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Reverse transcription, primer pools preparation and multiplex PCR steps for CHIKV serotype genomic sequencing

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Laís Ceschini¹, Luisa Maria Inácio da Silva¹, Gabriel Luz Wallau¹

¹Entomology department, Instituto Aggeu Magalhães (IAM), FIOCRUZ



Luisa Maria Inácio da Silva

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We use this protocol and it's working

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Abstract

This step-by-step protocol describes the cDNA synthesis, primer pools preparation and multiplex PCR conditions with the main goal to sequence the complete genome of CHIKV serotype strains.

Materials

Reverse transcription: SuperScript™ IV First-Strand Synthesis System. (200 reactions) Cat:18091200 Invitrogen

Multiplex PCR: Q5® High-Fidelity 2X Master Mix. Cat: M0492L NEB, H2O Ultrapure, primers described in table 1.



Reverse transcription

- 1 Using a 2mL tube prepare the **Mix 1** described below for 96 samples:

A	B	C
Mix 1 Reverse transcriptio n	Vol. (1x)	96 samples (+2 = 98 to keep some extra due to pipetting issues)
Random Hexamers (50μM)	1μL	98μL
dNTPs mix (10mM each)	1μL	98μL
Total	2 μL	194μL

- 2 Using 0,2mL PCR tubes or 96 wells plates add 11-16μL of extracted RNA from RT-PCR positive samples. Add **2μL** of **Mix 1** to the tube/well and take it to the thermocycler with the following set up:

65°C ----- 5 minutes

- 3 Take the tubes/wells to ice for 1 minute. (you can prepare a water bath with ice cubes to have a uniform temperature distribution).

- 4 Using a 2mL tube prepare **Mix 2**:

A	B	C
Mix 2 Reverse Transcriptio n	Vol. (1x)	96 samples (+2 = 98 to keep some extra due to pipetting issues)

	A	B	C
	5x SSIV Buffer	4µL	392µL
	100mM DTT	1µL	98µL
	RNaseOUT or RNase Inhibitor	1µL	98µL
	SSIV Reverse Transcriptase	1µL	98µL
	Total	7µL	686µL

- 5 Add **7µL of Mix 2** to the tubes containing the **Mix 1** plus RNA and take it to the thermocycler following the set up below:

Step1:

42°C ----- 50 minutes

70°C ----- 10 minutes

4°C ----- Hold

- 6 Store the cDNA at -20°C.

Observation:.. As a suggestion, to improve the final results only samples RT-PCR positive showing a Ct value of < 30 should be used for cDNA conversion and genomic amplification.

Pools of primers

- 7 Select two 0,6mL tubes for each pool.
- 8 Using the original 100uM primer solution eluted individually, put them together following the table below containing each primer volume.
- 9 Pool 1 will have a final volume of 469µl and pool 2 of 460µl.

- 10 In order to prepare the solution to use in the Multiplex PCR, dilute each pool 1:10. That is, 10µl of pool 1 and 90µl of ultrapure water.

TABLE 1: Primers and pool order

	A	B	C	D	E
	Primer	Sequence	Concentrati on inside of the pool *	Volume of primer within the pool	Pool
	400_1_LEFT_1	TGACACAC GTAGCCTA CCAGTT	0,015uM	10ul	1
	400_1_RIGHT_1	CGCATCGG GCAAACGC AGTGGTA	0,015uM	10ul	1
	400_3_LEFT_3	GCAGACGT CGCGATAT ACCAAG	0,015uM	10ul	1
	400_3_RIGHT_3	CCAGCTCT TAAGTAGC ATGCCG	0,015uM	10ul	1
	400_5_LEFT_0	GATGTGCA AGACTACC GACACG	0,015uM	10ul	1
	400_5_RIGHT_0	GACTGGGT ATCAGGCC TCTTGT	0,015uM	10ul	1
	400_7_LEFT_4	CAAGAAGC CCAGGATG CTGAAA	0,015uM	10ul	1
	400_7_RIGHT_4	GCTATGCG TACACGTC TTCCT	0,015uM	10ul	1
	400_9_LEFT_4	GCAGAGAG GACAGAAC ACGAGT	0,015uM	10ul	1
	400_9_RIGHT_4	CTCTCTGT CTCATCAC GTCGGT	0,015uM	10ul	1
	400_11_LEFT_0	AGCAGTGC GGCTTCTT CAATAT	0,015uM	10ul	1



