

# 1 Title: Rapid reconstruction of SARS-CoV-2 using a synthetic genomics platform

2 Short title: In yeast genome engineering of RNA viruses

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## 27 Abstract

28 Reverse genetics has been an indispensable tool revolutionising our insights into viral  
 29 pathogenesis and vaccine development. Large RNA virus genomes, such as from  
 30 Coronaviruses, are cumbersome to clone and to manipulate in *E. coli* hosts due to size and  
 31 occasional instability<sup>1-3</sup>. Therefore, an alternative rapid and robust reverse genetics platform  
 32 for RNA viruses would benefit the research community. Here we show the full functionality  
 33 of a yeast-based synthetic genomics platform for the genetic reconstruction of diverse RNA  
 34 viruses, including members of the *Coronaviridae*, *Flaviviridae* and *Paramyxoviridae* families.  
 35 Viral subgenomic fragments were generated using viral isolates, cloned viral DNA, clinical  
 36 samples, or synthetic DNA, and reassembled in one step in *Saccharomyces cerevisiae* using  
 37 transformation associated recombination (TAR) cloning to maintain the genome as a yeast  
 38 artificial chromosome (YAC). T7-RNA polymerase has been used to generate infectious  
 39 RNA, which was then used to rescue viable virus. Based on this platform we have been able  
 40 to engineer and resurrect chemically-synthesized clones of the recent epidemic SARS-CoV-2<sup>4</sup>  
 41 in only a week after receipt of the synthetic DNA fragments. The technical advance we  
 42 describe here allows to rapidly responding to emerging viruses as it enables the generation  
 43 and functional characterization of evolving RNA virus variants - in real-time - during an  
 44 outbreak.

## 45 Main Text

46 Population growth, increased travel and climate change foster epidemic and pandemic threats  
 47 by (re)emerging RNA viruses. Although black swan events cannot be predicted, our medical  
 48 sector should be ready to use the latest advancements in science to combat emerging viruses.  
 49 Within the past decade we have seen outbreaks of Middle-East Respiratory Syndrome  
 50 coronavirus (MERS-CoV)<sup>5</sup>, ZIKA virus (ZIKV)<sup>6</sup>, Ebola virus<sup>7</sup>, and at the end of 2019,  
 51 SARS-CoV-2 that was first detected in Wuhan, Hubei province, China<sup>4</sup>, but meanwhile  
 52 spread throughout China and beyond. Until now SARS-CoV-2 isolates from China have not  
 53 been shared with health authorities and the scientific community, although they are urgently  
 54 needed to build up serological diagnostics, to develop and assess antivirals and vaccines, and  
 55 to establish appropriate *in vivo* models. Generation of SARS-CoV-2 from chemically  
 56 synthesized DNA could bypass the limited availability of virus isolates and would  
 57 furthermore allow genetic modifications and functional characterization of individual genes.  
 58 However, although the workhorse *E. coli* did prove very useful for cloning many viral  
 59 genomes, it has drawbacks for a number of emerging RNA viruses, including coronaviruses,  
 60 in terms of assembling and stably maintaining full-length molecular clones.

61 Synthetic genomics is a field that was fuelled by the efforts to create a bacterial cell that is  
 62 controlled by a synthetic genome<sup>8</sup>. Genome-wide reassembly of the ~1.1Mb *Mycoplasma*  
 63 genome was first attempted using *E. coli* as an intermediate host<sup>8</sup>, but the maintenance of  
 64 these 100-kbp DNA fragments appeared to be very difficult in this host. Therefore, the yeast  
 65 *Saccharomyces cerevisiae* was chosen to clone, assemble and even mutagenize entire  
 66 *Mycoplasma* genomes<sup>9,10</sup>. This ultimately led to the construction of the Mycoplasma JCVI  
 67 Syn3.0, a cell carrying the minimal set of genes allowing its autonomous replication in  
 68 auxotrophic medium<sup>11</sup>. The rationale for using a yeast cloning system is the ability of yeast to

69 recombine overlapping DNA fragments *in cellulo*, which led to the development of a  
70 technique called “transformation-associated recombination” (TAR) cloning<sup>12</sup>.

71 More recently, TAR cloning was successfully applied for the assembly, genetic engineering  
72 and rescue of large DNA viruses such as cytomegalovirus and herpes simplex virus 1<sup>13,14</sup>.

73 While large DNA viruses such as herpesviruses can be rescued by transfecting the entire viral  
74 DNA genome into eukaryotic cells, the rescue of RNA viruses requires the transcription of a  
75 viral genomic RNA from the cloned DNA template. For coronaviruses that belong to a family  
76 of positive-stranded RNA viruses termed *Coronaviridae*, the generation of full-length  
77 molecular clones has long been hampered by the extraordinary genome size (27-31 kb) and  
78 occasional instability of cloned DNA in *E. coli*. It took until 2000/2001 before unconventional  
79 approaches, such as cloning in low copy bacterial artificial chromosomes (BACs) or vaccinia  
80 virus, or cloning of subgenomic DNA fragments followed by *in-vitro* ligation, were  
81 successful<sup>1-3</sup>. However, each system has caveats leaving the generation of recombinant  
82 coronaviruses cumbersome. We therefore aimed to assess the use of the yeast *S. cerevisiae* to  
83 assemble and maintain genomes of diverse RNA viruses in order to establish a rapid, stable,  
84 and universal reverse genetics pipeline for RNA viruses.

