



AviSeq™
Genomic DX

CFTR

REF: AVG300016 16 TESTS
REF: AVG300096 96 TESTS



USER GUIDE

AVG300_IFU_Rev02_NOV23



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1. AIM OF USE

Aviseq CFTR is a kit for first and second level analysis of CFTR gene. Analysis is performed with a molecular protocol based on Next Generation Sequencing (NGS) technologies.

Aviseq CFTR must be used together with the following Avicenna reagents (not included in the kit) containing the specific adapters and indexes or barcodes, and it's compatible with Illumina, Ion Torrent and Nanopore sequencers:

Illumina: Aviseq Index Set 16 (Ref.AVG404) or Aviseq Index Set1 (Ref. AVG411) or Aviseq Index set 384 (Ref. AVG405)

Ion Torrent: Aviseq Barcode Set 1-16 (Ref. AVG502) or Aviseq Barcode Set 17-32 (Ref. AVG503)

The kit is validated for germline analysis (SNPs, indel) of DNA extracted from blood and tissues (fresh, frozen).

TARGET GENES
CFTR

Table 1: Genes targeted.

2. KIT CONTENTS

The kit size is 16 or 96 tests.

Note: the excess volume present in the kit is calculated to allow the subdivision respectively into a maximum of 2 or 4 analysis sessions.

Tube	Description	Volume 16 tests	Volume 96 tests
1	CFTR Pool 1	20 µl	110 µl
2	CFTR Pool 2	20 µl	110 µl
Taq	Hot Start Hi-Fi Taq	12 µl	64 µl
BA	Buffer A	320 µl	1840 µl
BB	Buffer B	320 µl	1840 µl
NTP	dNTPs Mix 10 mM	36 µl	184 µl

Table 2: Kit contents.

3. STORAGE AND PRODUCT STABILITY

All reagents provided with our kits are ready to use and should be stored at -20°C.

The kit, intact and properly stored, will maintain high quality performance capacity until the expiry date marked on each single reagent tube and on the external jar/container.

When the reagents are thawed, they must be kept on ice throughout the process.

4. KIT FEATURES

- SPECIFICITY: >99%
- SENSITIVITY: >99%

5. REQUIRED MATERIAL NOT INCLUDED

5.1 Generic Material

- Computer with constantly updated and guaranteed secure internet connection.
- Micropipettes calibrated and periodically verified 0,2-2 µl, 2-20 µl, 20-200 µl or 100-1000 µl and filter tips.
- Vortex.
- Disposable Gloves without powder.
- Thermal cycler calibrated and periodically verified.
- Tubes and Caps or 96-wells plate, as needed, DNase and RNase free.
- Nuclease-free water.
- 1,5 ml tube magnetic separator or 96-wells plate compatible magnetic separator.
- Fresh 70% ethanol.
- Illumina sequencers calibrated and periodically verified.
- For Illumina sequencers: Aviseq Index Set 16 (Ref.AVG404) or Aviseq Index Set1 (Ref. AVG411) or Aviseq Index set 384 (Ref. AVG405)
- For Ion Torrent sequencers: Aviseq Barcode Set 1-16 (Ref. AVG502) or Aviseq Barcode Set 17-32 (Ref. AVG503)

5.2 Specific Material

The material listed below has been used and validated by Avicenna:

- Qubit™ 2.0 Fluorometer (Invitrogen Cod. Q32866) or Qubit™ 3.0 Fluorometer (Invitrogen Cod. Q33216) or Qubit™ 4.0 Fluorometer (Invitrogen Cod. Q33226) calibrated and periodically verified.
- Qubit™ assay tubes (Invitrogen Cod. Q32856).
- Qubit™ dsDNA HS Assay Kit (Invitrogen, cod. Q32851).
- Magnetic beads for the purification of genomic libraries, AMPure XP (Beckman Coulter, cod. A63880) or MAGTIVIO (magSi-NGSPREP Plus, cod. MDKT00010075).

QUALITATIVE ANALYSIS OF DNA (Optional)

- Agilent 2100 Bioanalyzer system with DNA reagent kit calibrated and periodically verified.

