

Report

Contents

Background	2
Introduction	2
Analyse	3
Results	6
Conclusions	7
References	7

Background

COVID-19 is a single-stranded positive RNA (+ss-RNA) virus from the betacoronavirus family, which also includes MERS-CoV, and SARS-CoV. Coronaviruses possess the largest genome (30 kb) of the RNA viruses, contributing to their complex genome regulation strategies. Once COVID-19 infects a human host cell, it makes use of the proteins and host machinery including human RNA binding proteins (RBPs) that bind to RNA for processing such as post transcriptional modifications, regulation of stability and degradation (Tay et al. 2020). Several human RBPs have been previously shown to directly interact with coronaviruses to regulate their transcription, replication and translation. The full range of RBPs that may interact with COVID-19 remain to be described. In order to make strides towards the development of therapeutic strategies for COVID-19 and to understand the pathology of the virus, it is important to study the interactions between the viral genome and RBPs. If an RBP is involved in COVID-19 replication, targeting it for inhibition may reduce the rate at which the virus reproduces and may yield better clinical results in patients. Previously, as part of BioHackathon 2020, Ferraini et al. (2020) identified a set of proteins predicted to bind to the SARS-CoV-2 genome in order to characterize host-virus interactions post-infection and suggest pharmaceutical interventions for COVID-19 patients. Their workflow used position-weight matrices to locate potential binding sites for human RBPs on the genomes of SARS-CoV-2, SARS-CoV-1, and RaTG13, as well as simulated genomes that shared the nucleotide compositions of the aforementioned viruses. The difference in number of binding sites for each RBP between real and simulated genomes was tested for statistical significance to identify enriched RBP binding sites, and these binding sites were compared between viruses. This analysis identified 38 human RBPs that bind with SARS-CoV-2’s genome, a handful of which do so exclusively.

Introduction

To build upon the work by Ferraini et al., we will first reproduce the pipeline presented in the hackathon to obtain baseline results. This includes differential expression analysis to identify human mRNAs that are significantly up or down-regulated in COVID-19 infected cells. Since the release of the original hackathon results, new publications have revealed datasets that may be useful for modeling COVID-19 and human RBP interactions. For example, a study from December 2020 (Schmidt et al. 2020) published a list of 104 human RBPs they found to be experimentally interacting with COVID-19. Similarly, another study from 2021 mapped COVID-19 RBP interactions (Kamel et al. 2021). Thus, we plan on building upon previous methods by integrating known interactions for training a classifier that could identify COVID-19 interacting RBPs. Further, once we obtain a high confidence candidate list of RBPs, we can further evaluate their druggability to suggest human RBPs for further study (Gordon et al. 2020). We will divide the list of published COVID-19 interacting RBPs into training and testing sets to train a classifier using viral sequence information and differential expression results for RBP mRNAs. This model is likely to yield additional interacting candidates that have not yet been described

Analyse

Project composition and functions

They first created a file (rbp_functions) with all the functions they needed: - to read the different type of file (readGFF, readFasta, readPWMsFromFasta), - to calculate PWM information content (GetPWMEntropy) - to scan sequence(s) with multiple PWMs (ScanSeqWithPWMs) - to simulate a DNA sequence (simulateSeq) - to do enrichment test for RBP (enrich_rbps_real)