85 In an effort to adopt a yeast-based reverse genetics platform to RNA viruses we first tested the  
86 Murine hepatitis virus strain A59 containing the gene for the green fluorescent protein (MHV-  
87 GFP), which is routinely used in our laboratory and has an established vaccinia virus-based  
88 reverse genetics platform<sup>15,16</sup>. The overall strategy is shown in Fig. 1a. Viral RNA was  
89 prepared from MHV-GFP-infected murine 17Cl-1 cells and used to amplify 7 overlapping  
90 DNA fragments by RT-PCR together spanning the MHV-GFP genome from nt 2024 to nt  
91 29672. Fragments containing the 5'- and 3'-termini were PCR-amplified from the vaccinia  
92 virus-cloned genome to include a T7-RNA polymerase promoter and a cleavage site (*PacI*)  
93 following the polyA sequence. Overlaps to the TAR vector pVC604 were included in the

94 primers used to amplify the 5'- and 3'-terminal fragments and the pVC604 backbone that was  
 95 also generated by PCR (Fig. 1b, Table 1, Extended Data Table 1, Extended Data Figure 1a).  
 96 All DNA fragments were used to transform *Saccharomyces cerevisiae* (strain VL6-48N) and  
 97 resulting colonies were screened for correct assembly of the YAC containing the MHV  
 98 genome by multiplex PCRs that covered the junctions of the different fragments. This screen  
 99 revealed that > 90 % of the clones tested were positive indicating a highly efficient assembly  
 100 in yeast (Extended Data Figure 1a). To rescue MHV-GFP we randomly chose two clones,  
 101 purified the YAC, linearized the plasmid using the restriction endonuclease *PacI* (Extended  
 102 Data Table 2) and subjected it to T7 RNA polymerase-based *in vitro* transcription to generate  
 103 capped viral genomic RNA. This RNA was transfected together with an *in vitro* transcribed  
 104 mRNA encoding the MHV nucleocapsid (N) protein into BHK-MHV-N cells as previously  
 105 described<sup>15</sup>. Transfected BHK-MHV-N cells were mixed with MHV-susceptible 17Cl-1 cells  
 106 and cytopathic effect (CPE), virus-induced syncytia, and GFP-expressing cells were readily  
 107 detectable for both clones within 48 hours, indicating successful recovery of infectious virus  
 108 (Fig. 1c). Finally, we assessed the replication kinetics of the recovered viruses, which were  
 109 indistinguishable compared to the parental MHV-GFP (Fig. 1d).

110 To address if the synthetic genomics platform can be applied to other coronaviruses, and if it  
 111 can be used for rapid mutagenesis, we used a molecular BAC clone of MERS-CoV<sup>17</sup>. We  
 112 PCR-amplified eight subgenomic overlapping DNA fragments covering the entire MERS-  
 113 CoV genome (Figure 2a, Extended Data Figure 1b, Extended Data Table 1). The 5'- and 3'-  
 114 terminal DNA fragments contained the T7-RNA polymerase promoter upstream the MERS-  
 115 CoV 5'-end and a restriction endonuclease cleavage site *MluI* downstream the polyA  
 116 sequence, and overlapping sequences with the TAR plasmid pVC604. To mutagenize the  
 117 MERS-CoV clone and introduce the GFP gene, fragment 7 was modified to overlap with the  
 118 PCR-amplified GFP gene. Following transformation almost all YAC clones were positive for  
 119 the junctions indicating a successful assembly (Figure 2a, Extended Data Figure 1b). Virus

rescue from cloned DNA was performed as described previously<sup>17</sup> and virus-induced CPE became apparent in VeroB4 cells inoculated with recovered recombinant MERS-CoV (Fig. 2b). Importantly, we could readily detect green fluorescent VeroB4 cells that had been infected with recovered MERS-CoV-GFP demonstrating that the synthetic genomics platform is suitable to genetically modify coronavirus genomes (Fig. 2b). As expected replication kinetics of recombinant MERS-CoV and MERS-CoV-GFP were slightly reduced compared to cell-culture-adapted MERS-CoV-EMC (Fig. 2c).

Having shown that the TAR cloning approach is robust and reproducible, we aimed to thoroughly evaluate the system concerning stability of cloned sequences, the range of applicability to other virus genomes, and if molecular clones can even be generated from clinical samples. The yeast clones containing YACs encoding MHV-GFP and MERS-CoV were passaged 15-17 times followed by re-sequencing. Results of this experiment were very supportive of the yeast-based platform (Extended Data Table 3), since the genomes could be stably maintained and no mutations arose. We then embarked on cloning a number of other coronaviruses (HCoV-229E<sup>2</sup>, HCoV-HKU1; GenBank: NC\_006577, MERS-CoV-Riyadh-1734-2015; GenBank: MN481979) and viruses of other families, such as ZIKV (family *Flaviviridae*, GenBank: KX377337) and human respiratory-syncytial-virus (hRSV; family *Paramyxoviridae*) (Table 1) that are known to be difficult to clone and stably maintain in *E. coli*. As shown in Extended data Figure 1d-h, we followed the same strategy to prepare the genome termini in cloned plasmids to include genetic elements that are later needed for virus rescue (e.g. T7-RNA polymerase promoter, cleavages sites or ribozyme elements) and produced a set of overlapping DNA fragments covering the rest of the genome. This strategy proved to be successful irrespective of the source of virus or nucleic acid template or the number of DNA fragments that were used for the one-step assembly in yeast. Of note, we were also able to clone hRSV-B, without any prior information on the virus genotype, directly from a clinical sample (nasopharyngeal aspirate) by designing RSV consensus primers to

amplify 4 overlapping DNA fragments (sequence submitted to GenBank). Collectively, these results demonstrate the synthetic genomics platform provides the technical advance to rapidly generate molecular clones of diverse RNA viruses by using viral isolates, cloned DNA, synthetic DNA, and even clinical samples as starting material.

