The advent of new methods to obtain large-scale surveys of gene expression in which transcript levels can be determined for thousands of genes simultaneously has facilitated the expansion of biological understanding from the analysis of individual genes to the analysis of systems of genes (and proteins). This change characterizes the movement into the era of functional genomics. Central to this movement is an appreciation of the gene’s role in cellular activity as it functions in the context of larger molecular networks.

Two salient goals of functional genomics are to screen for key genes and gene combinations that explain specific cellular phenotypes (e.g. disease) on a mechanistic level, and to use genomic signals to classify disease on a molecular level. Signals generated by the genome must be processed to characterize their regulatory effects and their relationship to changes at both the genotypic and phenotypic levels. Since transcriptional (and posttranscriptional) control involves the processing of numerous and different kinds of signals, mathematical and computational methods are required to model the multivariate influences on decision-making in complex genetic networks.

Historically, it has been within the domain of signal processing where such methodologies have been extensively studied and developed—in particular, estimation, classification, pattern recognition, automatic control, information theory, networks, computation, imaging, and coding. Moreover, signal processing is based on a holistic view of regulation and communication. As a discipline, signal processing involves the construction of model systems composed of various mathematical structures, such as systems of differential equations, graphical networks, stochastic functional relations, and simulation models. Therefore it is not surprising that the advent of high-throughput genomic and proteomic technologies is drawing a growing interest from the signal processing community in relation to attacking the fundamental issues of expression-based functional genomics.

The twin aims of tissue classification and pathway modeling require a broad range of signal processing approaches, including signal representation relevant to transcription and system modeling using nonlinear dynamical systems. To capture the complex network of nonlinear information processing based upon multivariate inputs from inside and outside the genome, regulatory models require the kind of nonlinear dynamics studied in signal processing and control. Genomics requires its own model systems, not simply straightforward adaptations of currently formulated models. New systems must capture the specific biological mechanisms of operation and distributed regulation at work within the genome. It is necessary to develop nonlinear dynamical models that adequately represent genomic regulation for diagnosis and therapy.

Genomic signal processing (GSP) is the discipline that studies the processing of genomic signals. The aim of GSP is to integrate the theory and methods of signal processing with the global understanding of functional genomics, with special emphasis on genomic regulation. Hence, GSP encompasses various methodologies concerning expression profiles: detection, prediction, classification, control, and
dynamical modelling of gene networks. Moreover, since RNA coding is controlled by DNA sequencing, the analysis of DNA sequences, treated as signals in their own right, can be considered within the domain of GSP. Overall, GSP is a fundamental discipline that brings to genomics the structural model-based analysis and synthesis that form the basis of mathematically rigorous engineering.

This special issue of EURASIP JASP contains some examples of GSP applications. The issue starts with three papers (Song et al., Chakravarty et al., and Sussillo et al.) on spectral analysis of DNA sequences. The next paper by Hero et al. treats statistical signal-processing-based gene selection. The following two papers (Wu et al. and Giurcâneanu et al.) develop signal processing techniques for gene clustering. The next two papers treat DNA sequence segmentation using statistical signal processing (Nicorici and Astola) and image processing (Hua et al.), respectively. Signal processing methods for gene prediction and regulatory network inference are developed in the papers by Fox and Carreira, Zhou et al., and Ivanov et al., respectively. The paper by Cristea deals with revealing large-scale chromosome features by analysis of genomic signals. In addition, the paper by Lennartsson and Nordin treats peptides identification using genetic programming. Finally, an invited tutorial by Dougherty et al. discusses key issues in GSP.

The guest editors would like to thank all the authors for contributing their work to this special issue. We would also like to express our deep gratitude to all reviewers for their diligent efforts in evaluating all submitted manuscripts.

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