Genome analysis

Advance Access publication August 5, 2010

A framework for oligonucleotide microarray preprocessing

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Associate Editor: John Quackenbush

ABSTRACT

Motivation: The availability of flexible open source software for the analysis of gene expression raw level data has greatly facilitated the development of widely used preprocessing methods for these technologies. However, the expansion of microarray applications has exposed the limitation of existing tools.

Results: We developed the oligo package to provide a more general solution that supports a wide range of applications. The package is based on the BioConductor principles of transparency, reproducibility and efficiency of development. It extends the existing tools and leverages existing code for visualization, accessing data and widely used preprocessing routines. The oligo package implements a unified paradigm for preprocessing data and interfaces with other BioConductor tools for downstream analysis. Our infrastructure is general and can be used by other BioConductor packages.

Availability: The oligo package is freely available through BioConductor, http://www.bioconductor.org.

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Supplementary information: Supplementary data are available at Bioinformatics online.

Received on March 21, 2010; revised on July 13, 2010; accepted on July 21, 2010

1 INTRODUCTION

Open source software significantly simplified the development and distribution of preprocessing methods for gene expression microarrays. The BioConductor project (Gentleman et al., 2004) is one hub of tools for the analysis of genomic data and distributes, among others, the affy package (Gautier et al., 2004), the most used tool for analysis of Affymetrix gene expression arrays. In addition to Affymetrix, other manufacturers (e.g. Illumina, NimbleGen and Agilent) also commercialize microarray solutions, increasing the number of applications of the technology.

With different microarray applications, the investigator can analyze genomic data from different perspectives: Gitan et al. (2002) use tiling arrays to identify, at high resolution, regions of DNA and histone modifications; The International HapMap Consortium (2003) uses genome-wide single nucleotide polymorphism (SNP) and copy number variant (CNV) arrays to obtain: (i) genotype calls, later used in association studies; and (ii) extract copy number estimates to assess chromosomal aberrations; Clark et al. (2007) use exon arrays to analyze alternative splicing. As new applications

became available and designs by other manufacturers became more popular (see Supplementary Material), users developed suboptimal solutions to allow the use of the existing code in affy on these new arrays. This strategy did not succeed because the new designs did not share the structure used by the Affymetrix gene expression arrays, such as density and array annotation standards.

Based on the BioConductor principles of transparency, reproducibility and efficiency of development, we developed the oligo package. Its infrastructure, presented in Section 2, is general and can be used by other BioConductor packages. The package natively supports feature-level data for different applications and manufacturers, as shown in Section 3. It implements a unified framework for preprocessing microarray data and interfaces with other BioConductor tools for downstream analysis.

2 INFRASTRUCTURE

The package contains structures to simplify usage and interaction with other packages. A clear distinction is made between featurelevel, summarized and annotation data, and this is reflected by the different classes that are implemented using the S4 scheme intrinsic to the R environment.

Before importing data, the researcher must have the respective annotation package already installed in the system. The annotation package provides array coordinates, feature types, sequences, feature names and other relevant information for preprocessing. Affymetrix shares these data using a number of file suffixes: CDF (expression and SNP arrays), BPMAP (tiling arrays) and PGF + CLF (exon and gene arrays). NimbleGen distributes their annotations through NimbleGen Design Files (NDFs), regardless of the array type, but an additional position (POS) file, containing up-to-date genomic coordinates, is not uncommon. Table 1 describes the suffixes used by some manufacturers, and we note that oligo currently supports Affymetrix and NimbleGen arrays. Using the annotation files provided by the manufacturer, the researcher can create the annotation package using the pdlnfoBuilder BioConductor package.

Bolstad et al. (2003); Carvalho et al. (2007, 2010); Irizarry et al. (2003a, 2006a, b, 2008); Ritchie et al. (2009); Scharpf et al. (2008) show significant improvements on the results when alternative algorithms are used as replacement for the solutions provided by the manufacturers. In fact, the novel methodologies described by Carvalho *et al.* (2007); Irizarry *et al.* (2008); Scharpf *et al.* (2008) use early versions of the oligo package hereby described to implement their solutions. Regardless of the application, these depend on two factors: (i) the ability to access the data at the rawest possible

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Table 1. File suffixes used by manufacturers for different array types

Manufacturer	Intensities	Design	Туре
Affymetrix	CEL	CDF PGF + CLF BPMAP	Expression/SNP exon/gene tiling
Agilent Illumina NimbleGen	GPR TXT XYS	GAL BPM NDF	All All All

The raw data files contain observed fluorescence intensities used in analysis; the design files provide information specific to the array, such as dimensions, physical locations, probe types and sequences. Currently, oligo supports Affymetrix and NimbleGen arrays.