The appearance of a novel CoV in China at the end of 2019 prompted us to go a step further and to test the applicability of our synthetic genomics platform to reconstruct the virus based on its released genome sequence and chemically synthesized DNA fragments. Notably, at the time when the first genome sequences of SARS-CoV-2 were released (Jan 10/11<sup>th</sup>, 2020) no virus isolate has been made available and it took until the end of January 2020 when successful isolation of SARS-CoV-2 was reported from patients in Australia (<https://www.the-scientist.com/news-opinion/australian-lab-cultures-new-coronavirus-as-infections-climb-67031>). We fragmented the genome into 12 subgenomic DNA fragments ranging in size between 0.5-3.4 kbp (Fig. 3a, Extended Data Figure 1i, Extended Data Tables 1, 4). In parallel, we also aimed to generate a SARS-CoV-2 expressing GFP that could be valuable to facilitate the establishment of serological diagnostics (e.g. virus neutralization assay) and detection in cell culture. To do this we divided fragment 11 into 3 sub-fragments (Fig. 3a, Extended Data Figure 1i, Extended Data Tables 1, 4) to include the GFP sequence, which was inserted in frame within the ORF7a. We noticed at the 5' end of the reported SARS-CoV-2 sequence that nucleotides 3-5 (5'-AUUAAAGG; Genbank MN996528.1) were different compared to the closely related SARS-CoV (5'-AUAUUAGG; GenBank AY291315) and to the even more closely related bat SARS-related CoVs ZXC21 and ZC45 (5'-AUAUUAGG)<sup>18</sup>, and that for all three the 5'-end has been experimentally determined by 5'-RACE<sup>4,18,19</sup>. Therefore, we designed three 5'-end versions of fragment 1 containing the reported SARS-CoV-2 sequence (fragment 1.3; 5'-AUUAAAGG), a version modified by 3 nucleotides (fragment 1.1; 5'-AUAUUAGG), and a version containing the 124 5'-terminal nucleotides spanning the first four 5'-terminal stem-loop structures of SARS-CoV (fragment



1.2; Fig. 3b). Notably, nucleotide differences between SARS-CoV-2 and SARS-CoV within this region are in agreement with the predicted RNA secondary structures (Fig. 3b).

We ordered the synthetic DNA constructs on January 14<sup>th</sup> (Fig. 3d, Extended data Table 4, Extended Data Document 1), and until February 4<sup>th</sup> 12 of 14 constructs were delivered as sequence-confirmed plasmids. However, fragments 5 and 7 turned out to be problematic to clone in *E. coli* and could not be delivered. Since we received at the same time SARS-CoV-2 viral RNA obtained from an isolate of a Munich patient (SARS-CoV-2/München-1.1/2020/929), we decided to amplify the regions of fragments 5 and 7 by RT-PCR. TAR cloning was immediately initiated, and for all 6 SARS-CoV-2 constructs we obtained correctly assembled molecular clones (3 versions of the 5'-end and each version with and without GFP; Extended Data Fig. 2). Since sequence verification was not possible within this short time frame, we randomly selected two clones for each construct for rescuing recombinant rSARS-CoV-2 and rSARS-CoV-2-GFP. YAC DNA were isolated, *in vitro* transcribed and the resulting RNAs electroporated together with an mRNA encoding the SARS-CoV-2 N protein into BHK-21 and, in parallel, into BHK-SARS-N cells expressing the SARS-CoV N protein<sup>20</sup>. The BHK-21 and BHK-SARS-N cells were seeded over VeroE6 cells and at two days post electroporation we observed green fluorescent signals in cells that received the GFP-encoding SARS-CoV-2 RNAs. Indeed supernatants transferred to fresh VeroE6 cells contained infectious recombinant viruses for almost all recombinant rSARS-CoV-2 and rSARS-CoV-GFP constructs (Extended Data Fig. 2). As shown in Figure 3c for SARS-CoV-2 clones 1.1, 2.2, and 3.1, plaques were detectable on VeroE6 cells that were inoculated with supernatants containing the RNA-electroporated cells, demonstrating the infectious virus has been recovered for recombinant rSARS-CoV-2 and for the variants containing modified 5'-termini. Similarly, we could detect green fluorescent cells on VeroE6 monolayers that were inoculated with supernatants from RNA-electroporated cells of rSARS-CoV-2-GFP clones 4.1, 5.2 and 6.1, demonstrating that recombinant GFP-expressing viruses

have been recovered for all 5'-end variants. This result demonstrates the full functionality of the SARS-CoV-2 reverse genetics system and also shows that the 5'-ends of SARS-CoV and bat SARS-related CoVs ZXC21 and ZC45 are compatible with the replication machinery of SARS-CoV-2.

We expect that the fast, robust and versatile synthetic genomics platform we describe here will provide new insights into the molecular biology and pathogenesis of a number of emerging RNA viruses and will specifically facilitate the molecular characterization of the novel coronavirus SARS-CoV-2. Although homologous recombination in yeast has already been used for the generation of a number of molecular virus clones in the past<sup>13,14,21</sup>, we present here a thorough evaluation of the feasibility of this approach to rapidly generate full-length cDNAs for large RNA viruses that have a known history of instability in *E. coli*. We show that one main advantage of the TAR cloning system is that the viral genomes can be fragmented to at least 14 overlapping fragments and re-assembled with remarkable efficacy (usually >90% of the clones are correct). This allowed us to complete the cloning and rescue of rSARS-CoV-2 and rSARS-CoV-2-GFP within one week. It should also be noted that we see considerable potential to reduce the time of DNA synthesis. Currently, synthetic DNA fragments are still routinely cloned in *E. coli*, which turned out to be problematic for SARS-CoV-2 fragments 5 and 7. We could however use 4 and 3 shorter dsDNA parts of fragments 5 and 7, respectively, that we received, to assemble the full-length fragments 5 and 7 by TAR cloning which is an additional proof of the superior cloning efficiency of our yeast system versus *E. coli*-based systems. This allowed us to generate a molecular clone of SARS-CoV-2 by using exclusively chemically synthesized DNA, and the rescue of this version is currently in progress.

