Estimating robustness of the tileShuffle method with repeated probes

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Abstract

Motivation:
In this paper the TileShuffle method is evaluated as a search method for candidate IncRNAs at 8q24.2. The method is run on three microarrays. Microarrays which all contained the same sample and repeated copies of tiled probes. This allows the coherence of the selection method within and between microarrays to be estimated by Monte Carlo simulations on the repeated probes.

Results: The results show poor consistency in areas selected between arrays containing identical samples. A crude application of the method can result in majority of the region to be selected, resulting in a need for further restrictions on the selection. Restrictions based on ranking internal tileShuffle test statistics do not increase precision.

As the tileShuffle method has been shown to have higher precision than the MAS and TAS software, one can conclude that methods giving unreliable results are in common use.

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1 Introduction

Gene expression is the process by which information from a gene is used in the synthesis of a functional gene product. These products are often proteins, but there are also non-protein coding genes where the product is a functional RNA. It has been predicted that more than 30,000 RNA genes are associated with the human genome (Kapranov and Laurent 2012).

Non-protein coding genes and their products can vary considerably in length. The shortest products, micro RNAs (miRNA), are on average only 22 bp, whereas long non-coding RNA (lincRNAs) are defined as transcribed RNA molecules longer than 200 nucleotides in length (Sana et al. 2012).

There have been several publications indicating that lincRNAs might play an important role in cancer development (Wang et al. 2011; Huarte and Rinn 2010; Tsai et al. 2011; Gibb et al. 2011b; Shore et al. 2012; Willard and Koochekpour 2012) and a good review of their functional role in human carcinomas is given in Gibb et al. (2011a).

lincRNAs are also thought to play a regulatory role in cancer-associated pathways governing mechanisms such as cell growth, invasion, and metastasis and have been seen to be expressed differently in primary and metastatic cancer (Tahira et al. 2011). lincRNAs might thus provide insights into the mechanisms underlying tumor development.

lincRNAs originate everywhere in the genome, but especially in long stretches where no protein-coding genes have been identified (Chung 2011). An example of such an area is 8q24, where multiple single nucleotide polymorphisms (SNPs) have been associated with risk of developing prostate cancer (Haiman et al. 2007; Yeager et al. 2007; Gudmundsson et al. 2007; Amundadottir et al. 2006). Currently, there are at least 11 databases which record lincRNAs (Dinger et al.)
Microarrays are frequently used to locate RNA genes. A microarray contains multiple copies of the same DNA oligonucleotides, known as probes, which are hybridized to a labeled RNA sample and the array is subsequently washed. Theoretically this will result in the labeled sample only remaining where the sample hybridized to probes. The signal intensities at the corresponding location on the microarray are used as a measure of the relative abundance of hybridization of each probe.

Typically a probe corresponds to a specific genomic region. Sometimes the probes overlap, referred to as tiling, and such arrays are called tiled microarrays. Tiled microarrays have been successful in assessing expression of non-coding RNAs. (Johnson et al. 2005; Mockler and Ecker 2005; Weile et al. 2007)

The ability to accurately detect the true gene-expression signal in microarrays is affected by several sources of variation (Pozhitkov et al. 2007; Churchill 2002). Further issues and different biases arise when using tiled microarrays, as opposed to other analysis of differential expression (Royce et al. 2005). It is therefore important to take technical variation into account when doing statistical analysis on microarray data (Wu et al. 2004; Royce et al. 2005).

Currently a variety of methods are available to analyse data from tiled microarrays, but as expression levels are generally lower for IncRNA than protein coding genes, (Gibb et al. 2011a) conventional methods for differential expression detection may have difficulties detecting them. A good overview of available methods is found in Otto et al. (Otto et al. 2012), where the tileShuffle method is introduced and shown to have higher precision than the commonly used TAS (Kampa et al. 2004) and MAT (Johnson et al. 2006) methods.

