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Construction of yellow fever virus subgenomic replicons by yeast-based homologous recombination cloning technique

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ABSTRACT
RNA replicon derived from Flavivirus genome is a valuable tool for studying viral replication independent of virion assembly and maturation, besides being a great potential for heterologous gene expression. In this study we described the construction of subgenomic replicons of yellow fever virus by yeast-based homologous recombination technique. The plasmid containing the yellow fever 17D strain replicon (pBSC-repYFV-17D), previously characterized, was handled to heterologous expression of the green fluorescent protein (repYFV-17D-GFP) and firefly luciferase (repYFV-17D-Luc) reporter genes. Both replicons were constructed by homologous recombination between the linearized vector pBSC-repYFV-17D and the PCR product containing homologous 25 nucleotides ends incorporated into PCR primers. The genomic organization of these constructs is similar to repYFV-17D, but with insertion of the reporter gene between the remaining 63 N-terminal nucleotides of the capsid protein and 72 C-terminal nucleotides of the E protein. The replicons repYFV-17D-GFP and repYFV-17D-Luc showed efficient replication and expression of the reporter genes. The yeast-based homologous recombination technique used in this study proved to be applicable for manipulation of the yellow fever virus genome in order to construct subgenomic replicons.

Key words: cloning technique, homologous recombination, replicon, reporter gene, yellow fever virus.

INTRODUCTION
The yellow fever virus (YFV) is a mosquito-borne agent transmitted mainly by species of the genera Aedes and Haemagogus that belongs to the Flavivirus genus of the Flaviviridae family. The YFV is the prototype Flavivirus, a genus of enveloped, single stranded, positive-sense RNA virus that is transmitted via arthropod vectors and include important human and veterinary pathogens as dengue virus, Japanese encephalitis and tick borne encephalitis virus (TBEV). The virus is endemic in tropical South America and sub-Saharan Africa and remains a serious threat to those countries. There is no antiviral therapy and vaccination with the attenuated YFV-17D strain has been the major strategy for yellow fever control (Gubler et al. 2007).
The virus genome consists of a positive-polarity, single-stranded RNA of approximately 11 Kb, encoding a single large polyprotein that is co- and post-translationally processed by viral and cell proteases to yield three structural proteins (C, prM and E) and seven non-structural (NS) proteins (NS1, NS2A, NS2B, NS3, NS4A, NS4B and NS5) from the N-terminus to the C-terminus, respectively (Jones et al. 2005, Kümmerer 2006).

Genetics analysis of positive-strand RNA viruses has been greatly facilitated by the use of recombinant DNA technology. In recent years it has been described the successful construction of full-length cDNA clones for several members of the flavivirus genus including YFV, dengue virus, West Nile virus and Kunjin virus (KUNV), among others (Bredenbeek et al. 2003, Imoto and Konish 2007, Khromykh and Westaway 1994, Polo et al. 1997, Puri et al. 2000, Sumiyoshi et al. 1992, Yamshchikov et al. 2005).

In order to extend a range of experimental tools in studies on flavivirus replication a self-replicating RNA (subgenomic replicon) derived of KUNV, a close relative of West Nile virus, was constructed (Khromykh and Westaway 1997). Thereafter, replicons were described for several flaviviruses including DENV-2 (Ng et al. 2007), DENV-3 (Mosimann et al. 2010), YFV (Jones et al. 2005), WNV (Widman et al. 2008) and TBEV (Kofler et al. 2004).

Subgenomic replicons of positive-stranded RNA viruses contain all of genetic elements needed to amplify themselves in susceptible host cells, but lack some or all of the genes coding structural proteins (Jones et al. 2005). Replicons of flaviviruses have been constructed by the introduction of in-frame deletions removing the coding regions for proteins prM and E and in some cases also the majority of C protein (Gehrke et al. 2003). Consequently, these RNAs are replicated in cells, but are not packaged into viral particles. Replicons have proven to be valuable tool for studying replication independently of virion assembly and maturation (Khromykh and Westaway 1997). Moreover, they have great potential as molecular tools for heterologous gene expression (Gehrke et al. 2003, Jones et al. 2005, Varnavski and Khromykh 1999).