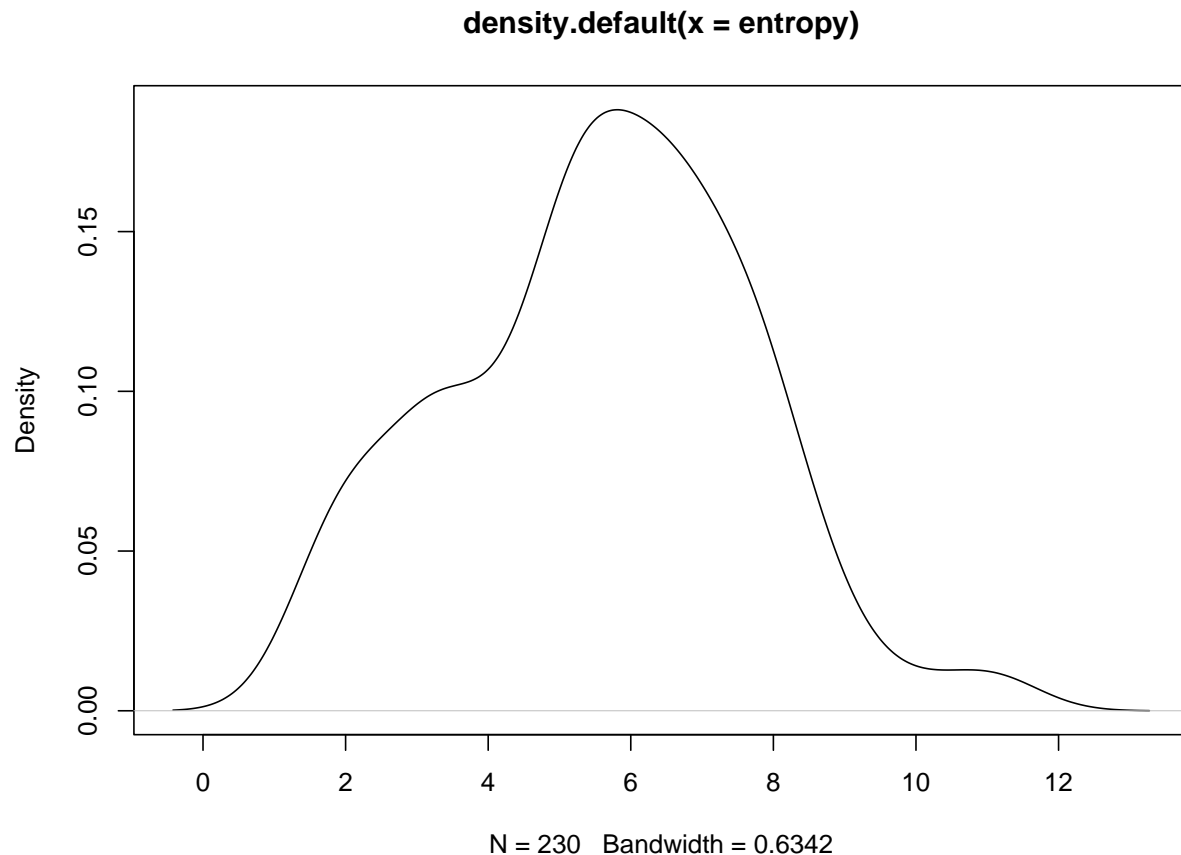
Then they created 7 seven files to analyse and to find binding sites enriched in the SARS-CoV-2 genome: - filter_ATtRACT - find_binding_sites - annotate_sites - simulate_genomes - find_sim_binding_sites - enrichment_analysis - filter_hits

1) RNA-binding proteins (filter_ATtRACT)

We start by filtering the RNA-binding proteins (RBP) coming from the daTabase of RNA binding proteins and AssoCiated moTifs (ATtRACT). Before filtration, there were 327 RBPs. We keep only human RBP (159 RBP) and we remove the ones consisting of only a single motif (106 RBP).

Then we filter the position weight matrices (PWM) which are used to represent motifs (patterns) in biological sequences by matching them with RBPs and by removing high entropy PWM. We reduced the number of PWMs from 1583 to 230 and the number of RBPs in the search from 106 to 102.

We can visualize the plot of the density vs the entropy.



2) Genome

We upload the SARS2 genome and we scan the genome with the PWMs. After the selection of unique RBP-PWM mappings and finding duplicate binding sites for the same protein and choose the one with the highest score, we reduced number of sites from 30597 to 29290.

