

# Genomic Time-Series Modeling Using Recurrent Neural Networks

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## Abstract

The advent of Next Generation Sequencing (NGS) has made it possible to study genomic data throughout time. This modern paradigm, Evolve-and-Resequencing (E&R), enables us to make more accurate and robust inferences, i.e. estimate model parameters, using multiple observations along generations. In this paper, we consider the recently repopularized Recurrent Neural Networks (RNN) to model the genomic time series of E&R. In fact, RNN is used as a generative model which for a initial estate and a choice of model parameter generates a sequence. Parameter estimation procedure involves a (non-convex) optimization of least square loss between observed sequence data and RNN-generated sequence with respect to model parameter. Backpropagation-in-time, is effectively used to compute gradients of objective function, and stochastic gradient descent with momentum algorithm is used for optimization. Experimental study on simulated data shows RNN provides significantly more accurate and robust estimates in shorter times.

## 1. Introduction

Until very recently, biological data analysis has been considered processing a snapshot of data. However, the emergence of NGS and related technologies has made it possible to not only create larger datasets but also to measure multiple observations of the same quantity in the course of time. In many cases, such as population genetics, it is of the great interest to model the evolutionary process and make inferences, predictions and retrospective studies. Indeed, a random process is better explained by time series data than a single observation.

In addition to inexpensive data availability, over last two decades, a large amount of efforts is dedicated to High-Performance Computing (HPC), which re-popularized and re-branded computationally intensive algorithms such as Neural Networks. The first properly proposed neural network model to exploit full potential of multi layer neural networks published by [Hinton and Salakhutdinov \(2006\)](#); [Hinton et al. \(2006\)](#) and its spectacular performance on image processing problems immediately spawned the field of Deep Neural Networks (DNN), aka Deep Learning. Shortly after, DNNs went beyond the tasks that they are initially indented to accomplish [Krizhevsky et al. \(2012\)](#) and had breakthroughs in time-series DNN models, aka RNNs, such as generative models [Sutskever et al. \(2011\)](#), speech processing [Hinton et al. \(2012\)](#) etc.

In this paper we aim to use the tools and machinery that has been developed for RNNs, to model times-series biological data. In particular, we consider the population genetics

problem of finding loci (locus) under selection given observation of allele frequency of a population in different generations of a Wright-Fisher model.

Terhorst et al. (2015) Stephan et al. (2006) Hudson (2002) Peng and Kimmel (2005) Sutskever (2013) Bergstra et al. (2010) (Rumelhart et al., 1986)

## 2. Methods

## 3. Experiments

## 4. Discussions

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