INTROL[®] CF Panel I Control v.02

INTENDED USE:

INTROL[®] CF Panel I Control v.02 is intended for *in vitro* diagnostic use as a quality control to monitor analytical performance of the extraction, amplification and detection steps of diagnostic assays used in the detection of the Cystic Fibrosis Transmembrane Conductance Regulator (CFTR) gene mutations and variants. This product is intended to be extracted and analyzed routinely with each CFTR assay run.

The INTROL[®] CF Panel I Control v.02 is designed to monitor the detection of 38 CFTR mutations associated with cystic fibrosis, including the 23 mutations recommended for testing by American College of Medical Genetics (ACMG) and American College of Obstetricians and Gynecologists (ACOG). The INTROLTM CF Panel I Control also monitors 3 polymorphisms (I506V, I507V, F508C) and the 5/7/9T variants.

PRODUCT DESCRIPTION:

INTROL[®] CF Panel I Control v.02 consists of synthetic CFTR DNA suspended in a matrix of carrier DNA of non-human species, preservatives, dye, and stabilizers. The synthetic DNA contains all 27 CFTR gene exons plus intronic borders, and contains specific mutations and polymorphisms which are divided among 3 bottles (bottles a, b, and c). The specific mutations present in each bottle are listed in Table 1; all other CFTR sequence is wild type. CFTR mutations that are not listed cannot be detected in the INTROL[®] CF Panel I Control v.02.

CFTR DNA is stabilized in the matrix and released when processed through common extraction methods as if it were a whole blood specimen. Following extraction, the released DNA can be used in common amplification based molecular assays techniques. Because INTROL[®] CF Panel I Control v.02 is designed to mimic the whole blood sample, the resulting copy number of the artificial CFTR segment, after extraction, will be similar to that found in a processed human whole blood sample (v/v).

INSTRUCTIONS FOR USE:

Extract and analyze INTROL® CF Panel I Control v.02 as you would a whole blood specimen:

- 1. Allow INTROL[®] CF Panel I Control v.02 to reach room temperature $(18^{\circ} 25^{\circ}C)$.
- 2. Thoroughly mix the controls prior to opening by inverting the bottle several times immediately before use, or by placing on an automated mixer.
- Extract INTROL[®] CF Panel I Control v.02 in the same manner as a whole blood clinical specimen. Use the same volume of INTROL[®] CF Panel I Control v.02 that would be used for a patient sample in your lab.

Note 1: Certain extraction methods may require additional processing of control material, such as dilution prior to analysis.

Note 2: The level of CFTR DNA present in the extracted control may not be detectable with certain quantitation methods and is not quantifiable by spectrophotometer measurements.

- 4. Analyze the extracted DNA as you would genomic DNA. If dilutions or other preparations of the extracted DNA are required as part of the testing procedure, handle the INTROL[®] CF Panel I Control v.02 DNA according to your standard laboratory protocol.
- 5. Tightly recap each bottle after use and store refrigerated $(2^{\circ} 8^{\circ}C)$.
- 6. Controls should be tested routinely as a matter of Good Laboratory Practice and according to guidelines or requirements of local, state, and/or federal regulations or accrediting organizations. The frequency of running the control material will depend on individual laboratory practice and may vary according to the analytical system being used.

STORAGE:

Upon receipt and after opening, the material should be stored at $2^\circ-8^\circ C. \ Do not freeze.$

STABILITY:

Unopened INTROL[®] CF Panel I Control v.02 material is stable through the expiration date printed on each bottle when stored refrigerated $(2^{\circ} - 8^{\circ}C)$. Opened material returned to the refrigerator $(2^{\circ} - 8^{\circ}C)$ shortly after use is stable for thirty (30) days from the date of opening. Contact MMQCI if control material was inadvertently frozen or exposed to high temperatures.

Table 1. Composition of INTROL[®] CF Panel I Control v.02 includes the following combinations of CFTR mutations and polymorphisms (plus wild type sequence covering 27 CFTR exons). Polymorphisms are in parentheses ().

Allele	Genotype		
Bot	tle a		
7T*	7T / 7T		
(I507V)*	I507V / WT		
(F508C)*	F508C / WT		
S549N/ S549R	Heterozygous		
S1251N	Heterozygous		
Bottle b			
E60X	Homozygous mutant		
G85E*	Homozygous mutant		
I148T	Homozygous mutant		
621+1G>T*	Homozygous mutant		
711+1G>T*	Homozygous mutant		
1078delT	Homozygous mutant		
R334W*	Homozygous mutant		
R347P*	Homozygous mutant		
9T*	9T / 9T		
A455E*	Homozygous mutant		
del F508*	Homozygous mutant		
V520F	Homozygous mutant		
1717-1G>A*	Homozygous mutant		
G542X*	Homozygous mutant		
G551D*	Homozygous mutant		
2184delA*	Homozygous mutant		
2789+5G>A*	Homozygous mutant		
3120+1G>A*	Homozygous mutant		
3199del6	Homozygous mutant		
D1152H	Homozygous mutant		
R1162X*	Homozygous mutant		
3659delC*	Homozygous mutant		
3849+10kbC>T*	Homozygous mutant		
3876delA	Homozygous mutant		
3905insT	Homozygous mutant		
W1282X*	Homozygous mutant		
N1303K*	Homozygous mutant		
Bot	tle c		
394delTT	Heterozygous		
R117H*	Heterozygous		
R347H	Heterozygous		
5T* / 7T*	Heterozygous		
(I506V)*	I506V / WT		
del I507*	Heterozygous		
R553X*	Heterozygous		
2183AA>G	Heterozygous		
*ACMG / ACOG Panel			