Multiple strategies have been used for the construction of subgenomic replicons. However, these strategies seem to be cumbersome to perform because so many steps are required. The observation that linear DNA fragments can efficiently stimulate recombination in Saccharomyces cerevisiae has led to the rapid development of powerful methods for DNA manipulation in yeast (Oldenburg et al. 1997). One of these powerful methodologies is a yeast-based homologous recombination cloning technique (Shanks et al. 2009). This technique is based on the ability of yeast cells to recombine sequences similar during the natural event of homologous recombination that occurs to repair DNA damage during replication (Kuzminov 2001, Orr-Weaver et al. 1981). In this methodology, a DNA fragment containing homologous ends sequences with the vector can be directly cloned using in vivo recombination in the linearized vector (Gibson 2009). The advantages of yeast recombination are that multiple pieces of DNA can be sewn together seamlessly without multiple rounds of amplification, and without the need for ligation in vitro or the use of restriction sites. This method is efficient, robust and simple to perform (Gibson 2009, Shanks et al. 2009).


In the the present report we use the homologous recombination in yeast to construct subgenomic replicons, derived from a full-length cDNA clone of 17D strain of YFV, that express biologically functional reporter genes green fluorescent protein (GFP) and firefly luciferase (Luc).
MATERIALS AND METHODS

CELL CULTURE

BHK-21 cells were maintained at 37°C in minimum essential medium (MEM) supplemented with 1% antibiotics (penicillin 10,000 U/mL and streptomycin 10,000 μg/mL, GIBCO) and 10% fetal bovine serum (FBS).

YEAST AND E. COLI HOSTS

The yeast *Saccharomyces cerevisiae* strain RFY206 (MATa, his3A200, leu2-3, lys2A201, ura3-52, trplA:hisG) was cultured with YPD broth (1% Bacto-yeast, 2% Bacto-peptone and 2% dextrose), and selections were done using tryptophan (trp) dropout medium (Yeast Nitrogen Base without tryptophan). It was made competent by the lithium acetate procedure (Sambrook and Russell 2001). Electrocompetent *E. coli* strain DH10B (used for DNA propagation) were cultured using Luria-Bertani medium.

PLASMID CONSTRUCTS

Homologous recombination in yeast was used for all plasmid constructions. The parental low copy number plasmid pBSC-repYFV-17D containing the prototype YFV-17D strain replicon (repYFV-17D) was previously characterized (Gil et al. unpublished data). The replicon repYFV-17D contains deletion of almost all of the structural proteins except 63 N-terminal nucleotides (nts) of the capsid protein and 72 C-terminal nts of the E protein. To facilitate cloning of foreign genes, a unique cloning site NarI was incorporated between these sequences.

In this study, two similar YFV-17D replicons were constructed, both with the same overall genetic structure, but expressing the sequence that code for two different reporter genes: the green fluorescent protein (repYFV-17D-GFP) and the firefly luciferase (repYFV-17D-Luc). The generation of the repYFV-17D-GFP were based on homologous recombination between one PCR product and the linearized vector in yeast (Figure 1).

![Figure 1](https://example.com/figure1.png)

**Figure 1** - Schematic representation of construction of subgenomic replicons repYFV-17D-GFP and repYFV-17D-Luc. (A) Homologous recombination in yeast between a PCR product containing Luc or GFP gene and the vector pBSC-repYFV-17D NarI digested. The PCR product is represented with a sequence identification. (B) Genome organization of subgenomic replicons repYFV-17D-GFP and repYFV-17D-Luc. * Homologous sequences (25 nts) for yeast recombination.
The PCR fragment containing the GFP gene was amplified from the version of the plasmid Green Lantern (pGL) from Stratagene using one primer pair that contained homologous sequences for recombination with the NarI digested vector pBSC-repYFV-17D (Table I).