Table 2. Raw data and annotation classes used by the oligo package

FeatureSet class	PDInfo class	Arrays	
ExpressionFeatureSet	ExpressionPDInfo	Expression	
ExonFeatureSet	ExonPDInfo	Exon	
GeneFeatureSet	GenePDInfo	Gene	
SnpFeatureSet	SnpPDInfo	SNP	
SnpCnvFeatureSet	SnpCnvPDInfo	SNP+CNV	
TilingFeatureSet	TilingPDInfo	Tiling	

level, after image processing (which is beyond the scope of this material); and (ii) the existence of an environment that provides analysis and visualization tools. The manufacturers whose products are currently supported by oligo provide these data using CEL (Affymetrix) and XYS (NimbleGen) files. These files supply array coordinates and observed intensities, which can be easily imported with oligo after installing the associated annotation package created by pdlnfoBuilder.

The oligo package implements multiple classes to manage raw intensities. They are used to differentiate data originating from different applications, such as gene expression versus exon data. Because the same method can behave differently when applied to objects of distinct applications, we make use of this feature to increase the flexibility of the package. From the *eSet* class defined in Biobase, we created the *FeatureSet* class, the template for feature-level data subclasses. Each application has its own *FeatureSet* subclass, as shown in Table 2. We make no distinction in terms of the manufacturer: data generated on chips of the same application, even from different manufacturers, will belong to the same *FeatureSet* extension. This unified framework increases productivity, because the preprocessing steps for arrays on the same application are essentially the same, regardless of the manufacturer.

This infrastructure is beneficial for both developers and end-users. Developers can use oligo solutions to facilitate the integration of their tools with BioConductor. The researcher benefits from the unified model that the package makes available, as the consistency in data delivery and handling improves efficiency.

3 APPLICATIONS

In this section, we use the oligo package with data of different types. We demonstrate its use to preprocess gene expression data in

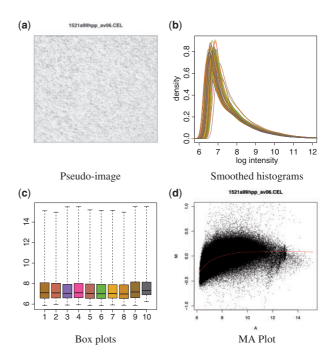


Fig. 1. The oligo package provides several tools for the visualization of raw data, represented in the package through the *FeatureSet* subclasses. In (a), the pseudo-image can be used to visually inspect the data for spatial artifacts. Using oligo, one can produce such figures using the *image* method. (b) shows the smoothed histogram, implemented in oligo via the *hist* method, providing a way to compare the distribution of intensities across multiple samples. In (c), we show boxplots generated with the *boxplot* method, also used to assess the data distribution. The *MAplot* method can be used to generate the MA plot shown in (d), used to assess the dependency of log-ratios on the average log-intensity of the data.

Section 3.1. Section 3.2 shows how it can be used to obtain genotype calls from SNP arrays. In Section 3.3, we show how it can be used to preprocess exon data at both exon and transcript levels. Section 3.4 uses data from tiling arrays to show how objects created by Oligo can be used with methods defined by other BioConductor packages. The Supplementary Material contains the actual code used in these examples.

3.1 Preprocessing expression arrays

After loading oligo, the user identifies the CEL or XYS files that represent the study in question. This is done with the list.celfiles or list.xysfiles functions. These functions accept the same arguments as list.files.

The read.celfiles and read.xysfiles functions import the CEL and XYS files into the R session. The returned object belongs to one of the *FeatureSet* classes shown in Table 2 and represents the raw data. These objects can be visualized through different strategies: Figure 1 shows, respectively, the pseudoimage, smoothed histograms, boxplots and MA plot for the Latin Square data on the Human Genome U95 array, made available by Affymetrix. Figure 1a–d are produced, respectively, with the *image*, *hist*, *boxplot* and *MAplot* methods.

The oligo package is tightly integrated with important BioConductor tools. Probe sequences are stored using the *DNAStringSet* objects, defined in the Biostrings package.

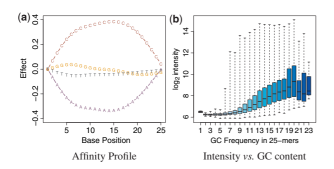


Fig. 2. The package is tightly integrated with other BioConductor tools to improve the user experience. (a) shows the affinity profile, which can be produced with oligo. In this figure, we can easily observe the clear interaction of nucleotide and position on the log₂-intensity. For (b), storing sequence information using the *DNAStringSet* class in Biostrings provides a compact representation of the data and allow efficient calculation, as shown above with the log₂-intensity boxplot stratified by GC content.

