A Leader Genes Approach-based Tool for Molecular Genomics: From Gene-ranking to Gene-network Systems Biology and Biotargets Predictions

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Abstract

Nanogenomics, being the interplay of nanobiotechnologies and bioinformatics, is emerging as an intriguing approach in the field of nowadays biomedicine. Microarrays can produce a wealth of data and details, but they need an algorithm for data reduction to be clearly understood and exploited. The Leader Genes approach, integrating the different available databases and genomics tools, enables the user to search for genes linked to a disease or a cellular process, and to visualize the class of the most important genes, that is to say those having the highest number of interconnections. In this manuscript, we will review the algorithm which has been validated with both experimental and clinical studies. We will describe the different steps that lead to its final version, and we will discuss future perspectives and developments.

Keywords: Bioinformatics; Gene prioritization; Gene-ranking; Leader gene approach; Molecular genomics; Nanogenomics; Systems biology

Introduction

Nanogenomics [1-3] is emerging as a promising discipline in the field of biomedicine, being the interplay of nanobiotechnologies and bioinformatics. Gene microarrays and the other biomolecular high-throughput tools of the omics sciences (such as linkage and linkage disequilibrium, GWAS or genome-wide association study, QTL or quantitative trait loci and its derivative, such as eQTL, expression QTL, phenome profile, transcriptome analysis for genome mapping, and so on) [4], can produce an incredible wealth of data. If on one hand, these “big data” [5,6] are necessary for a deep and profound underpinning of the entire molecular picture, which is extremely complex, on the other hand, researchers risk to be overwhelmed by these data, without a proper data reduction.

For this reason, gene prioritization or gene-ranking has been advocated as a crucial task in bioinformatics, leading to a list of few strong candidate genes that can be further validated with ad hoc experiments. Elucidating genetic traits is particularly useful when studying complex and multi-factorial diseases [7,8], such as Alzheimer’s dementia and other neurological pathologies [9], diabetes and other metabolic disorders [10], cancer [11], and cardiovascular disorders [12]. Usually these diseases are chronic, present a relatively mild phenotype, are slowly progressive, and have a tremendous burden for the society and impact on the population in term of quality of life (QOL). The physio-pathology of complex pathologies is characterized by various biologic pathways and networks, dependent upon the contribution of a large number of genes and gene products, and therefore, the knowledge of molecular mechanisms of complex multifactorial diseases must include and deal with a large array of genes. These genes form complex networks of interactions, which may be direct (that is to say, physical interactions between the proteins, confirmed by experimental techniques, such as NMR or crystallography), or indirect (involvement in the same metabolic pathway or co-expression under different conditions).

There are a lot of gene-ranking methods, which for example, exploit the Pearson correlation with clinical information (such as Kaplan-Meier survival time) ([4] and references therein, [13]), or are based upon gene expression time series or gene co-expression series [14], regulatory networks (protein-protein interaction, PPI [15-18], genetic interaction with gene-gene interaction maps [19]) (for further explanations and references, the reader is referred [20,21], ortholog mapping, biomedical literature mining (ab initio approach), or integrating and combining the above mentioned approaches [4]. The techniques that are usually employed are: support vector machine, neural networks, fuzzy algorithms, kernelized score functions [22], and other learning machine procedures, biomedical annotation [23] (such as LSI, Latent Semantic Indexing [24], PosMed or Positional PubMed [25]), and clustering [26,27].

The main purposes of gene-ranking [28] are to provide robust molecular signatures of the studied events, to discover reliable biomarkers, as well as to predict gene function (GFP), or further biological targets of potential pharmaceutical interest [29].

In this manuscript, we will review our algorithm, which has been validated with both experimental and clinical studies. We will describe the different steps that lead to its final version, and we will discuss future perspectives and developments.

Materials and Methods

The Leader Gene approach exploits a search-dependent statistical approach in the field of nowadays biomedicine.

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algorithm, which is based on the systematic, exhaustive and recursive search for the genes and their products involved in a given process or event (biological, such as a biochemical or biophysical reaction, or pathological, like a disorder), on the computation of their interaction map, and on their ranking according to the number of all experimentally established interactions, as derived from free and open source Web-available databases, such as STRING (Search Tool for the Retrieval of Interacting Genes, Heidelberg, Germany) [30].