<table>
<thead>
<tr>
<th>Primera</th>
<th>Sequence</th>
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<tbody>
<tr>
<td>Rep-YFV-GFP-Fb</td>
<td>CAATATGGTACGACGAGGAGTTCGCATGAGCAAGGGCGAGGAACTGTTCAC</td>
</tr>
<tr>
<td>Rep-YFV-GFP-Rc</td>
<td>TCAATGCTCATGGACATTGTCATGTTCTTGTACAGCTCGTCCATGCCATGTG</td>
</tr>
<tr>
<td>Rep-YFV-Luc-F</td>
<td>CAATATGGTACGACGAGGAGTTCGCATGGAAGACGCCAAAACATAAAG</td>
</tr>
<tr>
<td>Rep-YFV-Luc-R</td>
<td>TCAATGCTCATGGACAGGAGTTCGCATGGAAGACGCCAAAACATAAAG</td>
</tr>
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The homologous sequences for recombination in yeast are underlined.

The homologous sequences are underlined. The primers are identified according to the sequences that have and / or amplify. b F-forward. c R-inverse. The primers Rep-YFV-GFP-F and Rep-YFV-GFP-R were used to construct the replicon repYFV-17D-GFP, while the primers Rep-YFV-Luc-F and Rep-YFV-Luc-R were used to construct repYFV-17D-Luc.

The PCR product and the linearized vector pBSC-repYFV-17D were used to transform competent RFY206. After transformation, yeast cells were plated out on solid trp dropout medium. Individual clones were screened by PCR to confirm the presence and correct orientation of the reporter gene. Positive clones were grown in liquid trp dropout medium, and plasmid DNA extraction was performed using QIAprep Miniprep Kit according to manufacturer’s instruction.

The second replicon (repYFV-17D-Luc) was also constructed using homologous recombination of one PCR product. The DNA template used in PCR reactions was the plasmid LNBr-WT that contains the Luc gene. The primers sequence is showed in Table I.

**E. coli ELECTROPORATION**

In order to amplify the DNA after homologous recombination, the purified yeast DNA was used to transform competent *E. coli* strain DH10B. Plasmid DNA were electroporated using 2 mm cuvettes on an ECM BTX electro cell manipulator 830™ (BTX, Holliston, MA) with the following settings: 2.75 kV, 99 µsec and 5 pulses with 1 sec interval. Individual colonies were grown at 37°C overnight in Luria-Bertani broth medium containing chloranfenicol as resistance marker. Plasmid DNA was then purified using the QIAGEN Plasmid Mid kit according manufacturer’s instructions.

**IN VITRO TRANSCRIPTION**

DNA templates for *in vitro* transcription were produced by PCR amplification of the region containing the T7 promoter and the entire replicon of the relevant plasmids using KlenTaq polymerase (Clontech). Full length PCR product were purified by phenol-chloroform, ethanol precipitated and then *in vitro* transcribed using MegaScript T7 kit (Ambion) supplemented with a 7mG(PPP)G cap analogue (Ambion). The yield and integrity of transcripts were analyzed by gel electrophoresis under denaturing conditions. Aliquots of transcription reactions were used for electroporation of BHK-21 cells without any additional purification.

**RNA TRANSFECTION**

Approximately 2x10⁶ BHK21 cells in 100uL Cytomix solution (Ansari et al. 2004) were
electroporated with 5µg of in vitro transcribed RNA in 2 mm cuvettes using an ECM BTX electro cell manipulator 830™ (BTX, Holliston, MA) with five pulses at the setting of 600 V, 99 µsec and 1 sec of interval. After being at room temperature for 5 min, the cells were ressuspended in fresh medium. For immunofluorescence assays, transfected cells were spotted onto 24-well plate coated with 10-mm glass coverslips.

