

Instructions For Use

8247 Respiratory Control Panel (22 Targets)

Intended Use

The Respiratory Control Panel (22 Targets) is intended for use as non-viable, external, positive and negative control materials to evaluate the performance of nucleic acid amplification testing (NAAT) procedures that detect the analytes in Table 1. This product has no qualitative or quantitative assigned value. This control material is nonautomated and not intended to be used for screening, monitoring, or diagnosis. This control is not intended for any specific patient population or specimen.

Summary And Principles

The Respiratory Control Panel (22 Targets) can be used to monitor the extraction, amplification and detection process of molecular testing assays that include the analytes in Table 1. Routine use of quality controls monitor test variation, lot-to-lot test kit performance, operator performance, and aid in identifying random or systemic error.

Composition

The Respiratory Control Panel (22 Targets) consists of 6 individually packaged lyophilized positive control pellets and 6 individually packaged lyophilized negative control pellets. The kit also includes 12 vials of Molecular Hydration Fluid, each containing 300 µL of nuclease-free water. The analytes in Table 1 have been inactivated using irradiation, chemical, and/or thermal treatments.

The Respiratory Control Panel (22 Targets) is lyophilized in a PCR compatible matrix. The organisms are prepared in a buffered solution with materials of plant and animal origin, preservatives, and stabilizers. The solution is lyophilized into a ready-to-use pellet.

Table 1: Contents of the Respiratory Control Panel (22 Targets)


Analytes*	
Positive Control	
Viral Analytes	
Adenovirus Type 6	Influenza A subtype H1-2009
Coronavirus 229E	Influenza A subtype H3
Coronavirus HKU1 surrogate	Influenza B
Coronavirus NL63 surrogate	Parainfluenza Virus 1
Coronavirus OC43 surrogate	Parainfluenza Virus 2
Human Metapneumovirus surrogate	Parainfluenza Virus 3
Rhinovirus 1B	Parainfluenza Virus 4a surrogate
Influenza A subtype H1N1	Respiratory Syncytial Virus A
Influenza A subtype H3N2	SARS-CoV-2/USA/WA1/2020
Bacterial Analytes	
<i>Bordetella parapertussis</i>	
<i>Bordetella pertussis</i>	
<i>Chlamydomphila pneumoniae</i>	
<i>Mycoplasma pneumoniae</i>	
Negative Control	
Blank Pellets	

*All analytes are added at a target concentration of 10^3 - 10^5 copies per pellet. These are input concentrations and are not representative of recoverable concentrations or expected values.

Warnings And Precautions

- For In Vitro Diagnostic use only.
- For professional use only. To be used by personnel trained in the use of the assay.
- The inactivated lyophilized pellets are single-use only. Once hydrated, do not freeze for reuse.
- Do not open foil pouch until ready to use.
- Although this product has been inactivated, there is no known test or inactivation method that can assure that it will not transmit infection. This product must be treated as a potential biohazard. Wear appropriate personal protective equipment. Do not pipette by mouth. Do not smoke, eat, or drink in areas where specimens are handled. Disinfect any spills, and dispose of all materials in accordance with national and local regulations.
- Refer to the Safety Data Sheet (SDS) for more detailed information. The SDS can be located on the Microbiologics website at www.microbiologics.com or by contacting Customer Service at info@microbiologics.com.
- This product does not contain any hazardous substances listed in 1272/2008/EC.
- Report any serious incident that has occurred in relation to the device to Microbiologics and the local regulatory officials in which the user and/or the patient is established.

Storage And Expiration

 Store the Respiratory Control Panel (22 Targets) at 2°C-25°C in the original packaging up to the indicated expiration date. After opening the foil pouch, rehydrate and use immediately. In-use stability of the rehydrated pellet at room temperature (25°C) is 6 hours.

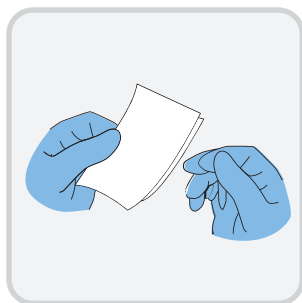
The Respiratory Control Panel (22 Targets) should not be used if:

- Stored improperly
- There is evidence of excessive exposure to heat or moisture
- The expiration date has passed
- Packaging is damaged

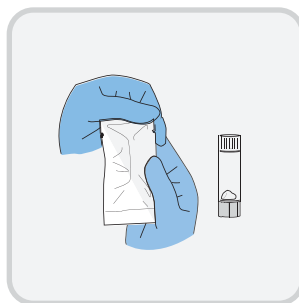
Materials Required But Not Provided

- Nucleic acid extraction kit, for assays that are not sample to result
- Instrumentation for detection
- Pipettors capable of delivering 0.5-1000 µl volumes
- Nuclease-free aerosol barrier pipette tips
- Vortex
- Microcentrifuge (optional)