The systematic search is usually carried out mining different databases (like PubMed using National Library of Medicine vocabulary of terms or MESH terms, NCBI OMIM, GeneBank, GeneCards, GeneAtlas, using the standardized nomenclature HUGO), and repositories containing DNA microarrays data (normalized with any normalization technique, such as LOWESS, or locally weighted scatterplot smoothing, LOESS, or local regression), integrating the obtained hits.

The calculation of an interaction map of the obtained list of genes is computed using the weighted number of links (WNLs). This measure is calculated for each gene using the program STRING [30], and this value is derived from the weighted sum of three types of interactions:

1. literature co-occurrence of the names of genes, and/or their products in abstracts of papers available on the Internet. The scores assigned are derived from benchmarked and validated scoring system based on the frequencies and distributions of gene/gene products names in the aforementioned abstracts. The benchmarks themselves are set from manual evaluation of predictions of gene and protein interactions by experts, and are typically below 0.5;
2. scores derived from databases dedicated to gene networks, containing data on induction and expression of a particular genes by other genes derived from microarray experiments, or other high-throughput omics techniques. The score of 1 is assigned if the link is already present in the database, while putative links have lower values (typically 0.6–0.8);
3. the same range of scores is assigned to gene interactions via physically observed interactions between proteins. The software used does not discriminate between in vivo or in vitro experiment derived data. Generally the scores are close to those of interaction type 2, but links of this type occur much rarely than of type 2.

The combined association scores Sij were summed for each gene i over its neighbors (j), giving the final weighted number of links for the gene i. Further, we applied clustering methods to the genes. All the clustering algorithms and techniques available databases, as well as derived also from repositories containing DNA microarray experiments. This systematic search resulted in the identification of 238 related genes. Later, after this step, software and tools predicting interactions among those genes were employed, assigning scores to each of the genes, according to the number of interactions for each gene weighted by the significance of each interaction (WNLs), and finally applying several types of clustering algorithms to the genes. All the clustering algorithms and techniques applied, either hierarchical or K-means, invariably selected the same six "leader" genes involved in controlling the cell cycle of human T lymphocytes (namely, CDK2, CDK4, MYC, CDC2, CDKN1A, CDKN1B). CDK2 (Cyclin-Dependent Kinase 2) is involved at the transition from G1 to S binding Cyclin A and Cyclin E, while CDK4 at the progression in G1 phase. MYC is a gene involved in the early phase of T cells proliferation, controlling G1 phase. CDC2 (Cell Division Control 2, also known as CDK1) is a phosphoprotein belonging to the Ser/Thr kinases, and is also a catalytic subunit of the Maturation-Promoting Factor (MPF) complex. It is responsible of the transition from G1 to S and G2 to M phases, binding both Cyclin A and Cyclin B. CDKN1A (Cyclin-Dependent Kinase inhibitor 1A or CDK-interacting protein 1A, also known as p21Cip1/WAF) and CDKN1B (Cyclin-Dependent Kinase inhibitor 1B, also known as p27Kip1) are inhibitors

Given a set S of n objects {x1, x2, ..., xn}, where each object is described by attributes xi=(x1i, x2i, ..., xmi), determine a classification that is most likely to have generated the observed objects.

Genes belonging to the highest rank are defined as "leader genes" or "hub genes" because they may be assumed to play an important role in the analysed processes. The "Leader Gene approach" can suggest a list of few, but strong candidate genes potentially relevant within a given cellular process or a pathology, according to the already available experimental data. Moreover, the interaction map among all the genes involved in the same process may be useful in interpreting the experimental and clinical results, and in planning new targeted experimentation. Interestingly, such experimentation may be simpler to be analysed than mass-scale molecular genomics, whose wealth of details may raise problems and complications [1,2]. This computational method gave promising results, when applied to the human T lymphocyte cell cycle, human kidney transplant, oral lichen planus and periodontitis. These results were also integrated with a targeted experimental analysis, to draw an overall picture of these processes, and are reviewed in the following paragraphs.

This interactive, automatic and user-friendly stand-alone tool has been written in house in Java, Javascript, PHP and HTML. The completely automated pipeline is performed via NCBI e-utils (e-search, e-fetch, for further information the author is referred to the NCBI site), and other similar facilities.

A pictorial scheme of this tool is given in Figure 1, while the algorithm is shown in Figure 2.

The clustering techniques the user can choose are: Clustering K-means and Chinese whispers (which has been thought specifically for graph clustering, [31]); as far as the number of clusters is concerned, the user can choose from heuristic number or provided by the user himself. A screen-shot of the output is given in Figure 3.

