

A flexible genome-scale SARS-CoV-2 clone resource

Dae-Kyum Kim^{1,2,3,11}, Jennifer J. Knapp^{1,2,3,11}, Da Kuang^{1,2,3,11}, Aditya Chawla^{1,2,3}, Patricia Cassonnet^{4,5,6}, Hunsang Lee^{1,2}, Dayag Sheykhkarimli^{1,2,3}, Payman Samavarchi-Tehrani³, Hala Abdouni³, Ashyad Rayhan^{1,2,3}, Oxana Pogoutse^{1,2,3}, Étienne Coyaud⁷, Sylvie van der Werf^{4,5,6}, Caroline Demeret^{4,5,6}, Anne-Claude Gingras^{2,3}, Mikko Taipale^{1,2,8}, Brian Raught⁹, Yves Jacob^{4,5,6,*}, Frederick P. Roth^{1,2,3,10,*}

¹ Donnelly Centre, University of Toronto, Toronto, Ontario, M5S 3E1, Canada

² Department of Molecular Genetics, University of Toronto, Toronto, Ontario, M5S 1A8, Canada

³ Lunenfeld-Tanenbaum Research Institute, Sinai Health System, Toronto, Ontario, M5G 1X5, Canada

⁴ Unité de Génétique Moléculaire des Virus à ARN, Département Virologie, Institut Pasteur, Paris, 75724, France

⁵ UMR3569, Centre National de la Recherche Scientifique, Paris, 75015, France

⁶ Université de Paris, Paris, 75015, France

⁷ Univ. Lille, Inserm, CHU Lille, U1192 - Protéomique Réponse Inflammatoire Spectrométrie de Masse - PRISM, F-59000 Lille, 59655, France

⁸ Molecular Architecture of Life Program, Canadian Institute for Advanced Research, Toronto, Ontario, M5G 1M1, Canada

⁹ Department of Medical Biophysics, Princess Margaret Cancer Centre, University of Toronto, Toronto, Ontario, M5G 2C1, Canada

¹⁰ Department of Computer Science, University of Toronto, Toronto, Ontario, M5T 3A1, Canada

¹¹ These authors contributed equally.

* Correspondence: yves.jacob@pasteur.fr (Y.J), fritz.roth@utoronto.ca (F.P.R.)

Summary

The world is facing a major health crisis, the global pandemic of COVID-19 caused by the SARS-CoV-2 coronavirus, for which no approved antiviral agents or vaccines are currently available. Here we describe a collection of codon-optimized coding sequences for SARS-CoV-2 cloned into Gateway-compatible entry vectors, which enable rapid transfer into a variety of expression and tagging vectors. The collection is freely available via Addgene. We hope that widespread availability of this SARS-CoV-2 resource will enable many subsequent molecular studies to better understand the viral life cycle and how to block it.

Keywords: SARS-CoV-2, coding sequence collection, Gateway-compatible

Introduction

A global pandemic of the coronavirus disease COVID-19, a severe respiratory illness caused by a novel virus from the family *Coronaviridae* (SARS-CoV-2), has infected millions and caused hundreds of thousands of deaths (World Health Organization, 2020a). COVID-19 manifestation in patients can range from asymptomatic (no symptoms) to severe pneumonia and death (Huang et al., 2020). Early analysis of the outbreak in China outlines symptoms that commonly include fever, dry cough, shortness of breath and myalgia (World Health Organization, 2020b). Person-to-person spread through respiratory droplets has been identified as a major source of transmission of the virus (Yu et al., 2020). To limit contagion, various measures from social distancing to nationwide lockdowns, have been imposed to contain and control the transmission of SARS-CoV-2 (Cohen and Kupferschmidt, 2020). Despite these measures, the number of confirmed COVID-19 cases has continued to rise (World Health Organization, 2020a), highlighting the need for an effective vaccine and antiviral agents. Furthermore, the extrapolations concerning the evolution of the pandemic are particularly alarming (Ferguson et al., 2020). It is therefore of intense and pressing interest to better understand this virus and its interaction with host cells on a molecular level.