6. IMPORTANT NOTES AND SAFETY INFORMATION

The user is required to apply the following provisions.

If the device or the results it generates, even in part, is transferred to a third party, the user must inform the end-user about the application of the specific provisions. The manufacturer is committed to constantly checking the possibilities of implementing the procedures, providing support to users.

- The kit is for professional use, it must be used by trained professionals in molecular biology.
- Do not use if package damaged.
- Biological samples and all reagents should be used in properly equipped rooms, clean and clear of potential contaminants. We suggest cleaning working areas frequently using a solution containing sodium hypochlorite 1-5%.
- Always use safety equipment such as laboratory coat, gloves and safety goggles during all steps described in the protocol.

- Check the risks and safety procedures associated with instruments, electricity, chemicals and other resources applied to the use of the device.
 - Prepare ways of detecting errors in the operation of the device, evaluating after each usage the quality of the results generated; in case of doubts or anomalies found, the supplier must be promptly contacted for support.
 - When the results produced are used in diagnostic or clinical processes, the user is required to consider the possible risks associated with diagnostic errors, to set up control mechanisms and to inform the medical personnel responsible for the diagnostic or clinical processes.
 - To avoid contamination of reagents we recommend using DNase/RNase free tubes, filter tips and to pay particular attention to keep all instruments clean and free of contaminants.
 - We suggest preparing a unidirectional workflow from the initial phase of DNA isolation following the PCR preparation phase, amplification and post-amplification phases in order to keep working areas separated for the different phases of the procedure using for each phase dedicated laboratory coats, micropipettes, tubes and filter tips.
 - Used reagents and biological samples must be wasted according to legal procedures.
- **Stopping point:** every time is present a stopping point you can proceed with the following step, or store the samples at 4°C for 24 hours or -20°C for a longer period.

7. OPERATIVE PROTOCOL

7.1 DNA SAMPLES PREPARATION

Use any commercial kit to obtain DNA from biological tissues. Determine the starting DNA concentration by fluorometric methods for accuracy. Dilute each DNA sample in PCR grade water to a final concentration of 5 ng/μl.

NB: The efficiency of the amplification process strongly depends on the quality of the DNA. For not fragmented DNA (i.e. fresh tissue, cytological samples...) 10 ng are in general sufficient for a good amplification process.

7.2 TARGET ENRICHMENT

Amplification of the target regions are performed with PCR in 2 reaction tubes, each containing one different CFTR pool. At the end of PCR, all the PCR reactions will be mixed into a single pool.

Refer to the table below for the quantities of reagents required to prepare each reaction of target enrichment:

Multiplex POOL 1 (Mix x 1 reaction)		Multiplex POOL 2 (Mix x 1 reaction)	
CFTR Pool 1	1μl	CFTR Pool 2	1μl
Buffer A	4μl	Buffer A	4μl
Buffer B	4μl	Buffer B	4μl
dNTPs Mix 10 mM	0.4μl	dNTPs Mix 10 mM	0.4μl
Hot Start Hi-Fi Taq	0.2μl	Hot Start Hi-Fi Taq	0.2μl
DNA	10ng	DNA	10ng
H2O	Up to 20μl	H2O	Up to 20μl
TOTAL	20μl	TOTAL	20μl

Table 3: Quantities of reagents for each target enrichment reaction.

The Reaction could be set up in 2 STEPS:

Step 1: Prepare AmpliMIX

In a clean and appropriated work area (e.g. laminar workflow) prepare the AmpliMIX, the amplification mix containing the PCR reagents common to all reaction tubes.

- Unfreeze Buffer A, Buffer B, dNTPs, CFTR Pool 1 and 2. Vortex and briefly centrifuge. Keep Hot Start Hi-Fi Taq in ice (do not vortex).
- Prepare AmpliMIX as described in the following table:

AmpliMIX	x 1
Buffer A	8 µl
Buffer B	8 µl
dNTPs Mix 10 mM	0.8 µl
Hot Start Hi-Fi Taq	0.4 µl
H2O	16.8 µl
TOTAL	34 µl

Table 4: AmpliMIX for 1 sample.