PRECAUTIONS AND WARNINGS:

- This product contains 23% ethanol (v/v) and could be flammable. Keep away from open flames.
- This product does not contain any biological material of human origin.
- The laboratory should follow Good Laboratory Practice (GLP) and establish its own performance characteristics for INTROL[™] CF Panel I in demonstrating adequate system performance.
- MMQCI CF products are not intended to be frozen and are shipped with a DO NOT FREEZE label.

LIMITATIONS:

- Interferences and cross-reactions may occur with some detection methods and confound interpretation of the test. Please refer to the kit manufacturers package insert to review possible cross-reactions and near neighbor interferences identified in the method.
- Recoveries may vary depending on extraction method, instrumentation, cycle time / temperature, reagents, method variation, and systematic or random errors.



INTROL® CF Panel I Control v.02

PERFORMANCE CHARACTERISTICS:

All INTROL[®] CF Panel I Control v.02 products are tested by an FDA-cleared CFTR mutation detection method before being released for market distribution. Any mutations not tested by the FDA-cleared method are sequenced bidirectionally before product is released. All mutations must be detected.

Clinical Evaluation - External Sites:

The clinical study performed at the external sites evaluated reproducibility of INTROLTM CF Panel I Control material with respect to within run, between run, between sites, between lots, and between methods.

Evaluation using different extraction methods:

# extraction methods	21
# laboratories	134
# successful laboratory extractions	129
percent successful	96% *

*Five laboratories didn't continue after DNA extraction because DNA quantitation method they used indicated that no DNA was extracted/isolated. Considering that the level of synthetic CFTR DNA present in the extracted control may not be detectable with certain quantitation methods, there is a possibility that extractions in these 5 laboratories may have been successful; however, this could not be assessed because the assays were not performed.

INTROL[®] CF Panel I Control v.02 material has been tested using CFTR assays at 10 external sites, 8 of which were clinical laboratories representing intended user. Samples from 11 different manufacturing lots were tested at minimally 3 external sites in at least 3 separate runs. Results are summarized in Table 2.

Table 2. External site evaluations.

Method	Site	# of Lots	# of Runs	Total Calls	% Correct Calls
Tag-It TM	1	10	9	223	100%
	2	3	9	138	100%
	3	1	1	6	100%
	4	1	1	7	100%
	5	1	1	4	100%
eSensor ®	6	5	1	30	100%
Other Amplification methods	7	5	1	38	100%
	8	3	1	31	100%
	9	1	2	7	100%
	10	5	40	649	100%
6 methods		11 Lots1	66 Runs	1133 Calls	100% Correct

1. Each bottle is processed independently and has its own lot number.

 $\rm INTROL^{\circledast}$ CF Panel I Control v.02 is protected by patents. It cannot be cloned, sold, or transferred to other laboratories without the explicit written consent of MMQCI.

Expected Results:

Expected results with the INTROL® CF Panel I Control v.02 using an FDAcleared CFTR assay are presented in Table 3.

Table 3. Results with the INTROL $^{\otimes}$ CF Panel I Control v.02 using an FDA cleared CFTR assay.

Method	Correctly Identified	No Call or	Not Tested
	Alleles	Other	by Assay
Tag-It [™]	All wt alleles	S549N: MUT ¹	S1251N,
	7T, I507V, F508C,		E60X,
	S549R, G85E, I148T,		2143delT
	621+1G>T, 711+1G>T,		3199del6,
	1078delT, R334W,		D1152H,
	R347P, 9T, A455E,		
	delF508, V520F,		
	1717+1G>A, G542X,		
	G551D, R560T,		
	1898+1G>A, 2184delA,		
	2789+5G>A,		
	3120+1G>A, R1162X,		
	3659delC,		
	3849+10kbC>T,		
	3876delA, 3905insT,		
	W1282X, N1303K,		
	394delTT, R117H,		
	R347H, 5T/7T, I506,		
	delI507, R553X,		
	2183AA>G		

Detected as homozygous mutant.

ORDERING INFORMATION:

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INTROL® Cystic Fibrosis Panel I Control v.02

Part Number:	G106ac
Kit Contains:	3 bottles x 2mL
	1 each G106a, G106b, and G106c
Part Number:	G106ac-1
Kit Contains:	3 bottles x 1mL
	1 each G106a-1, G106b-1, and G106c-1