3) Annotated sites

We load the gff file. We can visualize the head of the file.

	seqnames	start	end	width	strand	source	type	score	phase	ID	Dbxref	collection.date
1	NC_045512.2	1	265	265	+	RefSeq	five_prime_UTR	NA	NA	id-NC_045512.2:1..265		NA
2	NC_045512.2	266	21555	21290	+	RefSeq	gene	NA	NA	gene-GU280_gp01	GeneID:4...	NA
3	NC_045512.2	266	13468	13203	+	RefSeq	CDS	NA	0	cds-YP_009724389.1	Genbank:...	NA
4	NC_045512.2	13468	21555	8088	+	RefSeq	CDS	NA	0	cds-YP_009724389.1	Genbank:...	NA
5	NC_045512.2	266	805	540	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:1..180		NA
6	NC_045512.2	806	2719	1914	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:181..818		NA
7	NC_045512.2	2720	8554	5835	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:819..2763		NA
8	NC_045512.2	8555	10054	1500	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:2764..3263		NA
9	NC_045512.2	10055	10972	918	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:3264..3569		NA
10	NC_045512.2	10973	11842	870	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:3570..3859		NA
11	NC_045512.2	11843	12091	249	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:3860..3942		NA
12	NC_045512.2	12092	12685	594	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:3943..4140		NA
13	NC_045512.2	12686	13024	339	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:4141..4253		NA
14	NC_045512.2	13025	13441	417	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:4254..4392		NA
15	NC_045512.2	13442	13468	27	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:4393..5324		NA
16	NC_045512.2	13468	16236	2769	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:4393..5324		NA
17	NC_045512.2	16237	18039	1803	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:5325..5925		NA
18	NC_045512.2	18040	19620	1581	+	RefSeq	mature_protein_region_of_CDS	NA	NA	id-YP_009724389.1:5926..6452		NA

In bioinformatics, the general feature format (gene-finding format, generic feature format, GFF) is a file format used for describing genes and other features of DNA, RNA and protein sequences.

We can also visualize the head of the sites file.

Matrix_id	seqname	source	feature	start	end	score	strand	frame	seq	Gene_name	len
75	NC_045512.2	TFBS	TFBS	1	6	2.999063	+	.	ATTAAA	PPIE	6
75	NC_045512.2	TFBS	TFBS	1	6	2.999063	-	.	TTTAAT	PPIE	6
s63	NC_045512.2	TFBS	TFBS	1	4	3.450980	+	.	ATTA	TIAL1	4
s1	NC_045512.2	TFBS	TFBS	3	7	3.700284	-	.	CTTTA	ELAVL2	5
s6	NC_045512.2	TFBS	TFBS	3	7	3.700284	-	.	CTTTA	TIAL1	5
s9	NC_045512.2	TFBS	TFBS	3	7	3.737562	-	.	CTTTA	TIA1	5
143	NC_045512.2	TFBS	TFBS	4	8	2.997815	-	.	CCTTT	PTBP1	5
155	NC_045512.2	TFBS	TFBS	4	9	3.664262	-	.	ACCTTT	ZFP36	6
165	NC_045512.2	TFBS	TFBS	4	7	2.496259	-	.	CTTT	PTBP1	4
s21	NC_045512.2	TFBS	TFBS	6	11	4.189243	+	.	AGGTTT	ZRANB2	6
M033_0.6	NC_045512.2	TFBS	TFBS	7	14	5.822878	-	.	TATAAACC	KHDRBS3	8
s38	NC_045512.2	TFBS	TFBS	7	10	1.884956	-	.	AACC	HNRNP	4
s1	NC_045512.2	TFBS	TFBS	8	12	3.700284	+	.	GTTTA	ELAVL2	5
s6	NC_045512.2	TFBS	TFBS	8	12	3.558239	+	.	GTTTA	TIAL1	5
s9	NC_045512.2	TFBS	TFBS	8	12	3.737562	+	.	GTTTA	TIA1	5
75	NC_045512.2	TFBS	TFBS	9	14	2.999063	+	.	TTTATA	PPIE	6
75	NC_045512.2	TFBS	TFBS	9	14	2.999063	-	.	TATAAA	PPIE	6
s21	NC_045512.2	TFBS	TFBS	12	17	4.488048	-	.	AGGTAT	ZRANB2	6