The appearance of SARS-CoV-2 emphasizes the need for preparedness to rapidly respond to emerging virus threats. While the availability of SARS-CoV-2 isolates is still limited, full

virus genome sequences were disclosed very early during the outbreak. The rapidity of our synthetic genomics approach to generate SARS-CoV-2 and the applicability to other emerging RNA viruses make this system an attractive alternative to provide infectious virus to health authorities and diagnostic laboratories without the need to have access to clinical samples. Finally, as the SARS-CoV-2 outbreak is ongoing, we expect to see sequence variations and possibly phenotypic changes of evolving SARS-CoV-2 in the human host. With the synthetic genomics platform it is now possible to introduce such sequence variations into the infectious clone and to functionally characterize SARS-CoV-2 evolution in real-time.

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## Authors contributions

VT and JJ conceived the study. TT, NE and FL performed most of the experiments. PV, HS, JP, SSt, MH, AK, MG, LL, LH, MW, SP, DH, VC, SC, SSch, DM, DN, MM, CD, RD, did experimental work and/or provided essential experimental systems and reagents. TT, NE, FL, HS, JK, and RD performed sequencing including computational analyses. VT, JJ, TT, FL, NE wrote the manuscript and made the figures. All authors read and approved the final manuscript.

## Ethical statement

The authors are aware that this work contains aspects of Dual Use Research of Concern (DURC). The benefits were carefully balanced against the risks and the benefits outweigh the risks. Permission to generate and work with recombinant SARS-CoV-2 and SARS-CoV-2-

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### 323 [Competing interests](#)

324 The authors declare no competing interests

### 325 [Additional information](#)

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## Figure Legends

### **Figure 1. Application of yeast-based TAR cloning to generate viral cDNA clones and the recovery of recombinant MHV-GFP.**

**a**, Schematic representation of the general workflow of TAR cloning and virus rescue. In-yeast assembly require the delivery of overlapping DNA fragments covering the viral genome and a TAR vector into yeast in one transformation event. Transformed DNA fragments are joined by homologous recombination in yeast generating a YAC-cloned full-length viral genomic cDNA. The *in vitro* production of infectious capped viral RNA starts with the isolation of the YAC, followed by plasmid linearization to provide DNA template for run-off T7 polymerase-based transcription. Virus rescue starts with electroporation of BHK-21 and/or BHK-21 cells expressing the corresponding coronavirus N protein, followed by co-cultivation of electroporated cells with susceptible cell line for virus production and amplification. **b**, Schematic representation of MHV-GFP genome organisation with 9 viral subgenomic overlapping cDNA fragments used for TAR cloning. pT7, T7 RNA polymerase promoter; UTR, untranslated region; An, poly (A) tail; J represents assembly junction between two overlapping DNA fragments. **c**, Recovery of infectious recombinant MHV-GFP. Cell culture supernatants containing viruses produced after virus rescue from two MHV-GFP YAC clones (Clone1, Clone 2) were used to infect 17Cl-1 cells. At 48 hours post-infection, infected cells were visualised for GFP expression (left panels) and by bright field microscopy (right panels). Mock represents 17Cl-1 cells inoculated with supernatant from BHK-MHV-N cells electroporated without viral RNAs. **d**, Replication kinetics of parental MHV-GFP and recombinant MHV-GFP Clone 1 and Clone 2. L929 cells were infected at an MOI of 0.1. Cell culture supernatants were harvested at indicated time points post-infection and titrated by plaque assay. Data are representative of three independent experiments with two replicates per

virus in each experiment. Error bars show standard deviation (SD) from the mean. pfu/ml, plaque forming units per millilitre.

**Figure 2. Generation of viral cDNA clones and recovery of recombinant MERS-CoV and MERS-CoV-GFP.**

**a**, Schematic representation of MERS-CoV (upper panel) and MERS-CoV-GFP (lower panel) genome organization with 8 and 10 viral subgenomic overlapping cDNA fragments, respectively, used for TAR cloning. pT7, T7 RNA polymerase promoter; UTR, untranslated region; An, poly (A) tail; J represents assembly junction between two overlapping DNA fragments. **b**, Rescue of recombinant MERS-CoV and MERS-CoV-GFP. Following delivery of viral RNAs into BHK-21 cells via electroporation the cells were co-cultivated with VeroB4 cells and supernatants containing recombinant viruses produced were used to infect new VeroB4 cells. Infected cells were visualised by bright field microscopy for recombinant MERS-CoV (upper panel; 5 days post infection), and for GFP expression for recombinant MERS-CoV-GFP (lower panel, 3 days post infection). Mock represents VeroB4 cells inoculated with supernatant from BHK-21 cells electroporated without viral RNAs. **c**, Replication kinetics of the MERS-CoV-EMC (laboratory adapted isolate), rMERS-CoV and rMERS-CoV-GFP. VeroB4 cells were infected at an MOI of 0.01. Cell culture supernatants were harvested at indicated time points post infection and titrated by TCID<sub>50</sub> assay. Data are representative of three independent experiments with two replicates per virus in each experiment. Error bars show standard deviation (SD) from the mean. TCID<sub>50</sub>/ml, 50% tissue culture infectious dose per millilitre.

**Figure 3. Generation of viral cDNA clones and recovery of recombinant SARS-CoV-2 and SARS-CoV-2-GFP. a**, Schematic representation of SARS-CoV-2 (upper panel) and