This allows a compact representation of the data and simplifies the interfacing with other tools. The getBaseProfile and getAffinitySplineCoefficients functions can be combined to obtain the affinity profile shown in Figure 2a. The alphabetFrequency method in Biostrings is easily used with the pm method in oligo to show the dependency of log₂ intensities on GC content, as shown in Figure 2b.

Expression data can be preprocessed using RMA (Irizarry et al., 2003a, b) by applying the rma method to the ExpressionFeatureSet object. The data will be background corrected (if the background argument is set to TRUE), quantile normalized (if the normalize argument is TRUE) and summarized using the median-polish. The resulting object is an ExpressionSet, defined by the Biobase package. The Supplementary Material shows one detailed example of how to use Oligo to preprocess the expression data.

3.2 Obtaining genotype calls from SNP arrays

Carvalho *et al.* (2007) describe the Corrected Robust Linear Model with Maximum likelihood distance (CRLMM) algorithm to genotype SNP arrays. This method is implemented in oligo and, to use it, the investigator needs the annotation data package specific to the design used in the experiment. These annotation packages are available through the BioConductor website and, because they contain hand-curated data, we recommend users refrain from creating (with the pdlnfoBuilder package) their own annotation packages for SNP chips. Table 3 describes the supported designs and the respective annotation packages.

To demonstrate its genotyping capability, we use 269 CEL files on the XBA array, available on the HapMap website. The crlmm function requires the CEL file names and an output directory, where the results are stored. The output directory must not exist prior to the call and the software will take care of this task. The crlmm function is applied directly on the CEL files, to minimize the memory footprint. A detailed demonstration on how to use crlmm is shown in the Supplementary Material.

The getCrlmmSummaries function reads the results obtained by CRLMM back into the R session. The *calls* and *confs* methods are accessors to genotype calls and confidence values. Calls are coded as integers 1 (AA), 2 (AB) and 3 (BB). The confidence score

Table 3. SNP array designs currently supported by the oligo package and their respective annotation packages

Design	Annotation Package		
Mapping 50K XBA Mapping 50K HIND Mapping 250K NSP Mapping 250K STY Genomewide SNP 5.0 Genomewide SNP 6.0	pd.mapping50k.xba240 pd.mapping50k.hind240 pd.mapping250k.nsp pd.mapping250k.sty pd.genomewidesnp.5 pd.genomewidesnp.6		

These annotation packages are made available through the BioConductor website and contain hand-curated data, required by the CRLMM algorithm.

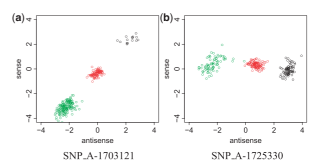


Fig. 3. Log-ratio data used by CRLMM for genotype calling, which can be seriously affected by probe effects. In this plot, genotype calls provided by Oligo are represented in different colors (black, AA; red, AB; green, BB) and each point represents one sample. SNP_A-1703121 shows significant discrimination on both strands and, as competing algorithms, CRLMM has excellent performance on similar scenarios. SNP_A-1725330 presents poor discrimination on the sense strand, because CRLMM does not average across strands, it can successfully predict the genotype calls. In comparable situations, competing algorithms are known to fail.

is the estimated probability that the algorithm made the right call. The plotM function provides the visualization of the results, as Figure 3a and b show.

3.3 Preprocessing exon arrays

The oligo package also supports the Affymetrix exon and gene arrays. Their annotation packages are available via BioConductor. These chips are extensions of the 3' IVT expression arrays and, as such, users are often interested in preprocessing them using the RMA algorithm. With these designs, the researcher can use oligo to obtain RMA summaries at exon and transcript levels.

The read.celfiles function imports any CEL file. The software identifies specifically if the files refer to exon or gene arrays and returns an object of the appropriate class. Raw data visualization can also be performed using the techniques presented in Section 3.1.

Similarly, the *rma* method provides RMA summaries and, in the case of the exon and gene arrays, accepts one extra argument: *target*. The possible values for *target* are probeset, core, extended and full. The first value will allow summarization to the exon level; the other three provide summarization to the gene level, using the Affymetrix definition of meta probesets.