The new file is more complete.

seqnames	start	end	width	strand	Matrix_id	source	feature	score	frame	seq	Gene_name	len	type	gene
NC_045512.2	1259	1265	7	+	s50	TFBS	TFBS	4.073349	.	CAGACGG	SRSF1	7	gene	ORF1ab
NC_045512.2	1259	1265	7	+	s50	TFBS	TFBS	4.073349	.	CAGACGG	SRSF1	7	CDS	ORF1ab
NC_045512.2	1259	1265	7	+	s50	TFBS	TFBS	4.073349	.	CAGACGG	SRSF1	7	CDS	ORF1ab
NC_045512.2	1259	1265	7	+	s50	TFBS	TFBS	4.073349	.	CAGACGG	SRSF1	7	mature_protein_region_of_CDS	NA
NC_045512.2	1259	1265	7	+	s50	TFBS	TFBS	4.073349	.	CAGACGG	SRSF1	7	mature_protein_region_of_CDS	NA
NC_045512.2	1260	1266	7	+	M020_0.6	TFBS	TFBS	4.757508	.	AGACGGG	FXR2	7	gene	ORF1ab
NC_045512.2	1260	1266	7	+	M020_0.6	TFBS	TFBS	4.757508	.	AGACGGG	FXR2	7	CDS	ORF1ab
NC_045512.2	1260	1266	7	+	M020_0.6	TFBS	TFBS	4.757508	.	AGACGGG	FXR2	7	CDS	ORF1ab
NC_045512.2	1260	1266	7	+	M020_0.6	TFBS	TFBS	4.757508	.	AGACGGG	FXR2	7	mature_protein_region_of_CDS	NA
NC_045512.2	1260	1266	7	+	M020_0.6	TFBS	TFBS	4.757508	.	AGACGGG	FXR2	7	mature_protein_region_of_CDS	NA
NC_045512.2	1262	1266	5	+	s19	TFBS	TFBS	3.113263	.	ACGGG	SRSF5	5	gene	ORF1ab
NC_045512.2	1262	1266	5	+	s19	TFBS	TFBS	3.113263	.	ACGGG	SRSF5	5	CDS	ORF1ab
NC_045512.2	1262	1266	5	+	s19	TFBS	TFBS	3.113263	.	ACGGG	SRSF5	5	CDS	ORF1ab
NC_045512.2	1262	1266	5	+	s19	TFBS	TFBS	3.113263	.	ACGGG	SRSF5	5	mature_protein_region_of_CDS	NA
NC_045512.2	1262	1266	5	+	s19	TFBS	TFBS	3.113263	.	ACGGG	SRSF5	5	mature_protein_region_of_CDS	NA
NC_045512.2	1263	1269	7	+	167	TFBS	TFBS	3.749375	.	CGGGCGA	SRSF1	7	gene	ORF1ab
NC_045512.2	1263	1269	7	+	167	TFBS	TFBS	3.749375	.	CGGGCGA	SRSF1	7	CDS	ORF1ab
NC_045512.2	1263	1269	7	+	167	TFBS	TFBS	3.749375	.	CGGGCGA	SRSF1	7	CDS	ORF1ab
NC_045512.2	1263	1269	7	+	167	TFBS	TFBS	3.749375	.	CGGGCGA	SRSF1	7	mature_protein_region_of_CDS	NA

4) Simulated genomes

3 sets of 5000 simulations - All the genome with proba of each letter calculated before - 3' - 5'

5) Find the binding sites of the simulated genomes

Pour chaque simulation

6) Enrichment analysis

Results

Conclusions

References