Results

Human T-lymphocytes cell cycle

Quiescent non-activated peripheral blood T cells, when stimulated by antigen-presenting cells (APCs), enter the cell cycle via the CD3/CD28 pathway. T cells cycle is a key biological event leading to proliferation, increase in size and production of molecules, such as cell surface receptors and cytokines/interleukins. Moreover, impairment and dysregulation of T cells cycle lead to several important diseases. The key genes involved in the cell cycle of human T lymphocytes stimulated by the mitogen compound phytohemagglutinin or PHA [32-34], were identified by iterative and recursive searches of different genomic databases, as well as derived also from repositories containing DNA microarray experiments. This systematic search resulted in the identification of 238 related genes. Later, after this step, software and tools predicting interactions among those genes were employed, assigning scores to each of the genes, according to the number of interactions for each gene weighted by the significance of each interaction (WNLs), and finally applying several types of clustering algorithms to the genes. The key genes involved in the cell cycle control of human T lymphocytes are CDK2, CDK4, MYC, CDC2, CDKN1A, CDKN1B. CDK2 (Cyclin-Dependent Kinase 2) is involved at the transitions from G1 to S binding Cyclin A and Cyclin E, while CDK4 at the progression in G1 phase. MYC is a gene involved in the early phase of T cells proliferation, controlling G1 phase. CDC2 (Cell Division Control 2, also known as CDK1) is a phosphoprotein belonging to the Ser/Thr kinases, and is also a catalytic subunit of the Maturation-Promoting Factor (MPF) complex. It is responsible of the transition from G1 to S and G2 to M phases, binding both Cyclin A and Cyclin B. CDKN1A (Cyclin-Dependent Kinase inhibitor 1A or CDK-interacting protein 1A, also known as p21Cip1/WAF) and CDKN1B (Cyclin-Dependent Kinase inhibitor 1B, also known as p27Kip1) are inhibitors

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of cyclin-CDK2 and -CDK4 dependent complexes, blocking Cyclin D and Cyclin E, and thereby, contribute to the control of G1/S transition and G1 progression.

Moreover, human lymphocytes gene expression was monitored before and after PHA stimulation after 48 hours in an experiment, and over 72 hours in an experiment, using a commercial Human Starter DNA microarray technology, and an in-house instrumentation (DNASER). Results were then compared with our previous bioinformatics predictions, and we found that experimental data were comparable with our bioinformatics findings. We studied and investigated in details the specific roles of PCNA (Proliferating Cell Nuclear Antigen) and CCNA2 (Cyclin A2) at all phases of the cell cycle, and of CHEK1 (Checkpoint Kinase 1) in regulating DNA repair and preservation, and their interactions with the leader genes.

**Periodontitis**

Periodontitis [35] is a complex, multifactorial, chronic and
progressive inflammatory disease affecting the periodontium, that is to say the surrounding and supporting tissues of the teeth. These tissues include the gum (or gingiva), the periodontal ligament, the cementum and the alveolar bone. Because of bacterial infection caused by microbes such as Porphyromonas gingivalis, Prevotella intermedia, Bacteroides forsythus, Aggregatibacter actinomycetemcomitans, Fusobacterium nucleatum, Capnocytophaga species, Treponema denticola, and Campylobacter rectus, the periodontal tissues become inflamed, and are slowly destroyed by the action of the inflammatory process. If left untreated, the teeth lose their ligamentous support to the alveolar bone, become mobile, and are eventually lost. A large majority of periodontitis patients respond well to conventional therapies. However, the results of some studies have suggested that a small percentage of patients present a poor response. Those patients are defined as “downhill patients” (loss of 4 up to 9 teeth), or as “extremely downhill patients” (loss of 12 up to 23 teeth).

Genes involved in human periodontitis were tentatively identified and ranked according to their number of interactions to obtain a preliminary, broader view of molecular mechanisms of periodontitis and plan targeted experimentation. Genes were identified with interrelated queries of several databases. The interactions among these genes were mapped and given a significance score. The weighted number of links was calculated for each gene. Genes were clustered according to this parameter. The genes in the highest cluster were termed leader genes. Sixty-one genes involved or potentially involved in periodontitis were identified. Only five were identified as leader genes (namely, NFKB1, CBL, GRB2, PIK3R1, RELA), whereas 12 others were ranked in an immediately lower cluster (namely, IKBKB, SRC, IL6R, TRAF2, TNF, PDGFRB, IL6ST, IL4R, IL6, TNFRSF1A, CRK, IL1R1). For 10 of 17 genes (58.8%), there is evidence of involvement in periodontitis; seven new genes that are potentially involved in this disease were identified. The involvement in periodontitis has been completely established for only two leader genes.