SARS-CoV-2-GFP (lower panel) genome organisation with 12 and 14 viral subgenomic overlapping cDNA fragments, respectively, used for TAR cloning. pT7, T7 RNA polymerase promoter; UTR, untranslated region; An, poly (A) tail; J represents assembly junction between two overlapping DNA fragments. **b**, Representation of predicted RNA stem-loop (SL) secondary structures formed in the 5'-UTR of SARS-CoV-2. The SARS-CoV-2 RNA secondary structures were manually adjusted based on RNA structure predictions by Chan et al.<sup>22</sup>. Black letters and numbering represents the SARS-CoV-2 5'-terminal sequence. Red letters depict nucleotides that are different within the SARS-CoV 5'-terminal sequence ( “-“ indicates nucleotide deletion in SARS-CoV versus SARS-CoV-2). N20 indicates 20 nucleotides. SL structures 1, 2, and 4 are shown according to Chan *et al.*<sup>22</sup>. Note that recombinant virus construct 1 (rSARS-CoV-2 clone 1) and construct 4 (rSARS-CoV-GFP clone 4) contain the SARS-CoV-2 5'-terminus with nucleotides 3-5 (UAA) exchanged by nucleotides 3-5 (AUU) of SARS-CoV; recombinant virus construct 2 (rSARS-CoV-2 clone 2) and construct 5 (rSARS-CoV-2-GFP clone 5) contain a 5'-terminus with the first 124 nucleotides from SARS-CoV; and recombinant virus construct 3 (rSARS-CoV-2 clone 3) and construct 6 (rSARS-CoV-GFP clone 6) contain the authentic SARS-CoV-2 5'-terminus. **c**, Rescue of recombinant rSARS-CoV-2 and rSARS-CoV-2-GFP. The experimental setting is illustrated in the upper panel. Two clones of each construct 1-6 were used to prepare YAC DNAs and *in-vitro* transcribed viral genome RNAs were electroporated into BHK-21 cells and/or BHK-SARS-N together with an mRNA encoding the SARS-CoV-2 N protein. Electroporated cells were co-cultivated with VeroE6 cells in a T25 flask and cell culture supernatants were transferred to 12-well plates to perform plaque assay for rSARS-CoV-2 constructs 1-3 and fluorescence microscopy for rSARS-CoV-2-GFP constructs 4-6. Plaque assays are shown in the lower panel for SARS-CoV-2 clones 1.1, 2.2, and 3.1. VeroE6 cells were infected with  $10^{-1}$ ,  $10^{-2}$ , and  $10^{-3}$  dilutions of 1 ml supernatant from the individual rescue experiments and are compared to non-infected cells (Mock). The expression of GFP in SARS-

CoV-2-GFP infected VeroE6 cells is shown for clones 4.1, 5.2, and 6.1. Mock represents non-infected VeroE6 cells. **d**, Summary of the timeline of events of the SARS-CoV-2 outbreak and the work leading to the reconstruction and recovery of rSARS-CoV-2.

# **Extended Data Figure 1**

**a**, Schematic representation of the MHV-GFP genome organisation (upper panel) and nine viral subgenomic cDNA fragments (F1-9) used for TAR cloning. Viral open reading frames (ORFs), the ORF for GFP and sequence elements at the 5'- and 3'-untranslated regions (UTRs) are indicated. Primers used to generate the fragments are listed in Extended Data Table 1. J2-9 represent the junctions, i.e. overlapping regions, between the subgenomic cDNA fragments. J1 and J10 represent junctions with the TAR vector. Gel images show results from two multiplex PCRs that were designed to detect the presence of correctly recombined junctions J1-10, confirming the proper assembly of the YAC containing the viral genome in 12 out of 12 clones. kb, kilobases; pT7, T7 RNA polymerase promoter; An, poly (A) tail.

**b**, Schematic representation of the MERS-CoV genome organisation (upper panel) and eight viral subgenomic cDNA fragments (F1-8) used for TAR cloning. Viral open reading frames (ORFs) and sequence elements at the 5'- and 3'-untranslated regions (UTRs) are indicated. Primers used to generate the fragments are listed in Extended Data Table 1. J2-8 represent the junctions, i.e. overlapping regions, between the subgenomic cDNA fragments. J1 and J9 represent junctions with the TAR vector. Gel image shows results from a multiplex PCR that was designed to detect the presence of correctly recombined junctions J1-9, confirming the proper assembly of the YAC containing the viral genome in 6 out of 8 clones. kb, kilobases; pT7, T7 RNA polymerase promoter; An, poly (A) tail.

**c**, Schematic representation of the MERS-CoV-GFP genome organisation (upper panel) and ten viral subgenomic cDNA fragments (F1-10 and F13) used for TAR cloning. Viral open reading frames (ORFs), ORF for GFP and sequence elements at the 5'- and 3'-untranslated regions (UTRs) are indicated. Primers used to generate the fragments are listed in Extended Data Table 1. J2-8 and J10-11 represent the junctions, i.e. overlapping regions, between the subgenomic cDNA fragments. J1 and J9 represent junctions with the TAR vector. Gel images show results from two multiplex PCRs that were designed to detect the presence of correctly recombined junctions J1-11, confirming the proper assembly of the YAC containing the viral genome in 5 out of 8 clones. kb, kilobases; pT7, T7 RNA polymerase promoter; An, poly (A) tail.

**d**, Schematic representation of the HCoV-229E genome organisation (upper panel) and thirteen viral subgenomic cDNA fragments (F1-13) used for TAR cloning. Viral open reading frames (ORFs) and sequence elements at the 5'- and 3'-untranslated regions (UTRs) are indicated. Primers used to generate the fragments are listed in Extended Data Table 1. J2-13 represent the junctions, i.e. overlapping regions, between the subgenomic cDNA fragments. J1 and J14 represent junctions with the TAR vector. Gel images show results from two multiplex PCRs that were designed to detect the presence of correctly recombined junctions J1-14, confirming the proper assembly of the YAC containing the viral genome in 7 out of 10 clones. kb, kilobases; pT7, T7 RNA polymerase promoter; An, poly (A) tail.

**d**, Schematic representation of the HCoV-HKU1 genome organisation (upper panel) and eleven viral subgenomic cDNA fragments (F1-11) used for TAR cloning. Viral open reading frames (ORFs) and sequence elements at the 5'- and 3'-untranslated regions (UTRs) are indicated. Primers used to generate the fragments are listed in Extended Data Table 1. J2-11

represent the junctions, i.e. overlapping regions, between the subgenomic cDNA fragments.