Shortly after the outbreak, the complete genome of two SARS-CoV-2 strains were published (Chan et al., 2020; Wu et al., 2020). Using the genome sequence as a reference, Chan *et al.* (Chan et al., 2020) identified 12 viral open reading frames (ORFs), including ORF1ab, a large polyprotein which is post-translationally processed into 16 proteins. More

recently, Wu *et al.* discovered two additional viral ORFs (ORF9Bwu and ORF10wu) with unclear functions (Wu et al., 2020). Progress on molecular characterization has been made on several viral proteins (Walls et al., 2020; Zhang et al., 2020), providing valuable insights into host-virus interaction. However, more research is necessary. The Gateway system offers efficient and high-throughput transfer of the viral coding sequences (CDSs) into a large selection of Gateway-compatible destination vectors used for protein expression in many biological systems, e.g. *Escherichia coli*, *Saccharomyces cerevisiae*, insect, or mammalian cells (Walhout et al., 2000). Broad availability of a collection of SARS-CoV-2 CDSs has the potential to enable many downstream biochemical and structural studies and thus a better understanding of processes within the viral life cycle, possibly yielding scalable assays for screening drug candidates that could disrupt these processes.

Results and Discussion

A total of 94 clones are currently included in the Gateway-compatible collection, covering 28 out of 29 total annotated CDSs in the SARS-CoV-2 genome. NSP11 was omitted due to its 36 base pair length, which makes it incompatible with the Gateway cloning system (ThermoFisher). All 28 of these CDS regions are available in clones with and without termination codons. The 'no-stop' collection was further extended to include six clones encoding different cleaved products of the spike (S) protein — S-fragment 1–6. We also included two CDS variants with in-frame deletions (S-24nt and E-27nt) and one truncated CDS variant (ORF8B-truncated) that were detected by recent viral transcriptome mapping efforts (Davidson et al., 2020; Kim et al., 2020).

Although our collection facilitates tagging of SARS-CoV-2 proteins for various functional studies, certain applications require removal of tags at some stage, for example, after protein purification. Fusion proteins can potentially interfere with the yield, structure, and function of purified proteins, such as during large scale production and crystallography studies. To address this we have expanded our collection to include clones containing an N-terminal recognition sequence for nuclear inclusion protease from tobacco etch virus (TEV) (Carrington and Dougherty, 1987; Carrington and Dougherty, 1988). The TEV sequence is one of the best characterized and widely used endoproteolytic reagents due to its stringent sequence

specificity, ease of production, and ability to tolerate a variety of residues at the P1' position of its recognition site (Waugh, 2011).

To promote open-access dissemination of the collection, all clones have been deposited to Addgene (Kamens, 2015). Table S2 summarizes all CDSs in the collection, together with their nucleotide sequences, nucleotide and amino acid lengths and direct links to Addgene.

We hope that this SARS-CoV-2 CDS-clone collection will be a valuable resource for many applications, including study of how coronaviruses can exploit host cellular processes for the viral replication cycle (de Wilde et al., 2018), understanding virus-host protein-protein interactions (Gordon et al., 2020; Lasso et al., 2019), production of recombinant virus proteins for structural studies (Edavettal et al., 2012), mapping of protein subcellular localization using N-terminal fluorescent reporters (Tanz et al., 2013), or development of vaccines or other therapeutics (Jing et al., 2012; McDonald et al., 2007).

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Authors Contribution

Conceptualization, D.-K.K., B.R., Y.J., and F.P.R.; Methodology, D.-K.K., J.J.K., H.L., D.S., M.T., and F.P.R.; Investigation, D.-K.K., J.J.K., A.C., P.C., H.L., D.S., P.S.-T., H.A., A.R., and O.P.; Writing – original draft, D.-K.K., D.K. and D.S.; Writing – review & editing, D.-K.K., J.J.K., D.K., H.L., A.R., Y.J., and F.P.R.; Supervision, É.C., S.V.D.W., C.D., A.-C.G., M.T., B.R., Y.J., and F.P.R.; Funding acquisition, D.-K.K., A.-C.G., B.R., Y.J., and F.P.R.

Declaration of Interests: The authors declare no competing interests.

Table 1. The genome-scale SARS-CoV-2 coding sequence clone collection.