**IMMUNOFLUORESCENCE ASSAYS**

Indirect immunofluorescence assay were used to detect viral protein expression in replicon-transfected cells. For this purpose, transfected cells on coverslips were rinsed with PBS and fixed with cold acetone at -20°C for 5 min. Then, cells were incubated at 37°C for 1 hour with a 1:100 dilution of a hyperimmune ascitic fluid raised against group B flaviviruses (Instituto Evandro Chagas), rinsed with PBS, followed by incubation with 1:100 dilution of a fluorescein isothiocyanate (FITC)-conjugated goat anti-mouse IgG antibody (Sigma-Aldrich) at 37°C for 1 hour. Following final washing, drying, and mounting, cells were visualized with a DMI 4000B fluorescence microscope (Leica).

**LIVE FLUORESCENCE ANALYSIS**

To observe GFP expression into live cells, repYFV-17D-GFP-transfected cells were rinsed with 1X PBS buffer and visualized by fluorescence microscopy using a SP2 AOBS inverted fluorescence microscope (Leica). Green lantern plasmid DNA transfected cells were used as positive control.

**LUCIFERASE ACTIVITY ASSAY**

To measure luciferase activity, naive and repYFV-17D-LUC-transfected BHK-21 cells were counted, washed with PBS and lysed by the addition of cell culture lysis buffer (Promega) according manufacturer’s instructions. Cell lysate were then centrifuged, and 20 uL of the resulting supernatant were mixed with 100 uL of luciferase substrate prior measurement of luciferase activity using a Mithras LB 940 Bioanalyzer (Berthold).

**RESULTS**

**CONSTRUCTION OF YFV REPLICONS**

The subgenomic replicons derived from a full-length cDNA clone of 17D strain of YFV expressing biologically functional reporter genes (GFP and LUC) were constructed using a yeast-based homologous recombination cloning method.

The replicon repYFV-17D constructed from the plasmid pBSC-repYFV-17D has an in-frame deletion of almost entire structural protein coding sequence, but retained the first 63 nts N-term of the capsid protein coding sequence and the 72 nts C-term of the E-protein. These important sequences for flavivirus replication includes the sequence of cyclization replication and signal sequence for NS1 protein during flavivirus protein synthesis, respectively (Khromykh and Westaway 1997, Lindenbach et al. 2007). To allow the expression of reporter genes from flavivirus replicons, we used a strategy that involved the in-frame fusion of the reporter genes GFP and LUC, following the 21 codons encoding the N-terminus of the capsid protein.

The use of the yeast-based homologous recombination cloning technique allowed the construction of repYFV-17D-GFP and repYFV-17D-Luc replicons in one step. Digestion of pBSC-repYFV-17D with NarI restriction enzyme and dephosphorylation of the ends yielded a linear plasmid unable to replicate in yeast, unless the plasmid was again circularized. Circularization was facilitated by recombination with overlapping DNA fragments, generating chimeric repYFV-17D sequences. After recombination in Saccharomyces cerevisiae four clones of each replicon construct were selected for confirmation of recombination by PCR (Figures 2A and 3A), using specific primers that amplified the inserted region, related to each...
The green fluorescent reporter protein expression was detectable by fluorescence analysis of live GFP in BHK-21 cells as early as 48 hours after transfection with repYFV-17D-GFP replicon RNA as with the green lantern plasmid DNA (Figure 2C). Also, the replication of the replicon was indirectly confirmed by the accumulation of yellow fever virus specific proteins, detected by immunofluorescence staining of transfected cells using specific antibodies against flavivirus proteins.
Luciferase activity was detected in cellular extracts prepared from BHK-21 cells electroporated with \textit{in vitro} transcribed repYFV-17D-Luc RNA for seven days post-electroporation. As shown in Figure 3B, the level of luciferase activity reached a maximum value at 48 hours post-electroporation. After this point time, the luciferase activity was also detected, but in a lower level until the minimum value had been reached at day seven. The autonomous replication of YFV replicon expressing the luciferase reporter gene (repYFV-17D-Luc) was also demonstrated by immunofluorescence analysis with specific antibodies to flavivirus proteins, 48 hours after transfection of BHK-21 cells (Figure 3C). The results showed that the expression of the reporter gene was dependent on the amplification of the YFV replicon by RNA replication and that its expression may be the transient.