Step 2: Prepare Master Mixes

- Accurately mix the AmpliMIX (do not vortex), then divide the volume in 2 x 1,5 ml tubes and add a specific CFTR Pool to each tube, as to the table below:

MMixes	x 1	
	MMix 1	MMix 2
AmpliMIX	17 µl	17 µl
CFTR Pool 1	1 µl	-
CFTR Pool 2	-	1 µl
TOTAL	18 µl	18 µl

Table 5: Preparation of MasterMixes.

- For each sample, dispense 18 µl of each MasterMix per tube, change working area then add 2 µl of DNA (concentration 5 ng/µl) in each of the 2 PCR tubes. Briefly centrifuge the samples.

N.B. During all time keep the mixes and reagents on ice when not in use.

Place PCR tubes in the thermocycler and then start the program according to the following thermal profile:

Stage	Temperature (°C)	Time	Ramping	Cycles
Stage 1	98	2 min	4°C/s	1
Stage 2	98	10 sec	4°C/s	27
	60	2 min	1°C/s	
	72	20 sec	1°C/s	
Stage 3	72	5 min	4°C/s	1
Stage 4	10	HOLD	4°C/s	1

Table 6: PCR thermal profile.

At the end of PCR, mix the PCR products deriving from the same sample in a unique tube to obtain a library pool, and then proceed with the protocol.

➤ STOPPING POINT

INDEXING/BARCODING PROTOCOL

7.3 INDEXING/BARCODING PCR MIX PREPARATION

7.3.1 Defrost, quickly vortex and centrifuge the reagents supplied. Keep the Hot Start Hi-Fi Taq on ice (do not vortex).

N.B. For PCR use Buffer A, Buffer B, dNTPs and Hot Start Hi-Fi Taq contained in panel kit; during all time keep the mixes and reagents on ice when not in use.

Illumina: Aviseq Index Set 16 (Ref.AVG404) or Aviseq Index Set1 (Ref. AVG411) or Aviseq Index set 384 (Ref. AVG405)

Ion Torrent: Aviseq Barcode Set 1-16 (Ref. AVG502) or Aviseq Barcode Set 17-32 (Ref. AVG503)

7.3.2 For each sample, prepare a mix according to the following scheme:

ILLUMINA		ION TORRENT	
Master Mix	x1	Master Mix	x1
DNA (library)	5 µl	DNA (library)	5 µl
Buffer A	10 µl	Buffer A	10 µl
Buffer B	10 µl	Buffer B	10 µl
dNTPs Mix 10 mM	1 µl	dNTPs Mix 10 mM	1 µl
Hot Start Hi-Fi Taq	0.2 µl	Hot Start Hi-Fi Taq	0.2 µl
Indexes	2 µl	Barcode Oligo	2 µl
H ₂ O	21.8 µl	H ₂ O	21.8 µl

Table 7: Master Mix preparation volumes.

The Reaction could be set up in 2 STEPs.

Step 1: Prepare AmpliMIX:

AmpliMIX	x1
Buffer A	10 µl
Buffer B	10 µl
dNTPs Mix 10 mM	1 µl
Hot Start Hi-Fi Taq	0.2 µl
H ₂ O	21.8 µl
TOTAL	43 µl

Table 8: PCR mix preparation volumes.

Step 2: Prepare Master Mixes

For Ion Torrent: assign to each sample a specific barcode and add to each sample 2 µl of the selected barcode.

Prepare Master Mix (Table 9).

ION TORRENT	
Master Mix	x1
AmpliMIX	43 µl
Barcode	2 µl
TOTAL	45 µl

Table 9: Master Mix preparation.

7.3.3 In a post-PCR area, add 5 µl of the library (point 7.2) Please note that the total volume of the PCR should be 50 µl.

Proceed with point 7.3.4.

For Illumina: for each sample add 2 µl of indexes to AmpliMix, by picking from 96-well plate.

Note: for the conversion table refer to the IFU of each index set kit.

Prepare Master Mix (Table 10).

ILLUMINA	
Master Mix	x1
AmpliMIX	43 µl
Indexes	2 µl
TOTAL	45 µl

Table 10: Master Mix preparation.