Gene Symbol	CDS Name	Putative Function/Domain	AA Length
ORF1AB	NSP1	Suppress antiviral host response	180
	NSP2	Unknown	639
	PLPRO (NSP3)	Putative PL-pro domain	1,946
	NSP4	Complex with NSP3 & 6 for DMV (double-membrane vesicle) formation	501
	NSP5	3CL-pro domain	307
	NSP6	Complex with NSP 3 & 4 for DMV formation	291
	NSP7	DNA primase subunits	84
	NSP8		199
	NSP9	RNA/DNA binding activity	114
	NSP10	Complex with NSP14: Replication fidelity	140
	RNA-pol (NSP12)	RNA-dependent RNA polymerase	919
	Heli (NSP13)	Helicase	602
	NSP14	ExoN: 3'-5' exonuclease	528
	NSP15	XendoU: poly(U)-specific endoribonuclease	347
	NSP16	2'-O'-MT: 2'-O-ribo methyltransferase	299
S	S	Spike glycoprotein trimer that binds to host cell receptors (e.g. ACE2)	1,273
S	S-24nt	Spike glycoprotein trimer (minus 8 amino acids)	1,265
S	S-fragment1	Entire Ectodomain	1,213
S	S-fragment2	Entire Ectodomain without the signal peptide	1,199
S	S-fragment3	N-term fragment after the furin cleavage	686
S	S-fragment4	N-term fragment after the furin cleavage without the signal peptide	672
S	S-fragment5	C-terminal Ectodomain from the furin cleavage site	528
S	S-fragment6	C-terminal Ectodomain from the Tmpress 2 priming site	399
ORF3A	3A	Induce inflammatory response and apoptosis	275
ORF3B	3B	Induce inflammatory response and inhibit the expression	58

		of IFN β	
E	E	Envelope protein pentamer	75
E	E-27nt	Envelope protein pentamer (minus 9 amino acids)	66
M	M	Membrane protein	222
ORF6	6	Antagonize STAT1 function and IFN signalling, and induce DNA synthesis	61
ORF7A	7A	Induce inflammatory response and apoptosis	121
ORF7B	7B	Induce inflammatory response	43
ORF7B	7B-truncated	Induce inflammatory response (with N terminus truncated)	20
ORF8	8	Induce apoptosis and DNA synthesis	121
N	N	Facilitate viral RNA packaging	419
ORF9B	9B	Induce apoptosis	98
ORF9Bwu	9Bwu	Unknown	73
ORF10wu	10wu	Unknown	38

STAR METHODS

RESOURCE AVAILABILITY

Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Frederick P. Roth (fritz.roth@utoronto.ca).

Materials Availability

Plasmids generated in this study have been deposited to Addgene (see Table S2 for links), available with a completed Materials Transfer Agreement.

Data and Code Availability

This study did not generate any unique datasets or code.

METHOD DETAILS

Synthesis of viral coding sequences

Based on the published annotation of the genome sequence of the HKU-SZ-005b (Chan et al., 2020) and Wuhan-Hu-1 (Wu et al., 2020) isolates of SARS-CoV-2, we requested the synthesis of viral coding sequences (GenScript, IDT), including termination codons and *attB* recombination sequences, with optimization of codon usage to reduce GC content and optimize expression in human and insect cells. A start codon is added to NSP2–16 to allow independent transcription, as they are prophetically cleaved from ORF1 post translation in host cells. ORF9Bwu, an alternative ORF within the N gene from the SARS-COV-2 (Wu et al., 2020), was subsequently amplified by Polymerase Chain Reaction (PCR) from the viral N gene with primers listed in Table S1.

Generation of Gateway-compatible viral coding sequence clone collections

Synthesized viral coding sequences were then incorporated into Gateway Entry plasmids: either pDONR207 (Invitrogen) or pDONR223 (Rual et al., 2004). To enable C-terminal fusion constructs, we also generated an equivalent set of Gateway-compatible clones without termination codons.. These clones were made by either PCR-amplifying the whole plasmid with primers that eliminated the stop codon, or by amplifying CDS regions from the first collection, using downstream primers with complementary regions that were internal to each stop codon, and which simultaneously incorporated the flanking sequences necessary for incorporation into a Gateway Entry plasmid (pDONR207, pDONR221 and pDONR223). We further expanded our collection to include clones containing N-terminal recognition sequence for nuclear inclusion protease from tobacco etch virus (TEV) to enable the *ad-hoc* removal of fusion tags. TEV sequences were incorporated by amplifying CDS regions from the first collection using forward primers containing TEV sequences and original reverse primers.

Each SARS-CoV-2 CDS bacterial clone was isolated from a single colony, and its inserted CDS was confirmed by full-length Sanger sequencing (TCAG DNA sequencing facility, Toronto, Canada). All clones with a pDONR221 or pDONR223 backbone were sequenced with M13F and M13R primers. Clones with a pDONR207 backbone were sequenced with customized forward and reverse primers. All primer sequences are available in Table S1.

Supplementary information

Table S1. Primers used for amplifying and sanger sequencing viral coding sequences.

Table S2. Clones in the genome-scale SARS-CoV-2 coding sequence collection, together with their nucleotide and amino acid lengths, coding sequence and direct links to Addgene.

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