The efficient expression of the reporter genes GFP and Luc reveals the correct insertion of the gene cassette into the replicon, the correct translation/processing of the protein and the efficient replication of the YFV-replicon.

**DISCUSSION**

In yeast, homologous recombination is an important strategy to repair double-stranded DNA breaks and create genetic diversity (Orr-Weaver et al. 1981, Kuzminov 2001). Overlapping DNA fragments undergo recombination readily when co-transformed into yeast cells (Oldenburg et al. 1997, Gibson et al. 2008, Panicali and Paoletti 1982, Post and Roizman 1981). So, the homologous recombination method can be effectively used to clone a desired gene into a plasmid and to generate chimeric genes with precise fusion junctions without the need of appropriate restriction sites.

A DNA fragment whose ends are identical to plasmid sequences can be directly inserted into a linearized vector by \textit{in vivo} homologous recombination in yeast, alleviating the need for ligation \textit{in vitro}. This method relies on sufficient homology at the ends of the DNA fragment and plasmid to engage the recombination machinery. At first, it were thought that 100 nts of homologous was required in order to recombinantion take place. Nowadays, it is known that 15 nts of homologous sequence is sufficient to mediate homologous integration (Manivasakam et al. 1995). The present study shows that homologous ends of 25 nts incorporated into primers get PCR products to efficient yeast-based homologous recombination. This cloning strategy has yet a limited application in the flavivirus genome manipulations (Puri et al. 2000, Polo et al. 1997, Pang et al. 2001) and their use are encouraged mainly by the stability of these genomes when manipulated in yeast instead of bacteria. Difficulties are encountered to construct flavivirus infectious clones because of the apparent genomic instability when using prokaryotic cells, as \textit{Escherichia coli} (Kapoor et al. 1995, Lai and Monath 2003, Polo et al. 1997, Sumiyoshi et al. 1992). For this reason, the use of the yeast \textit{Saccharomyces cerevisae} for cloning strategies has also proved to be efficient to by-pass the deleterious effects and instability in \textit{E. coli} (Polo et al. 1997).

Replicons of the flavivirus express all the non-structural proteins in order to mimic the expression during a viral infection and has proven to be a powerful tool to study the role of non-structural proteins and viral replication (Khromykh et al. 1998, Lindenbach and Rice 1997). Moreover, amplification of replicon RNA in the cytoplasm of cells makes them excellent vectors for the expression of heterologous genes at high levels, so transient and stable (Khromykh 2000). Featuring one or more structural proteins deleted, replicons are competent for autonomous replication, but unable to form infectious viral particles (Jones et al. 2005, Kofler et al. 2004, Widman et al. 2008).

Cloning of genes reporters have been widely used in the manipulation of viral genomes (Ng et al. 2007, Fayzulin et al. 2006, Scholle et al. 2004, Rossi...
et al. 2005, 2007) and is a valuable tool for evaluation of vector competence. The expression of these YFV-based replicons that encode reporter genes was found to be dependent on the amplification of input replicon RNA via RNA replication.

Most of the construction strategies of flavivirus reporter replicons previously proposed aim at the inclusion of the 3'-UTR reporter gene downstream of the IRES element of encephalomyocarditis virus (IRES-ECMV) to direct cap-independent translation of the reporter gene (Fayzulin et al. 2006, Khromykh and Westaway 1997, Shi et al. 2002, Scholle et al. 2004, Rossi et al. 2005). Insertions at the N-terminal of the NS1 of flavivirus replicons are followed by 2A protease sequence of the foot and mouth disease virus (2A-FMDV) to ensure proper processing of the heterologous protein (Shustov et al. 2007) or IRES-ECMV (Jones et al. 2005, Ng et al. 2007). All these strategies are based on standard cloning methods with the need for ligation in vitro, the use of restriction sites and a long time consuming. Such methods generally involve a number of steps, and the transfer of viral genetic material to different plasmids for growth in various bacteria, for the different stages of the overall process. In addition, the location and number of particular restriction sites is often a limitation to the generation of specific site-directed mutants.

