

Sequence analysis

INSECT 2.0: a web-server for genome-wide *cis*-regulatory modules prediction

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Abstract

INSECT is a user-friendly web server to predict the occurrence of *Cis*-Regulatory Modules (CRMs), which control gene expression. Here, we present a new release of INSECT which includes several new features, such as whole genome analysis, nucleosome occupancy predictions, and which provides additional links to third-party functional tools that complement user capabilities, CRM analysis and hypothesis construction. Improvements in the core implementation have led to a faster and more efficient tool. In addition, this new release introduces a new interface designed for a more integrative and dynamic user experience.

Availability and implementation: <http://bioinformatics.ibioba-mpsp-conicet.gov.ar/INSECT2>

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1 Introduction

The complexity of multicellular organisms is principally a product of the intricate regulatory networks through which they control cell expression, in a range of spatiotemporal conditions. The high specificity required for gene expression regulation is acquired mainly through the physical interaction of many Transcription Factors (TFs) that bind to groups of *cis* DNA elements (Transcription Factor Binding Sites, TFBSs) in a coordinated manner. These elements are often located close to each other and follow specific and highly conserved patterns, generally known as *Cis*-Regulatory Modules (CRMs).

Mathematical models of many TFBSs have been obtained and incorporated into publicly available databases. These models, usually represented as Position Weight Matrices (PWMs), can be used to predict new instances of regulatory sites over specific sets of sequences. However, the major obstacle for prediction of TFBSs, when aligning PWMs to gene regulatory sequences, is the vast amount of artifactual findings due to random and non-functional sites with high PWM scores (i.e. false positives).

INSECT first version (Rohr *et al.*, 2013) was presented as a CRM detector that outperforms most of the available tools with similar characteristics in terms of sensitivity and specificity, by the implementation of several strategies to lower the rate of false positives. The main advantage of our method is a more realistic semantic representation of CRMs structures. INSECT offers several features that allow users to easily extract information related to the genes of interest from the Ensembl annotations and other third-party tools, such as the UCSC Genome Browser. The tool interface minimizes usage complexity, allowing a highly integrative analysis to be performed.

INSECT 2.0 is significantly faster than its previous version, allowing for genome-wide analysis. New third-party tools have been included to help in the processes of further analysis and finding of CRMs, which can maximize the likelihood of representing true regulatory elements. INSECT 2.0 is a new online tool of great value to every researcher interested to predict the occurrence of CRMs at genome level and over long lists of genes, with no requirement to install prior infrastructure.

2 New features

Increased speed

A redesigned search core and data architecture have significantly boosted search speed. While the previous version was relatively fast when analyzing several hundred genes, INSECT 2.0 can perform genome-wide analysis in the same amount of time or faster, depending on the structure complexity of the CRMs and stringency parameters.

Genome-wide analysis and improved results dissection

Genome-wide analysis can be performed for 14 genomes from Ensembl. By simply defining distance limits relative to the TSS of each gene, INSECT 2.0 automatically retrieves the corresponding sequences and performs the search, providing several genomic annotations and links to third-party tools along with the results.

It has been shown that transcriptional regulation constrains the organization of genes across and within chromosomes (Janga et al., 2008). For this reason, INSECT 2.0 allows CRM searches on specific chromosomes, and the filtering of the resulting lists of genes by chromosome, when genome-wide analyses are performed.

The protein-coding genes-centric vision for gene expression regulation has until now underestimated the importance of many other gene categories present in genomes. Non-protein-coding genes are critically involved in the regulation of cellular processes, and have been proven to be as tightly regulated as protein-coding genes, as recently demonstrated for lnc-RNA genes (Alam et al., 2014). In addition, the implementation of a biotype filter provides the capability to perform analyses that are focused on non-protein-coding genes.

Chromatin structure and nucleosomes positioning within DNA may play a major role in allowing TFs–DNA interaction to proceed with transcriptional regulation. While high nucleosomes occupancy in DNA mainly acts as a barrier for most TFs to target the DNA, some others, called pioneer factors, may be biased to target silent chromatin and initiate cell-fate changes (Soufi et al., 2015). We have included a new feature for nucleosomes occupancy calculations for CRMs (Kaplan et al., 2009), offering more accurate estimation of whether the predicted site is a real regulatory element or not.

Additional PWM libraries

For users that do not have their own PWM for their TFs of interest, having a database of known PWMs is essential. In this new release, we have added several PWMs from Hocomoco (655 PWMs) (Kulakovskiy et al., 2013), SwissRegulon (189 PWMs) (Pachkov et al., 2013) and FlyFactorSurvey (486 PWMs) (Zhu et al., 2011). Taking into consideration those already included in our previous version, INSECT 2.0 has currently 2526 PWMs integrated in its PWM database.

Summary statistics and filters

Genomic CRMs search analysis could yield long lists of genes with several CRM hits on each of them. For this reason, we have implemented visualizations and filters that help to dissect and explore the search results.

TFs bind to specific elements at the regulatory regions of genes following certain patterns. Some TFBSs are distributed very close to the TSS of the target genes, while others show a decreased presence at those regions, shifting their occurrence to further regions (Chen et al., 2015). The distribution of distances of the predicted CRMs relative to the TSS for each gene and the PWM score for each TFBS

in the CRMs are calculated and visualized as histograms in the results page, in order to give the user additional information about the resulting binding patterns.

Dynamic filters for both TSS-relative distances and PWM scores are implemented. By adjusting these filters, results can be filtered on the fly, avoiding the need to run the job again with different parameters. This dynamism helps users to better handle the results and obtain as much information as possible by progressively changing search rules from flexible to more strict conditions, in order to identify those candidate genes with the most reliable CRM predictions.

Additional interaction with third-party tools

Prediction of CRMs is a difficult task given the amount of false positives that can arise during its search and evaluation. Despite PWMs being a useful representation of TFBSs, cells have several levels of complexity that account for transcriptional regulation beyond the occurrence of representative sites in the DNA. For this reason and to improve the working hypothesis, we have integrated several external tools to provide additional biological features about the genes that are predicted to contain potential CRMs. INSECT 2.0 is linked to GeneMANIA (Mostafavi et al., 2008), enabling users to select genes with predicted CRMs to inspect whether or not they are related, according to available bibliography. Furthermore, we have included a link to the Ensembl Regulatory Build (Zerbino et al., 2015), which aims to annotate regulatory elements in human and mouse genomes, by merging both experimental and computational derived data.

3 Conclusion

INSECT 2.0 is a powerful tool for researchers requiring search and analysis capabilities for potential CRMs presence at a genomic scale. This new release has added several features to help users further explore predicted CRMs based on the biological aspects of genome structure. The implementation of search filters reduces the search space to specific sets of genes of interest, minimizing the rate of false positives. New visualizations and statistical histograms of the global results have also been included. By linking the genes with predicted CRMs to additional third-party resources such as the UCSC Genome Browser, GeneMANIA or Ensembl Regulation, the powerful functionalities of its interface make INSECT 2.0 a unique integrative platform for CRMs prediction. The combined feature set allows users to exploit CRMs information that otherwise may remain hidden in huge tables, or that would require programming, data gathering or computational skills which would be barriers for non-specialized users.

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