	A	B	C	D	E
	400_11_RIGH T_0	TGCCTAAC TGCGTAAA CTCCTTT	0,015uM	10ul	1
	400_13_LEF T_2	ATTAAGGA GTGGGAGG TGGAGC	0,015uM	10ul	1
	400_13_RIG HT_2	TCTAGAAT GGACGCTG CCTCAG	0,015uM	10ul	1
	400_15_LEF T_4	GGGGAAAG AATGGAAT GGCTGG	0,015uM	10ul	1
	400_15_RIG HT_4	CGTTCACT GGTTCTAT CTGCGT	0,015uM	10ul	1
	400_17_LEFT _0	AACTGAAC GCAGCCTT TGTAGG	0,015uM	10ul	1
	400_17_RIGH T_0	ACACCTGT GGAGAGGA GAGGTA	0,015uM	10ul	1
	400_19_LEF T_1	CATACAGAT GCGGACCC AAGTG	0,015uM	10ul	1
	400_19_RIG HT_1	GTTCAGGA GTCATGGC ATAACGG	0,015uM	10ul	1
	400_21_LEF T_2	GCGCGTAA GTCCAAGG GAATAC	0,015uM	10ul	1
	400_21_RIG HT_2	GTCTCCGC TGTTTCTT GTACGG	0,015uM	10ul	1
	400_23_LEF T_1	ACTTTCGG AGACTTCC TACCCG	0,015uM	10ul	1
	400_23_RIG HT_1	ACAGCCTC TCTTTAGTC TCTGGA	0,015uM	10ul	1
	400_25_LEF T_2	ACCAAATC ACCGATGA GTATGATGC	0,015uM	10ul	1



	A	B	C	D	E
	400_25_RIG HT_2	TCGTTATC CTGATAGG GCTGGC	0,015uM	10ul	1
	400_27_LEF T_4	AGGCCTAA GGTGCAGG TTATACA	0,015uM	10ul	1
	400_27_RIG HT_4	GCAGGTGA CAGCTGGA AATCTC	0,015uM	10ul	1
	400_29_LEF T_0	CGATGAAT TGATGGCA GCCAGA	0,015uM	10ul	1
	400_29_RIG HT_0	GCAAAGGT GGCCATGG ACATTA	0,015uM	10ul	1
	400_31_LEF T_1	TTCTACAAT AGGAGGTA CCAGCCT	0,015uM	10ul	1
	400_31_RIG HT_1	TTCATGCA CATTCTCT CTCTGCG	0,015uM	10ul	1
	400_33_LEF T_3	GATACCCG TGCACATG AAGTCC	0,015uM	10ul	1
	400_33_RIG HT_3	TTTTTCGTA GCAGCAGG GTGTG	0,015uM	10ul	1
	400_35_LEF T_0	CCACAAGA CCGTACCT AGCTCA	0,015uM	10ul	1
	400_35_RIG HT_0	TGGTGAAA TGGGTGCG TACATG	0,015uM	10ul	1
	400_37_LEF T_3	AATGTCAC AACAGTCC GGCAAT	0,015uM	10ul	1
	400_37_RIG HT_3	TTGGGTGG TCAGGATA CAGCAA	0,015uM	10ul	1
	400_39_LEF T_1	GGCCACCC GCATGAGA TAATTC	0,015uM	10ul	1



	A	B	C	D	E
	400_39_RIG HT_1	ATAGGACA ATCAGGGC TGCCAG	0,015uM	10ul	1
	400_41_LEF T_0	CTTGGAAC CAACGCTA TCGCTT	0,015uM	10ul	1
	400_41_RIG HT_0	AGCAGCCA CAGTGATAT TATTTCT	0,015uM	10ul	1
	400_43_LEF T_0	ACCAGGAC AATTTGGC GACATC	0,015uM	10ul	1
	400_43_RIG HT_0	ATACCTCA CACGACAT GTCCGT	0,015uM	10ul	1
	400_45_LEF T_3	CTACACAA GTACACTG TGCAGCC	0,015uM	10ul	1
	400_45_RIG HT_3	TGTTATTCA GGGGTTGT TCAGCC	0,015uM	10ul	1
	400_2_LEFT _0	CCAGCAAG GAGGATGA TGTCGGAC	0,015uM	10ul	2
	400_2_RIGH T_0	TGTGTCGA ACCCTACC CAGTAC	0,015uM	10ul	2
	400_4_LEFT _0	TGTTCTCA GTAGGGTC AACGCT	0,015uM	10ul	2
	400_4_RIGH T_0	GGATGCCG GTCATTTG ATCACA	0,015uM	10ul	2
	400_6_LEFT _1	TGAGAAGC TTTTGGGG GTCAGA	0,015uM	10ul	2
	400_6_RIGH T_1	ACATCTTC CTGTGCTG CCTGTA	0,015uM	10ul	2
	400_8_LEFT _0	ACTTTCCC CGCAGACC GTATTA	0,015uM	10ul	2



	A	B	C	D	E
	400_8_RIGH T_0	CAGCTTCT TCCTTCTT GCAGCA	0,015uM	10ul	2
	400_10_LEF T_0	ACCTGGTG ACTAGCGG AAAGAA	0,015uM	10ul	2
	400_10_RIG HT_0	GACGACAC AATGGCAG TCACAG	0,015uM	10ul	2
	400_12_LEF T_0	GAGGGTGG GTAAACA ACTGCA	0,015uM	10ul	2
	400_12_RIG HT_0	TTATCCCC GCTGTTTC GAGGAT	0,015uM	10ul	2
	400_14_LEF T_1	ACGCGGAT AACCACTG GGATAA	0,015uM	10ul	2
	400_14_RIG HT_1	TTATAGCC GCTAACCA GGAGCA	0,015uM	10ul	2
	400_16_LEF T_0	AGGTGACT CACTGAGA CTGCTC	0,015uM	10ul	2
	400_16_RIG HT_0	ATCGTTCT TCGCGATG TCCATG	0,015uM	10ul	2
	400_18_LEF T_2	GGACCAAA CTTCTCAA ATTACACG GA	0,015uM	10ul	2
	400_18_RIG HT_2	CCAAACTA CTGTCAGG GTGCAC	0,015uM	10ul	2
	400_20_LEF T_4	CAGAAATG CCCGGTGG ATGATG	0,015uM	10ul	2
	400_20_RIG HT_4	ATCGGCGC TTAGATCAA ACTGAC	0,015uM	10ul	2
	400_22_LEF T_0	GAGGGAGA AACCTGAC CGTGAT	0,015uM	10ul	2