In this work we have demonstrated that yellow fever virus replicon vectors constructed by homologous recombination in yeast can be used for expression of foreign genes in mammalian cells. The GFP and Luc expression by replicons repYFV-17D and repYFV-17D-GFP-Luc (Figures 2C and 3C), respectively, fall within the competence and replicative vector of these replicons. The efficiency of homologous recombination achieved was satisfactory (Figures 2A and 3A) and reinforces the applicability of this technique in studies of gene cloning.

YFV replicons lacking 2A protein sequence between the reporter gene and the NS1 signal sequence were previously found to be deficient in reporter gene expression (Jones et al. 2005). In contrast, we have constructed by homologous recombination in yeast functional replicons with insertion of a heterologous gene in this region. The expression of non-structural proteins in cells electroporated with RNA transcribed in vitro from both replicons, repYFV-17D-GFP (Figure 2B) and repYFV-17D-Luc (Figure 3B) revealed the replicative efficiency of these replicons. These results show that it is possible to insert heterologous genes between C-terminal end of the cyclization sequence of C protein and N-terminal end of the signal sequence of the envelope protein of the flavivirus replicons without interfering with viability.

As vaccine vectors or expression vectors for heterologous genes, replicons of the flavivirus present several advantages over conventional vectors, including: high levels of gene expression; capacity of autonomous replication; replication exclusively cytoplasmic, eliminating any possibility of chromosomal integration; inability to perform more than one cycle of infection, making them a safe tool for vaccine development, and small genome that allows easy manipulation (Anraku et al. 2002). Another advantage of the replicons are the different ways in which they can be used as immunizing: DNA, RNA or defective particles (Varnavski et al. 2000, Aberle et al. 2005, Ishikawa et al. 2008, Suzuki et al. 2009, Widman et al. 2008).

We present here an approach to the generation of replicons that reduces limitations of standard techniques for cloning and the number of steps needed to modify the viral genomes. The successful recovery of functional subgenomic replicons derived from RNA transcribed from an infectious clone YFV-17D has implications in developing new vaccine strategies and tools for studies of viral replication and pathogenesis. Moreover, the genomic stability conferred by recombination system in yeast makes this tool even more reliable and safe for the use of replicons for different purposes.
RESUMO

O replicon de RNA derivado do genoma de Flavivirus é uma ferramenta valiosa para o estudo de replicação viral independente da montagem e da maturação do virion, além de possuir um grande potencial para expressão de genes heterólogos. Neste estudo nós descrevemos a construção de replicons subgenômicos do vírus da febre amarela utilizando a técnica de recombinação homóloga. O plasmídeo contendo o replicon do vírus febre amarela cepa 17D (pBSC-repYFV-17D), caracterizado anteriormente, foi manipulado para a expressão heteróloga dos genes repórteres green fluorescent protein (repYFV-17D-GFP) e firefly luciferase (repYFV-17D-Luc). Ambos os replicons foram construídos por recombinação homóloga entre o vetor pBSC-repYFV-17D linearizado e o produto de PCR contendo 25 nucleotídeos terminais homólogos incorporados aos oligonucleotídeos iniciadores. A organização genômica dos construídos é semelhante ao repYFV-17D, com inserção de um gene repórter entre os restantes 63 nucleotídeos N-terminais da proteína do capsídeo e 72 nucleotídeos C-terminais da proteína E. Os replicons repYFV-17D-GFP e repYFV-17D-Luc mostraram uma eficiente replicação e expressão dos genes repórteres. A técnica de recombinação homóloga em levedura usada neste estudo demonstrou ser aplicável à manipulação do genoma do vírus da febre amarela para a construção de replicons subgenômicos.

Palavras-chave: técnica de clonagem, recombinação homóloga, replicon, gene repórter, vírus da febre amarela.

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