The flexibility of oligo and its annotation packages allows integration with other BioConductor tools. Below, we use the

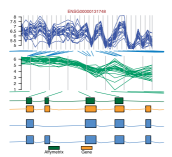


Fig. 4. Visual representation of the observed log₂-intensities and summarized data at the exon level for a fragment of gene ENSG00000131748. On the top panel of the figure, each line represents one different sample; the vertical bins represent the start and end positions for each probe (first subfigure) and probeset (second subfigure). On the bottom panel, the block diagram shows the probes, gene and transcript, respectively, in green, orange and blue. Here, the oligo, biomaRt, Biostrings, BSgenome and GenomeGraphs packages were used together to provide an improved visualization of the data at a specific genomic location.

biomaRt, Biostrings, BSgenome and GenomeGraphs packages to obtain more information on probesets and to visualize the results (see Fig. 4) at a specific genomic location. The Supplementary Material shows detailed information on how to combine sequence information obtained through oligo to get updated genomic coordinates by sequence alignment using the Biostrings and BSgenome packages.

3.4 Interfacing with ACME to find enriched regions using tiling arrays

The oligo package handles tiling data from both Affymetrix and NimbleGen, as long as the annotation packages are created through the pdlnfoBuilder package and installed in the system.

The functions read.celfiles2 and read.xysfiles2 can be used to import the data into the R session. The difference between read.celfiles/read.xysfiles and read.celfiles2/read.xysfiles2 is that the former reads in the data as one-channel data and the latter reads it in as two-channel data. The getNgsColorsInfo function parses the names of the XYS files and returns an object with suggested channels and sample names that can be combined with read.xysfiles2.

Using ChIP-chip data provided by NimbleGen, we use oligo to import the XYS files and combine their contents to create an object that can be used with the ACME package. The ACME package calculates enrichment, using algorithms that are insensitive to normalization strategies and array noise. We refer the interested user to the Supplementary Material, which contains detailed information on the use of oligo interfacing the ACME package. For this example dataset, we show below some enriched regions (flagged with TF=TRUE) found on Chromosome 1 for Sample 1:

Length	TF	StartInd	EndInd	Start	End
2179	FALSE	1	2179	56753	7925574
8	TRUE	2180	2187	7943079	7943879
18	FALSE	2188	2205	7943979	8009243
8	TRUE	2206	2213	8009343	8010043
251	FALSE	2214	2464	8010143	9893203
6	TRUE	2465	2470	9893303	9893803

4 DISCUSSION

The integration of data management, visualization and analysis is essential in current research. Weaknesses in existing tools are more evident today that more applications and array manufacturers are available. To overcome the deficiencies introduced by suboptimal solutions and improve the delivery of original strategies, we developed the Oligo package. It uses the commonalities of oligonucleotide microarray designs and applications to provide an open-source solution that centralizes the preprocessing tasks under a solid framework that can be reused by other developers, improving the consistency between packages within the BioConductor project.

The structure used by oligo is flexible and its objects inherit the properties of Biobase objects, using the standards set by the BioConductor project. Because it is implemented in R, every feature of this statistical software is at the researcher's disposal. This simplifies the interface with many other packages from both projects (R and BioConductor), widening the options during analysis. Classes used by its companion, the pdlnfoBuilder package, are based on a broader class, offering inheritability properties and transparency to the user, who benefits from the fact that annotation packages use SQL databases to minimize their memory footprint.

The oligo and affy packages are closely related: the former uses the knowledge acquired by the latter to provide solutions for limitations found so far. The main improvements offered by oligo are: (i) support to multiple vendors and platforms; (ii) efficient storage and access schemes for annotation of current high-throughput arrays, whose metadata have become significantly large; and (iii) native support to manufacturer files.

Using oligo, one can handle data from different applications (expression, tiling, SNP/CNV, exon and gene) of two manufacturers (Affymetrix and NimbleGen), using their native file schemes, avoiding potential problems introduced by conversion tools. We demonstrated how one can use our package to preprocess and visualize oligonucleotide microarray data. We show how the package can serve as an interface between the data files and methodologies implemented by other BioConductor packages. These features define a unified framework that allows the efficient use of the environment set by both R and BioConductor projects and increase the productivity of novel methods and algorithms.

ACKNOWLEDGEMENTS

Robert Gentleman, Wolfgang Huber, Martin Morgan, Seth Falcon, Marc Carlson, Vincent Carey, Robert Scharpf and James MacDonald for insights, comments and lengthy discussions on the package implementation. Ming-Wen An for suggestions that significantly improved the readability of the manuscript. Marvin Newhouse and Jiong Yang for all the help with the computational environment.

Funding: Doctoral scholarship awarded by the Brazilian Funding Agency CAPES (Coordenação de Aprimoramento Pessoal de Nível Superior) and National Institutes of Health grants R01RR021967 and P41HG004059.

Conflict of Interest: none declared.

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