J1 and J12 represent junctions with the TAR vector. Gel images show results from two multiplex PCRs that were designed to detect the presence of correctly recombined junctions J1-12, confirming the proper assembly of the YAC containing the viral genome in 14 out of 14 clones. kb, kilobases; pT7, T7 RNA polymerase promoter; An, poly (A) tail.

**e**, Schematic representation of the MERS-CoV-Riadh-1734-2015 genome organisation (upper panel) and eight viral subgenomic cDNA fragments (F1-8) used for TAR cloning. Viral open reading frames (ORFs) and sequence elements at the 5'- and 3'-untranslated regions (UTRs) are indicated. Primers used to generate the fragments are listed in Extended Data Table 1. J2-8 represent the junctions, i.e. overlapping regions, between the subgenomic cDNA fragments. J1 and J9 represent junctions with the TAR vector. Gel images show results from a multiplex PCR and a simplex PCR, both of which were designed to detect the presence of correctly recombined junctions J1-9, confirming the proper assembly of the YAC containing the viral genome in 5 out of 8 clones. kb, kilobases; pT7, T7 RNA polymerase promoter; An, poly (A) tail.

**f**, Schematic representation of the ZIKA virus genome organisation (upper panel) and six viral subgenomic cDNA fragments (F1-6) used for TAR cloning. Viral open reading frames (ORFs) and sequence elements at the 5'- and 3'-untranslated regions (UTRs) are indicated. Primers used to generate the fragments are listed in Extended Data Table 1. J2-5 represent the junctions, i.e. overlapping regions, between the subgenomic cDNA fragments. J1 and J6 represent junctions with the TAR vector. Gel image shows results from a multiplex PCR that was designed to detect the presence of correctly recombined junctions J1-6, confirming the proper assembly of the YAC containing the viral genome in 3 out of 15 clones. kb, kilobases; pT7, T7 RNA polymerase promoter; HDR, hepatitis delta virus ribozyme.

477

478 **g**, Schematic representation of the human RSV-B virus genome organisation (upper panel)  
 479 and six viral subgenomic cDNA fragments (F1-6) used for TAR cloning. Viral open reading  
 480 frames (ORFs) and sequence elements at the 5'- and 3'-untranslated regions (UTRs) are  
 481 indicated. Primers used to generate the fragments are listed in Extended Data Table 1. J1-5  
 482 represent the junctions, i.e. overlapping regions, between the subgenomic cDNA fragments.  
 483 J1 and J5 contain junctions with the TAR vector. Gel image shows results from a multiplex  
 484 PCR that was designed to detect the presence of correctly recombined junctions J1-5,  
 485 confirming the proper assembly of the YAC containing the viral genome in 7 out of 8 clones.  
 486 kb, kilobases; pT7, T7 RNA polymerase promoter; HHrb, hammerhead ribozyme, Rb-T7ter,  
 487 Ribozyme and T7 RNA polymerase terminator sequence.

488

489 **h**, Schematic representation of the SARS-CoV-2 genome organisation (upper panel) and  
 490 twelve viral subgenomic cDNA fragments (F1-12) used for TAR cloning. Viral open reading  
 491 frames (ORFs) and sequence elements at the 5'- and 3'-untranslated regions (UTRs) are  
 492 indicated. Primers used to generate the fragments are listed in Extended Data Table 1. J2-12  
 493 represent the junctions, i.e. overlapping regions, between the subgenomic cDNA fragments.  
 494 J1 and J13 represent junctions with the TAR vector. Gel images show results from two  
 495 multiplex PCRs that were designed to detect the presence of correctly recombined junctions  
 496 J1-13, confirming the proper assembly of the YAC containing the viral genome in 6 out of 6  
 497 clones. kb, kilobases; pT7, T7 RNA polymerase promoter; An, poly (A) tail.

498

499 **i**, Schematic representation of the SARS-CoV-2 genome organisation (upper panel) and  
 500 fourteen viral subgenomic cDNA fragments (F1-10 and F12-15) used for TAR cloning. Viral  
 501 open reading frames (ORFs), ORF for GFP and sequence elements at the 5'- and 3'-

untranslated regions (UTRs) are indicated. Primers used to generate the fragments are listed in Extended Data Table 1. J2-12 and J14 represent the junctions, i.e. overlapping regions, between the subgenomic cDNA fragments. J1 and J13 represent junctions with the TAR vector. Gel images show results from two multiplex PCRs and a simplex PCR, all of which were designed to detect the presence of correctly recombined junctions J1-13 and J14 respectively, confirming the proper assembly of the YAC containing the viral genome in 6 out of 6 clones. kb, kilobases; pT7, T7 RNA polymerase promoter; An, poly (A) tail.

