

# Studying three-dimensional genome organization using Hi-C

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## Summary

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*Keywords:* Hi-C , 3D genome organization , evaluation

## Introduction

### Background information

Background information goes here ...

### Motivation

Motivation goes here ...

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## **Materials-Methods**

### **Results**

#### **Complexity-Timing Analysis**

##### **Focus area**

I focused on the use of Python packages (Numpy, Pandas) as well as on visualization (with matplotlib and Pandas).

## **Conclusion-Discussion**

It is clear from the analysis presented in this study that even state-of-the-art Hi-C analysis techniques do not give extremely reproducible results. When two different restriction enzymes are used with the same sample as source, while the correlation is relatively high, it is not ideal. Moreover, when the resolution is increased (128kb bins instead of 4096kb bins), the correlation even in the case of technical replicates treated with the same enzymes is very low. Thus, new more robust analysis techniques that lead to more reproducible results, are required. This is going to be a large part of my PhD thesis work.

While I optimized the code for speed, by using Numpy instead of native Python to deal with matrices and list comprehension instead of for loops when possible, the real bottleneck is the I/O operations (reading input and writing output). This is critical as the matrix files that I have to deal with, are really large. One solution to the problem would be to minimize I/O operations by using Numpy for everything, but this would require much

better knowledge of Numpy than I currently have. Moreover, Numpy may be not as versatile or efficient as R in certain circumstances which means that I may be unable to fully replace R with Numpy. Another approach would be to use rpy or rpy2 for R/Python integration but I have been always facing problems with rpy/rpy2 installation on my system.

As far as the visualization is concerned, both Pandas and matplotlib look interesting but:

1. They need time to learn.
2. There are many requirements (dependencies etc) and in many cases it is difficult to run code using these packages on the cluster.
3. The resulting graphs, especially in the case of Pandas, do not seem to be extremely customizable. Moreover, I had to write more code to achieve the same result I would get with writing less code in R.

I will ceratinly try to explore Numpy, matplotlib, Pandas and other packages further, as it is always possible that the drawbacks I see right now compared to R may be just due to lack of expertise on all these Python packages.

## References