The tileShuffle method identifies transcribed segments in terms of significant differences from the background distribution, using a permutation test statistic, called a window score. All probes within a sliding window have a window score assigned (arithmetic mean trimmed by median or max and min value). Further, probes are subdivided into bins by GC content and processed independently.
The significance of a window score is assessed by permuting probes across the array, but always within the same bin. Empirical p-values are estimated by counting the number of permuted windows with higher score (Otto et al. 2012).

The aim of this study is to assess the robustness of the tileShuffle method on the expressed regions level. It utilizes a special array-design where every probe is repeated ten times on each tiled array. This enables Monte-Carlo simulations of expression signals which are used to estimate expression on pseudo-arrays, whose differences lie in a variability that is usually neglected in microarray experiments. Further, a single biological sample was split in three and used on repeated arrays, providing estimates of another variability that is commonly neglected. The consistency in regions selected across the pseudo-arrays, that should ”in theory” give identical results, will be used as a measure of the robustness of the tileShuffle method.

2 Methods

The data discussed in this publication are RNA expression data from custom designed Nimblegen microarray experiments where the same prostate tissue sample was used on three arrays. The data have been deposited in NCBI’s Gene Expression Omnibus (Edgar et al. 2002) and are accessible through GEO Series accession number GSE45934.

The arrays contained 50 nucleotide probes from chr8:127640000-129120000 at locus 8q24, tiled at a 20 base interval. The whole region was tiled evenly, but probes with blat score greater than 5 (Kent 2002) or blast score greater than 40 (Altschul et al. 1990) were excluded. That left in total 54236 (out of 74000) probes, each of which was replicated 10 times on the array.

Spatial artifacts in the expression signal were minimized by aggregating the wells of the microarray into ten non-overlapping logical virtual ”containers”, allocating each of the ten replicates of a probe to a different container.

These ten replicate spots for each probe, evenly spread across the array,
permitted Monte-Carlo simulation of the expression signals. In that way, for each of the three microarrays, 1000 pseudo replicate arrays were produced, with only one repetition of each probe, selected at random. The pseudo-arrays were made in triplets, such that the same set of replicates was used to produce pseudo-arrays for all three microarrays within every simulation.

The arrays were normalized by the quantile normalization method (Bolstad et al. 2003) and consequently analysed with the tileShuffle method under various conditions, detailed later, but always one at a time. The window size, the minimum length of selected areas, was set as 1000 bases, as the aim was to detect relatively long areas. Three GC-content bins were used and the number of permutations was 1000. All statistical analyses were performed in the R statistical package (R Development Core Team 2011) and graphics are generated with the ggplot2 library (Wickham 2009).

The tileShuffle method assesses significance on minimal expected transcriptional units rather than on a single probe level. (Otto et al. 2012) Therefore the tiled region was split up into areas of length 100 bases. These areas will be underlying when addressing genomic locations that are expressed. An area will be deemed as expressed if all corresponding 100 bases were within an expressed region.

3 Results

3.1 Consistency within an array

As each probe was repeated ten times on every array, the consistency of the method could be estimated by Monte-Carlo simulations on the probe sets. In that way, 1000 pseudo-arrays were produced, each by randomly selecting one repetition of each probe. All experimental sources of variation of these pseudo-arrays are identical, except the physical location of the probesets within the microarray.

In order to investigate the effect of this probe-to-probe variation on calls of expression, the tileShuffle method was run on each of the 1000 pseudo-arrays.
Figure 1: A smoothed plot of on how many of the 1000 pseudo-arrays each genomic location was called expressed by the tileShuffle method. This is shown for the first 300,000 bases of the tiled area.

Figure 1 shows on how many of the pseudo-arrays each genomic location was ”called” ”expressed” (p <0.05, adjusted for multiplicity). For clarity, the figure shows only the first 300,000 bases of the tiled area or about one-fifth of the tiled region. Graphs for the remaining regions were similar. The figure shows that a great majority of the underlying tiled region is selected on at most 25% of the pseudo-arrays, whereas a few areas are selected consistently in near all cases.