## Extended Data Figure 2

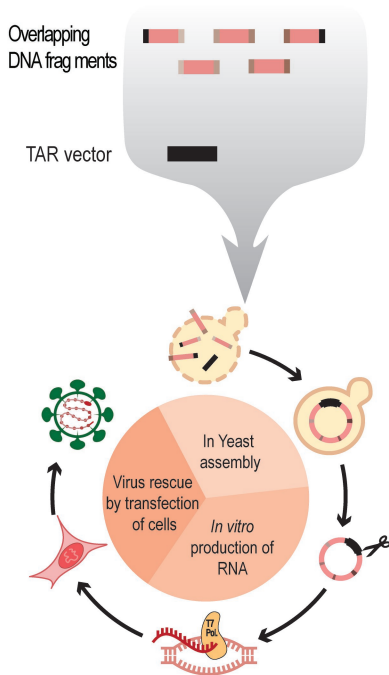
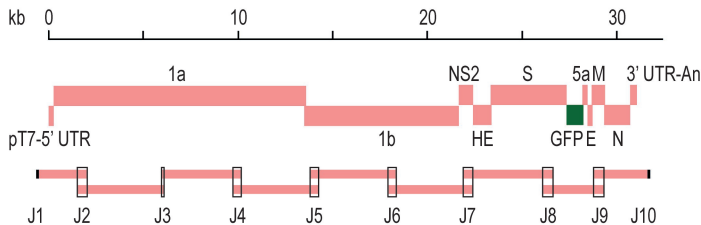
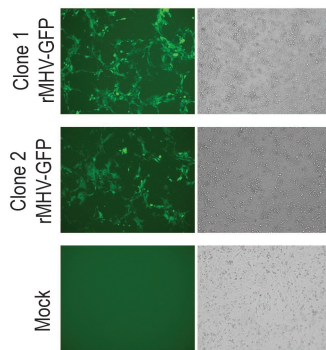
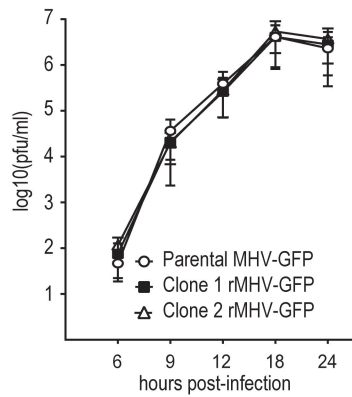
**a**, Six constructs were initially designed based on 3 different 5'-UTR regions. These regions contain i) a modified sequence of the 5'-UTR region of the SARS-CoV-2 (5'- ATAUUAGG; 5'-terminal 8 nucleotides are shown, modified nucleotides underlined) were nucleotides 3-5 (UAA) of SARS-CoV-2 were changed to AUU to match the SARS-CoV nucleotides 3-5 (synthetic fragment 1.1 for constructs 1 and 4), ii) a SARS-CoV-2 5'-terminus with the first 124 nucleotides exchanged by the corresponding 5'-terminal sequence of SARS-CoV (synthetic fragment 1.2; construct 2 and 5) and iii) the reported sequence of the SARS-CoV-2 (GISAID: EPI\_ISL\_402119) (synthetic fragment 1.3; constructs 3 and 6). After yeast transformation, 10 colonies were randomly picked for each of the six constructs and all the junctions bridging the overlapping fragments were verified by multiplex PCR. For each construct, two clones (x.1 and x.2) were randomly selected and YAC DNAs were isolated (12 clones in total). RNAs were generated by *in vitro* transcription and electroporated together with an mRNA encoding the SARS-CoV-2 N protein either into BHK-21 cells (12 clones) or BHK-SARS-N cells (cells expressing the SARS-CoV N protein) (6 clones). Electroporated cells were then co-cultivated with susceptible VeroE6 cells to rescue the recombinant viruses. P.0 supernatants were collected at different time points after electroporation (from 2 dpe to 5

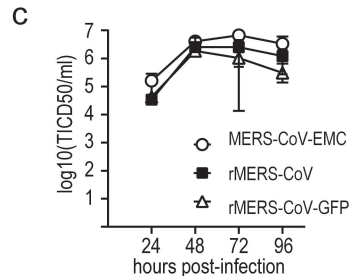
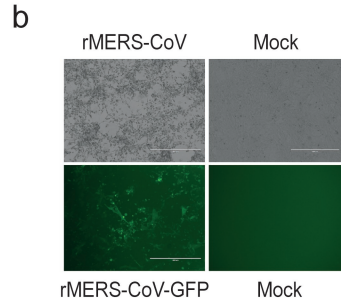
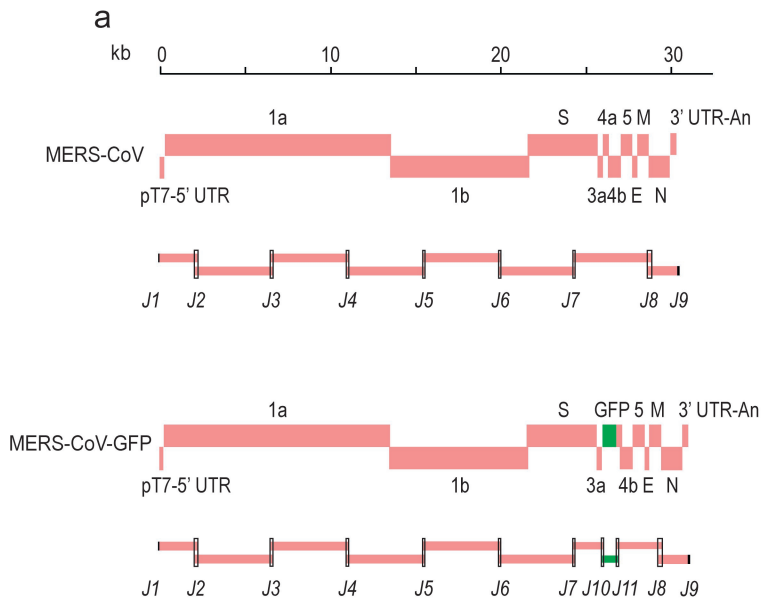