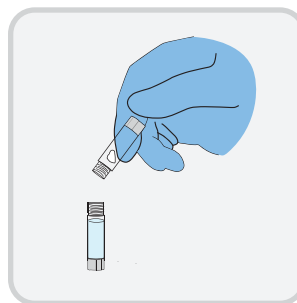
Instructions For Use



1. Read package insert, instructions for use or lab protocol for the applicable assay. Some instruments and assays are equipped with special QC settings. In these instances, it may be necessary to use the special setting when using QC sets and panels.



2. Tear open pouch at notch. Remove vial from pouch and ensure the pellet is at the bottom of the vial before opening.



3. Hydrate the lyophilized pellet into the provided vial of molecular hydration fluid.



4. Recap the vial and vortex for 10 seconds at full speed to mix.

5. If a centrifuge is not available, tap the capped vial on a rigid surface to collect material at the bottom of the vial.
a. Alternatively, centrifuge briefly to collect any droplets clinging to the cap or upper walls of the vial.

6. Use the appropriate volume for the assay being performed and follow laboratory protocols or manufacturer instructions for processing a sample.


























Limitations

- There are no known extrinsic factors or interfering substances specific to this product. Please consult assay instructions for any extrinsic factors or interfering substances.
- This product is unassayed control material. It may not be suitable for use with all kits and procedures as not all instruments and assays are compatible with multi-target controls. Customer is responsible for verifying the performance of this product with their chosen instrumentation and assay(s). As a third-party control manufacturer, Microbiologics' provides quality controls that deliver an independent, unbiased assessment of performance with any instrument or method. While not intended to replace control materials provided by the assay/instrument supplier, third-party control materials should be considered.
- Target concentrations of each analyte are specific to Microbiologics' assay method and procedures. These analytes are non-viable and may be used with any PCR-based test or assay. Microbiologics guarantees each nucleic acid is present and can be amplified but does not guarantee specific analyte concentrations. Each laboratory should establish its own range of acceptable values on their assay system per their internal quality assurance procedure/program. Nucleic acid reactivity, which may vary over time, is dependent on a laboratory's instrumentation, assay method, procedures, calibration, or technician. Microbiologics' molecular controls are not calibrators and should not be used for assay calibration or as an absolute reference material.

Microbiological State

This product was prepared using suitable inactivation methods. While the product has been tested for innocuity, universal laboratory precautions are recommended, and material should be treated as though it was a viable specimen.

Key Of Symbols

	Authorized Representative in the European Community / European Union		In vitro diagnostic medical device
	Batch code (Lot)		Manufacturer
	Biological risks		Negative control
	Catalog number		Positive control
	Caution		Quantity
	CE mark		Swiss Authorized Representative
	Consult instructions for use or consult electronic instructions for use		Telephone number
	Contains sufficient for <n> tests		Temperature limit
	Device for near-patient testing		UK Conformity Assessed mark
	Do not re-use		UK Responsible Person
	Do not use if package is damaged and consult instructions for use		Use-by-date
	Health hazard		Water; Fluid
	EU Authorized Representative		

Please refer to product labels for applicable symbols.

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Website

Visit our website, www.microbiologics.com, for current technical information and product availability.

Bibliography

- Zumla A, Al-Tawfiq JA, Enne VI, et al. Rapid point of care diagnostic tests for viral and bacterial respiratory tract infections - needs, advances, and future prospects. *The Lancet Infectious Diseases*. 2014; 14(11):1123-1135. Published 2014 Sep 1. doi:10.1016/S1473-3099(14)70827-8
- Boers SA, Peters CJA, Wessels E, Melchers WJG, Claas ECJ. Syndromic panels or “panel syndrome”? A perspective through the lens of pre-analytical and post-analytical phases of microbiological diagnosis. *Clinical Microbiology and Infection*. 2019;25(9):1077-1083. doi:10.1016/j.cmi.2019.03.011.
- Barenfanger J, Drake C, Leon N, Mueller T, Trout T. 2000. Clinical and Financial Benefits of Rapid Detection of Respiratory Viruses: an Outcomes Study. *J Clin Microbiol* 38. https://doi.org/10.1128/jcm.38.8.2824-2828.2000.

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Revision History

Publication History		
Revision	Date	Description of Change
A	2026-01	Initial release