	A	B	C	D	E
	400_22_RIG HT_0	AGTCATAA CTCGTCGT CCGTGT	0,015uM	10ul	2
	400_24_LEF T_4	CACGGCCA ATAGAAGC AGGTATC	0,015uM	10ul	2
	400_24_RIG HT_4	TTGACGGA TTGAATGT CGCTCG	0,015uM	10ul	2
	400_26_LEF T_0	ACCCACTT TGGACTCA GCAGTA	0,015uM	10ul	2
	400_26_RIG HT_0	AGGACGGC GTTCAATC TCCTAA	0,015uM	10ul	2
	400_28_LEF T_0	CCAGGATG ATTCACTT GCGCTT	0,015uM	10ul	2
	400_28_RIG HT_0	GGAGCTTT CTGGGATA CAACTGC	0,015uM	10ul	2
	400_30_LEF T_1	GATGGCAA CGAACAGG GCTAAT	0,015uM	10ul	2
	400_30_RIG HT_1	GGTCTGGG TCTGATGA CTTGGA	0,015uM	10ul	2
	400_32_LEF T_3	CCCCCAA AAGAAACC GGTTCA	0,015uM	10ul	2
	400_32_RIG HT_3	GAGTACTG TACTGCTC CGTGGT	0,015uM	10ul	2
	400_34_LEF T_0	CGTCACGA AAATCACC CCTGAG	0,015uM	10ul	2
	400_34_RIG HT_0	TCTGTCGC TTCGTTTC TGATGC	0,015uM	10ul	2
	400_36_LEF T_0	CCGTGCAC GATTACTG GAACAA	0,015uM	10ul	2

	A	B	C	D	E
	400_36_RIG HT_0	CACAATTG CACTTGTA CCGCAC	0,015uM	10ul	2
	400_38_LEF T_1	TCCTCTGG CAAATGTG ACATGC	0,015uM	10ul	2
	400_38_RIG HT_1	CACCCACC ATCGACAG GAGTAT	0,015uM	10ul	2
	400_40_LEF T_1	TATACCTGT GGAACGAG CAGCA	0,015uM	10ul	2
	400_40_RIG HT_1	TGTACCGC AGCATTTTC ACGTAC	0,015uM	10ul	2
	400_42_LEF T_4	TCAGCATA CAGGGCTC ATACCG	0,015uM	10ul	2
	400_42_RIG HT_4	GACGGTCT CTGCAGTA CCAGTT	0,015uM	10ul	2
	400_44_LEF T_0	ATCTCCAT CGACATAC CGGACG	0,015uM	10ul	2
	400_44_RIG HT_0	TGTGACGC CGGGTAAT TACTA	0,015uM	10ul	2
	400_46_LEF T_1	TCCCTAAA GAGACACA CCGCAT	0,015uM	10ul	2
	400_46_RIG HT_1	TCTTAGCT ATATATGGT GTGTCTCT TAG GG	0,015uM	10ul	2

*approximate concentration of each primer in the 25µl PCR reaction.

Note: The primers were designed using the <https://primalscheme.com> based on the KP164576, KP164571, KP164572, KP164568, KP164570 and KP164569 reference genomes (Machado, 2019).

Multiplex PCR



- 11 Prepare the **Mix 1** for a Multiplex PCR for each **Pool 1 and Pool 2** using a Falcon tube of 15mL (~96 amostras) or a 2mL tube.

	A	B	C	D
	Mix 1 Multiplex PCR	Vol. Pool 1 (1x)	Vol. Pool 2 (1x)	96 samples (+2) (pool1 ou pool2)
	Q5 Master Mix High fidelity 2X	12,5 µl	12,5 µl	1.225 µl
	Pool primers (Pool1 or Pool2) /Use concentrati on	1,7 µl	1,7 µl	166,6 µl
	Ultra Pure Water	8,3 µl	8,3 µl	813,3 µl
	Total	22,5µl	22,5µl	2205µl

- 12 Add **2,5µl of cDNA** (totalling 5µl) in 22,5µl of the pool1 and pool2 reaction and take it to the thermocycler following the conditions bellow:

Step1:

98°C ---- 30 seconds

Step2: (45 cycles)

98°C ---- 15 seconds

58°C ---- 30 seconds

72°C ---- 5 minutes

Step3:

72°C ---- 2 minutes

Hold 4°C