### 3.2 Consistency between arrays

To compare across arrays, pseudo arrays were simulated in triplicates so that a single set containing probes from the same physical location on the microarray was generated for all three arrays at a time. Thus, within each triplicate of pseudo-arrays, all probes have the same internal physical location of probes on the original microarrays, blocking the location effect of probes within a microarray. This emulates real situations where probes are not replicated.

The performance of different methods for selecting subsets of the areas deemed expressed is compared in Table 1. The three columns show the areas selected on exactly one, two or all three pseudo-arrays within a triplicate, as a proportion of areas that are selected on some pseudo-array within the triplicate.
Table 1: The average proportion of areas that are selected on exactly one, two or all three pseudo-arrays within a triplicate of all areas that are selected on some triplicate. From top-down: 1) Only the 30 areas with the highest window score are selected in every simulation, 2) all the probeset, one replication of each probe, 3) half of the probeset was used with two replicates of each probe, 4) the median score over every 10 probes was calculated a priori and fed to the method, 5) only select areas that are deemed expressed in at least 99% of the replications 6) all 10 replicates were fed to the method.

| Condition                                                                 | One array | Two arrays | Three arrays |
|---------------------------------------------------------------------------|-----------|------------|--------------|
| thirty highest window scores                                             | 0.58      | 0.19       | 0.22         |
| one replicate of all probes                                              | 0.49      | 0.23       | 0.28         |
| two replicates of half of the probes                                     | 0.47      | 0.23       | 0.30         |
| median of every ten probes                                               | 0.46      | 0.23       | 0.31         |
| selected in at least 99% of simulations                                  | 0.52      | 0.16       | 0.32         |
| all ten replicates                                                       | 0.23      | 0.14       | 0.63         |

Ideally one would like to maximize the proportion of instances where a location is expressed on either none, or all of the three arrays. The results are shown by increasing proportion of areas that are selected on all three pseudo-arrays.

Finally, Supplementary Fig. 1 shows the relationship between the average proportion of the total underlying genomic area that is chosen in each simulations against the number of replicates used in every simulation. The relationship is shown for the proportion that is selected on at least one array, at least two arrays and all three arrays and a number of replicates running from 1 up to 10 replicates of each probe per array.

4 Discussion

This paper is based on an experimental setup using three tiled microarrays containing the same biological sample, each using ten repetitions of each probe. Monte-Carlo simulations from real data are used to investigate the robustness of the tileShuffle method when targeting areas on locus 8q24 that
are expressed in prostate cancer.

This study raises several concerns regarding the consistency of areas selected. First of all the method shows considerable variability depending on which of the 10 replicates of each probe the method is applied to. Ideally, every area on Figure 1 should be expressed in either all or none of the Monte-Carlo simulations, resulting in the proportion being close to 0 or 1. As shown on the figure, these proportions span the whole spectrum from zero to one. Most probes which are ”called” are only called in fewer than 25% of all simulations, indicating a serious lack of repeatability. A few areas are selected consistently on nearly all pseudo-arrays, but as probes are not repeated in the common situation, this plot is not available and one cannot identify locations that are consistently expressed across pseudo-arrays.

Table 1 shows poor between-array consistency in choice of areas, which should ideally be identical. Probes which show consistency across repetitions within an array (selected in at least 99% of simulations) do not show more consistency across arrays.

The difference in results by applying the method on all ten replicates v.s. first calculating the median of every ten probes and then applying the method is somewhat counter-intuitive. A few further points should also be noted: Selecting a fixed number of areas with the highest window score is an unrobust method. The difference in using repeated probes rather then denser tiling is small, although in favour of repeated probes.

Finally, as shown on Supplementary Fig. 1, the average proportion of the underlying genomic region that is selected increases rapidly as the number of replicates of each probe increases. With ten replicates of each probe, almost 60% of the underlying region is selected on at least one array out of three. This might suggest that majority of the underlying region is ”expressed” by definition and the lack of consistency is in caused by small power.
5 Conclusion

This paper shows poor consistency of the tileShuffle method both between selection of replicates of probes within a microarray and also between microarrays containing the same sample. As the tileShuffle method has shown to have higher precision than the MAS and TAS software, one can conclude that methods giving unreliable results are in common use.

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