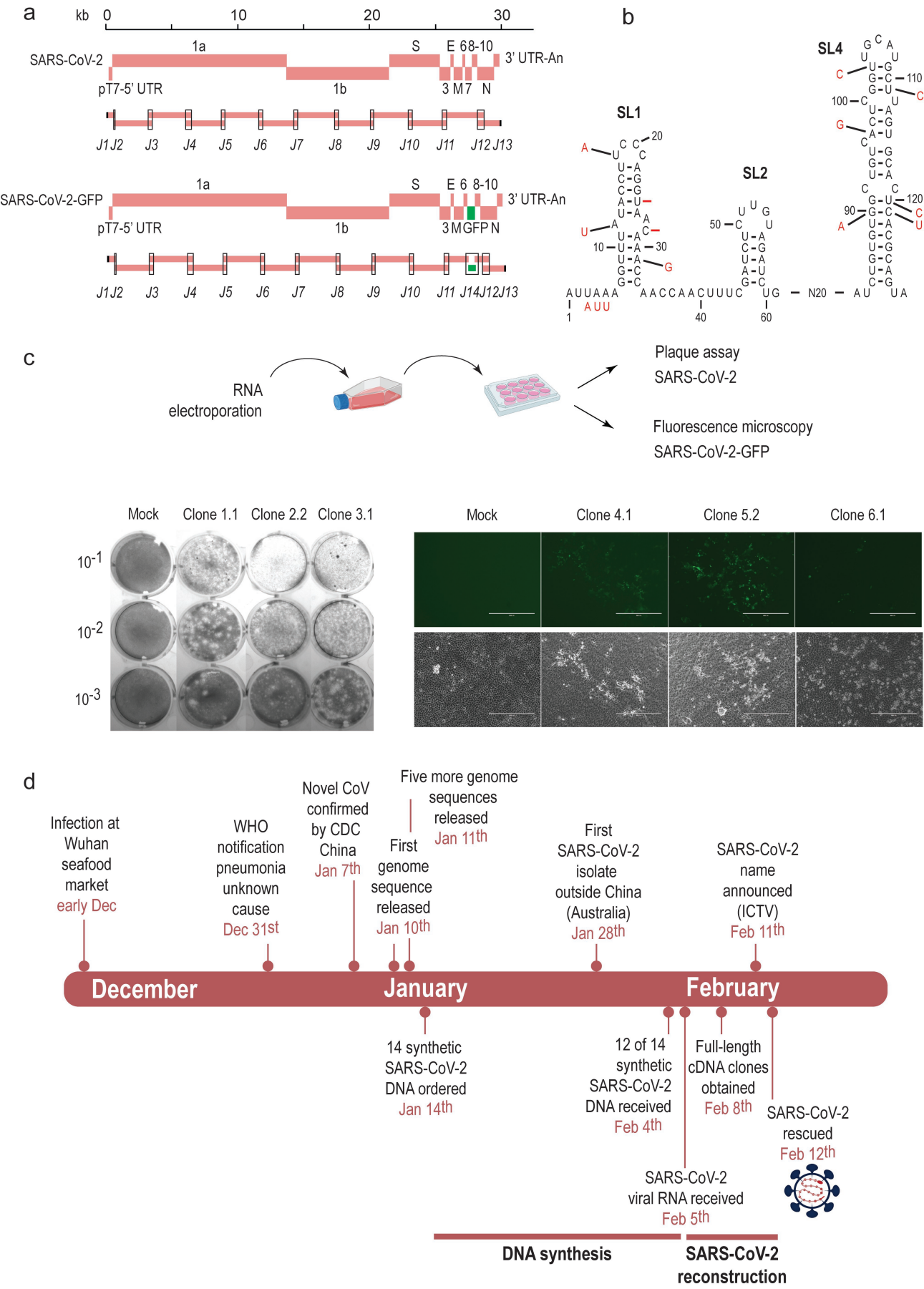
527 dpe) and transferred to VeroE6 cells to generate P.1 virus stocks, and in parallel to  
 528 demonstrate presence of infectious virus by plaque assay (virus clones that do not encode  
 529 GFP) or fluorescence microscopy (GFP-encoding virus clones). YAC, yeast artificial  
 530 chromosome; h.p.e, hours post-electroporation; d.p.e, days post-electroporation; CPE,  
 531 cytopathic effects; GFP, green fluorescent protein; pfu/ml, plaque forming units per millilitre.

532

533

**a****b****c****d**





**Table 1: RNA virus genomes cloned using the synthetic genomics platform**

| <b>Virus</b>              | <b>Family</b>   | <b>RNA type</b> | <b>Size (kbp)</b> | <b>Template</b>                                          | <b>Fragment generation</b>         | <b>Number of assembled fragments*</b> | <b>Virus rescue</b> |
|---------------------------|-----------------|-----------------|-------------------|----------------------------------------------------------|------------------------------------|---------------------------------------|---------------------|
| MHV-GFP                   | Coronaviridae   | (+)RNA          | 31.9              | Viral RNA (laboratory virus); DNA clone (vaccinia virus) | RT-PCR/PCR                         | 9                                     | yes                 |
| MERS-CoV                  | Coronaviridae   | (+)RNA          | 30.1              | DNA clone (BAC)                                          | PCR                                | 8                                     | yes                 |
| MERS-CoV-GFP              | Coronaviridae   | (+)RNA          | 30.7              | DNA clone (BAC)<br>GFP plasmid DNA                       | PCR                                | 10                                    | yes                 |
| HCoV-229E                 | Coronaviridae   | (+)RNA          | 27.3              | Viral RNA (laboratory virus); DNA clone (vaccinia virus) | RT-PCR/PCR                         | 13                                    | in progress         |
| HCoV-HKU1                 | Coronaviridae   | (+)RNA          | 29.9              | Synthetic DNA<br>viral RNA (virus isolate)               | PCR/RT-PCR                         | 11                                    | in progress         |
| MERS-CoV-Riyadh-1734-2015 | Coronaviridae   | (+)RNA          | (30)              | Viral RNA (virus isolate)                                | RT-PCR                             | 8                                     | in progress         |
| ZIKA virus                | Flaviviridae    | (+)RNA          | 10.8              | Viral RNA (isolate)                                      | RT-PCR                             | 6                                     | in progress         |
| Human RSV-B               | Paramyxoviridae | (-)RNA          | 15                | Clinical sample                                          | RT-PCR                             | 4                                     | in progress         |
| SARS-CoV-2                | Coronaviridae   | (+)RNA          | 30                | Synthetic DNA/viral RNA                                  | plasmid DNA restriction/ RT-PCR    | 12                                    | yes                 |
| SARS-CoV-2-GFP            | Coronaviridae   | (+)RNA          | 30.5              | Synthetic DNA/viral RNA                                  | plasmid DNA restriction/RT-PCR/PCR | 14                                    | yes                 |

\*excluding the TAR vector fragment