**Kidney transplant**

A good rate of survival in patients receiving solid organ transplantation, which is the main objective of the research in the field, depends on life-long immunosuppression (such as corticosteroids and other immunosuppressive drugs), which may result in increased figures of infection and tumors. Induction of tolerance to allograft would represent the optimal solution for controlling both chronic rejection (CR), and side effects of immunosuppression. Although spontaneous “operational tolerance” can occur in human kidney or liver transplantation, the lack of noninvasive peripheral blood biological markers of this rare phenomenon precludes the identification of potentially tolerant patients in whom immunosuppression could be tapered, as well as the development of new tolerance inducing strategies. Here, the potential of high throughput microarray technology to decipher complex pathologies allowed us to study the peripheral blood specific gene expression profile, and corresponding Expression Analysis Systematic Explorer (EASE) molecular pathways associated to operational tolerance in a cohort of human kidney graft recipients. In comparison with patients with CR, tolerant patients displayed a set of 343 differentially expressed genes, mainly immune and defense related genes, in their peripheral blood mononuclear cells (PBMC), of which 223 were also different from healthy volunteers. Using the expression pattern of these 343 genes, we were able to classify correctly >80% of the patients in a cross-validation experiment, and classified correctly all of the samples throughout the years. Collectively, this study identifies a unique PBMC gene signature associated with human operational tolerance in kidney transplantation, using both by a classical statistical microarray analysis [36], and a nonstatistical analysis [37]. The findings obtained [37] were replicated using a murine model [38], which confirmed the importance of AKT pathway in the tolerance network.

Moreover, recent findings [39], carried out following the protocol shown in Figure 4 indicated that the SMILE/TMTC3 gene may be
Figure 4: Protocol of SMILE experiment.

Figure 5: LOWESS normalized and log-transformed gene expression values for the SMILE knock-down experiment: no treatment group, PMA activation group, SMILE silencing group and SMILE knock-down plus PMA activation group (up) and scatter plot of PMA activation, SMILE silencing and SMILE knock-down plus PMA activation genes versus no treatment genes (bottom).
involved in kidney graft operational tolerance in human. This gene was found to be up-regulated in blood from patients with a well functioning kidney transplant, in the absence of immunosuppression compared to other transplanted recipients with clinically different status. A microarray study of SMILE knock-down and phorbol 12-myristate 13-acetate (PMA) activation in HeLa cells (Figures 5-7) was herein compared to our earlier analysis based on microarray data of kidney allograft tolerance (Figure 8 and 10) and rejection (Figure 9 and 11) in humans, and in a rat model of allograft transplantation (Figure 12 and 13), to determine possible new genes and gene networks involved in kidney transplantation. The nearest neighbors at the intersection of the SMILE knock-down network with the human tolerance/rejection networks are shown to be NPHS1 and ARRB2, the former (Nephrin) being involved in kidney podocyte function, and the decrease of the latter ( Arrestin β2) being recently shown to be involved in monocyte activation during acute kidney allograft rejection in rat. Moreover, another one of the neighbors at the intersection of SMILE network and tolerance/rejection networks is XBP-1 (X-box binding protein 1), that we report previously to be increased, at a transcript level, after ER stress in SMILE silenced cells (Figure 14). Finally, we were also able to show that topological properties (both local and global) of joint SMILE knock-down network-tolerance/rejection networks and joint PMA activation network-tolerance/rejection networks in rat and human are essentially different, likely due to the inherent nature of the gene SMILE, and the mitogen compound PMA, that do not act the same way on genes and do not interfere the same way on networks. We also showed that interestingly SMILE networks contain more feed-forward loop (FFL) motifs, and thus, SMILE calls for a more fine-tuned genetic regulation.

