SOS: Online probability estimation and generation of T and B cell receptors

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Recent advances in modelling VDJ recombination and subsequent selection of T and B cell receptors provide useful tools to analyze and compare immune repertoires across time, individuals, and tissues. A suite of tools—IGoR [1], OLGA [2] and SONIA [3]—have been publicly released to the community that allow for the inference of generative and selection models from high-throughput sequencing data. However using these tools requires some scripting or command-line skills and familiarity with complex datasets. As a result the application of the above models has not been available to a broad audience. In this application note we fill this gap by presenting Simple OLGA & SONIA (SOS), a web-based interface where users with no coding skills can compute the generation and post-selection probabilities of their sequences, as well as generate batches of synthetic sequences. The application also functions on mobile phones.

Availability and implementation: SOS is freely available to use at sites.google.com/view/statbiophysens/sos with source code at github.com/statbiophys/sos

INTRODUCTION

The adaptive immune systems recognises pathogens through the generation of a highly diverse repertoire of T and B cell receptors (TCR and BCR) which have the potential to recognise even unknown pathogen and initiate an immune response. In order to produce this diversity it exploits a highly stochastic process named V(D)J recombination. In addition, to block possible auto-reactive receptors, a selection process is mounted in the thymus for T cells, and a similar process of central tolerance is implemented for B cells. Probabilistic models of TCR and BCR have been proposed [3–5] based on immune repertoire sequencing data [7–10]. Software has been developed to infer the probability of generation of any B- or T-cell receptor (IGoR [1], OLGA [2] and SONIA [3])—have been publicly released to the community that allow for the inference of generative and selection models from high-throughput sequencing data. However using these tools requires some scripting or command-line skills and familiarity with complex datasets. As a result the application of the above models has not been available to a broad audience. In this application note we fill this gap by presenting Simple OLGA & SONIA (SOS), a web-based interface where users with no coding skills can compute the generation and post-selection probabilities of their sequences, as well as generate batches of synthetic sequences. The application also functions on mobile phones.

FEATURES

As explained in the introductory “About” tab, the web tool evaluates the generation and post-selection probability of single naive T and B cell receptors in different species based on the specific sequence the user inputs manually. The engine is based on two pieces of python software, OLGA and SONIA, and shipped with pre-trained models of recombination and selection for the following loci: human alpha and beta chains or TCR (TRA and TRB), human heavy and light chain of unmutated BCR (IGH, IGK, and IGL), and mouse TRB.

After choosing the species and receptor chain in the “Evaluate” tab, the user inputs a Complementary Determining Region 3 (CDR3), either as a nucleotide or an amino acid sequence, and optionally V and J germline genes from dropdown lists. The server outputs the generation probability (\(P_{gen}\), conditioned on sequence productivity), and the post selection probability (\(P_{post}\), as shown in Fig. 1 (left)). When V and J are not specified, the program sums over all possibilities for these segments to calculate the total probability of the CDR3.

To help interpret the result and assess how the sequence of interest compares to others, \(P_{gen}\), \(P_{post}\), and the selection factor \(Q = P_{post}/P_{gen}\) are plotted as green vertical lines on histograms of random sequences taken pre- (blue line) and post- (orange line) selection (Fig. 1 right). That feature only works when V and J are specified. The tool also provides an estimation of the probability to observe the sequence in a generic repertoire. The user inputs the size \(N\) of the sequenced repertoire (unique productive nucleotide sequences), and the tool outputs the probability of observing the sequence within a repertoire of that size, given by \(1 - (1 - P_{post})^N\).

Using the “Generate” tab, the user can synthetize a specified number of receptor sequences from \(P_{gen}\) or \(P_{post}\), after choosing the species and chain type from dropdown lists. The file with the generated sequences, composed of the CDR3 sequence (nucleotide and amino acid translation), V and J segments, is available for download as a CSV file. The user may fix the seed of the random number generator for reproducibility.
FIG. 1: **SOS web interface.** The user inputs a CDR3 sequence (amino acid or nucleotides) and V and J segments. The program outputs the generation probability $P_{gen}$, the probability in the periphery $P_{post}$, and evaluates a p-value corresponding to the probability of finding that sequence by chance in a repertoire of size $N$ (input by user). An additional tab allows for the generation of synthetic repertoires.

**DISCUSSION**

The interface can be used by investigators to evaluate how surprised one should be to find a given sequence in one or multiple repertoires. It could help distinguish receptors with a specific function from chance detections. The tool can also be used to evaluate the potential of certain receptors (in particular antibodies, albeit in their unmutated version) for vaccination or therapeutic purposes. The web interface is also available on mobile phones without the plotting options.

**Acknowledgments.** This work was partially supported by the European Research Council Proof of Concept Grant n. 824735

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\[ P_{gen} = 1 - (1 - P_{post})^N = 1 - e^{-Np_{post}} = 2.4\times10^{-9} \]