7.3.3 In a post-PCR area, add 5 µl of the library (point 7.2) Please note that the total volume of the PCR should be 50 µl.

Proceed with point 7.3.4.

7.3.4 Briefly centrifuge samples.

7.3.5 Keep tubes on ice until placement into the thermocycler.

7.3.6 Place the reaction tubes/plate in the thermal cycler and run the following PCR protocol:

Stage	Temperature (°C)	Time	Ramping	Cycles
Stage 1	98	1 min	4°C/s	1
Stage 2	98	10 sec	4°C/s	15
	54	40 sec	1°C/s	
	72	3 min	1°C/s	
Stage 3	72	2 min	4°C/s	1
Stage 4	10	HOLD	4°C/s	1

Table 12: Thermal-cycler profile.

➤ **STOPPING POINT**

7.4 PCR PURIFICATION

In appropriate post-PCR area, proceed with the library purification:

N.B. Fresh ethanol 70 % is required for a proper wash low molecular weight molecule. Low ethanol concentrations can lead to loss of sample.

- a) For each sample, distribute 40 µl (0.8X) of magnetic beads (previously resuspended) in low binding 1,5 ml tubes.
- b) Add 50 µl of each amplified library pool; pipet up and down 5 times to thoroughly mix the beads suspension with the DNA.
- c) Incubate samples at room temperature for 5 minutes.
- d) Place tubes on magnetic support and incubate for 2 minutes or until the mixture appears clear; remove supernatant without touching the pellet.
- e) Add 200 µl of fresh ethanol 70% and rotate tubes twice to wash the beads; remove supernatant without touching the pellet.
- f) Repeat the washing step for a total of two times.
- g) Ensure that all ethanol is removed; put the tube on the thermal bath at 37°C for 3 minutes, then remove the tube from the thermal bath and incubate 1 or 2 minutes at room temperature, paying attention to not letting crack the beads pellet.
- h) Elute the DNA in 42 µl of RNase/DNase free distilled water; pipet up and down 5 times to thoroughly mix the beads suspension with the DNA.
- i) Incubate 5 minutes at 37°C.
- j) Place tubes on magnetic support and incubate for 2 minutes or until the mixture appears clear; carefully collect 40 µl of the supernatant containing the desired amplicons without touching the pellet.

➤ **STOPPING POINT**

7.5 QUALITATIVE AND QUANTITATIVE EVALUATION OF PCR PRODUCTS (OPTIONAL)

A quick estimation of DNA products size may be obtained running samples on agarose gel. However, we strongly recommend using Agilent 2100 Bioanalyzer (refer to manufacturers manual for protocol).

7.6 QUANTITATIVE EVALUATION OF PCR PRODUCTS

Quantify PCR products using a fluorometer, according to the manufacturer instructions, to prepare an equimolar pool of samples for sequencing.














For Illumina: dilute each library according to the Illumina sequencer procedure.

For Ion Torrent: dilute each library according to the Ion Torrent sequencer procedure.

8. TROUBLESHOOTING

PROBLEM	POSSIBLE CAUSE	SUGGESTION
Absence of bands - gel electrophoresis	Wrong PCR thermal profile	Verify the PCR thermal profile and calibration then repeat the PCR reaction
	Mistakes in master mix preparation	Verify PCR mix components and repeat the PCR reaction
	Degraded reagents	Verify expiry date and storing conditions of the products
	Presence of inhibitors	Verify concentration and quality of DNA extracted using a spectrophotometer. If necessary, repeat DNA extraction.
	Low amount of DNA	Verify concentration and quality of DNA extracted using a spectrophotometer. If necessary, repeat DNA extraction.
Absence of bands - gel electrophoresis	Wrong PCR thermocycling settings	Verify the PCR thermal profile and calibration then repeat the PCR reaction
Presence of low molecular weight fragments	Mistakes in master mix preparation	Verify PCR mix components and repeat the PCR reaction

9. SYMBOLS

	According to 98/79/CE Directive		Catalogue number
	In Vitro Diagnostic Medical Device		Batch code
	Expiration date		Temperature limitation
	Consult instruction for use (IFU)		Sufficient for n. tests
	Manufacturer		Do not use if package damaged
	Representative		