**Oral lichen planus**

Oral Lichen Planus (OLP) is a chronic inflammatory oral mucosal disease that has an important global burden, both epidemiological and economic, affecting about 0.5-2% of the world population. In the *ab initio* theoretical study, genes involved in human OLP pathogenesis [40] are identified and ranked according to their number of interactions, in order to obtain a broader view of its molecular mechanisms, and to plan targeted experiments. Genes involved or potentially involved in OLP were identified by systematically querying several databases, until the identification of a final set of genes. Interactions among these genes were mapped and given a significance score using STRING database. For each gene, significance scores were summed to obtain a weighted number of links (WNL), and subsequently, genes were clustered according to this parameter. The genes in the highest cluster were termed as leader genes; the other ones were ranked as class B genes, class C genes, and so on. This study was complemented by a topological analysis of the network, carried out using Cytoscape, BinGO and FANMOD software. The interactions in the obtained network showed power law behaviour, in agreement with the scale-free topology theory of the biological graphs.

![Figure 6](image_url): Scatter plot of PMA activation log-transformed genes values (a) and scatter plot of SMILE knock-down log-transformed genes values (b).
Figure 7: Volcano plot for PMA activation (a) and Volcano plot for SMILE knock-down (b). These graphs can help in visualizing differentially expressed genes.

Figure 8: Computational properties and scale-free behavior of human kidney transplant network are shown. Human kidney transplant pro-tolerance bionetwork (in green) is shown together with (in red) PMA activation for HeLa cells.
132 genes were identified and five of them (namely, JUN, EGFR, FOS, IL2, ITGB4) were classified as leaders. JUN encodes a protein which interacts directly with specific target DNA sequences to regulate gene expression. This gene is mapped to 1p32-p31, a chromosomal region involved in both translocations and deletions in human malignancies. EGFR is a transmembrane glycoprotein that is a member of the protein kinase superfamily, and is a cell surface protein that binds to epidermal growth factor. Binding of the protein to a ligand leads to cell proliferation. FOS encodes leucine zipper proteins that can dimerize with proteins of the JUN family, thereby forming the transcription factor complex AP-1. FOS has been implicated as regulators of cell proliferation, differentiation and transformation. In some cases, expression of the FOS gene has also been associated with apoptotic cell death. IL2 is a secreted cytokine important for the

Figure 9: Computational properties and scale-free behavior of human kidney transplant network are shown. Human kidney transplant pro-rejection bionetwork (in green) is shown together with (in red) PMA activation for HeLa cells.

Figure 10: Computational properties and scale-free behavior of human kidney transplant network are shown. Human kidney transplant pro-tolerance bionetwork (in green) is shown together with (in red) SMILE knock-down for HeLa cells.
Figure 11: Computational properties and scale-free behavior of human kidney transplant network are shown. Human kidney transplant pro-rejection bioworknet (in green) is shown together with (in red) SMILE knock-down for HeLa cells.

Figure 12: Computational properties and scale-free behavior of rat kidney transplant network are shown. Rat kidney transplant bioworknet (in green) is shown together with (in red) PMA activation for HeLa cells.
proliferation of both T and B lymphocytes. ITGB4, a receptor for the laminins, mediates cell-matrix or cell-cell adhesion, and transduced signals that regulate gene expression and cell growth. It’s likely to play a pivotal role in the biology of invasive carcinoma. Interestingly, all of them but EGFR were up-regulated, and were widely distributed in the network (in terms of topological parameters such as stress, eccentricity and radiality), and showed higher topological coefficients than the other genes (Figure 15).

Conclusions
In conclusion, our Leader Genes approach-based tool provides a general executive framework for identifying and ranking genes associated with complex and multifactorial diseases. However, the approaches and techniques that were systematically implemented...
in the present study are general, and not confined by specific trait or species, and therefore, can be applied to various biological events, as well as diseases in different organisms. For this reason, the tool we presented is a general-purpose tool.

Moreover, in the last studies—specifically in the investigations on the role of SMILE/TMTC3 gene in kidney transplant operational tolerance, and on the etiology of OLP—we recently complemented the leader gene approach with a systems biology and topological analysis of the obtained graphs and networks. This is preliminary for further bioinformatics analysis and disease simulations using ad-hoc software. Topological analysis, in fact, can shed light on how molecular pathways work, and how a disease develops and evolves. Our analysis showed that our network exhibits a power law behaviour in agreement with the Scale-free theory of bio-networks, and has more FFL (feed-forward loops) than one would expect to find in a random graph (the so-called Erdős–Rényi random graph). The topological properties of leader genes and their role in controlling each pathway emerged from onthological analysis confirm our results.

For these reasons, we believe this tool can foster further research in the genomics field. Further developments are currently under study.

Figure 15: Oral Lichen Planus genes network (a), and its topological properties, namely the betweenness centrality coefficient (b), the number of shared neighbors (c) and the shortest path (d).

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