. If \( p/C_0 \) SNR and \( p/C_0 \) SNR’ are similar, then the two parameters have been chosen appropriately.

We emphasize that our method works excellently if \( p/C_0 \) SNR is high. However, if \( p/C_0 \) SNR is low, then one may choose threshold values that roughly yield TPR + FPR = 1, where TPR and FPR define the ROC curve. In this case, however, one should bear in mind that the classification may be incorrect with a probability of FPR.

DISCUSSION

In this paper, we have examined noise in array CGH data of three resolution, the BAC array data, the 19 k and 385 k oligo array data, and found that noise is highly non-Gaussian and possesses long-range spatial correlations. We have also developed a novel method for processing array CGH data. The method is a suitable combination of smoothing and segmentation, and has fully taken into account the characteristics of noise in array CGH data. We have shown that the method is as accurate as the best smoothing-based methods for detecting aberration regions, and as accurate as the best segmentation-based methods for finding boundary points. Furthermore, we have proposed a new concept, \( p/C_0 \) SNR, to quantify the confidence level of aberration regions and boundaries detected. We have found that \( p/C_0 \) SNR for the 15 publicly available BAC array CGH data are all quite large, indicating it is a relatively easy matter to accurately detect aberrations and boundaries from those data. However, \( p/C_0 \) SNR for the four 19k oligo array data are quite small, suggesting it is considerably more challenging to detect aberrations from such array CGH data.

Although we have found that array CGH noise is highly non-Gaussian with long-range spatial correlations, we do...
not clearly know the mechanisms. A challenging task for future research would be to understand the biological mechanisms of such noise, as well as understand whether those mechanisms are differentially related to different types of diseases.

Being able to identify smaller copy-number changes that affect only a few probes is of particular importance in the field of copy number polymorphism. This is because inherited, germ-line copy number variants are typically much smaller than rearrangements in cancer genomes. For example, two recent papers, one by McCarroll et al. (29), another by Conrad et al. (30), identified a large class of inherited, multi-kilobase deletion polymorphisms that are predominantly smaller than 20 kb in size. We emphasize that in order to detect small copy number changes, the key is to improve the resolution of the array technology so that at least 2 points can be sampled for the region of interest. If only a single isolated point can be sampled, then it would be impossible by any analysis method to classify it as a true copy number change or just an outlier or noise.

Finally, readers interested in this method are strongly encouraged to contact with the authors (e.g. gao@ece.ufl.edu) to obtain the source Matlab code.

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