

Comprehensive and rapid detection of respiratory pathogens and associated resistance genes using the MiniSeq[™] System

A 24-hour, sample-to-result sequencing workflow using Illumina RNA Prep with Enrichment and the Respiratory Pathogen ID/AMR Panel.

Analysis powered by



Introduction

Respiratory tract infections are a public health concern, with viral, bacterial, and fungal pathogens known to cause respiratory illnesses. Upper respiratory tract infections are typically less severe and include the common cold (viral rhinitis), sinusitis, pharyngitis, and others. Lower respiratory tract infections comprise bronchitis, bronchiolitis, and pneumonia. These illnesses, particularly pneumonia, can be severe and require hospitalization, or even be fatal.^{1,2}

Accurate identification of respiratory pathogens can be challenging, particularly in the case of mixed or co-infections. Conventional methods for detection include *in vitro* culture, antibody- or antigenbased assays, and PCR.³ PCR has become a preferred method for pathogen identification, due to its speed, high sensitivity, and specificity.⁴ However, due to the limited number of targets per assay, PCR can require multiple, sequential assays to test for even common pathogens. Compounding the COVID-19 pandemic with a flu season increases the potential for co-infections and complications with SARS-CoV-2 and other respiratory pathogens,⁵ highlighting the need for rapid, accurate, and broad-spectrum pathogen detection.

Next-generation sequencing (NGS) provides an effective way to analyze samples and detect known and emerging respiratory pathogens from a variety of sample types, including those with multiple infectious agents, in a single assay. To that end, Illumina offers the

Respiratory Pathogen ID/AMR Panel, which targets ~1500 respiratory pathogens, including viruses, bacteria, and fungi, and associated antimicrobial resistance (AMR) markers. Combining the Illumina Respiratory Pathogen ID/AMR Panel with Illumina RNA Prep with Enrichment and sequencing with the MiniSeq Rapid Reagent Kit offers several advantages, including:

- Sensitive detection of both DNA- and RNA-based respiratory pathogens in a single assay in under 24 hours
- Comprehensive genome coverage of critical viral pathogens, including SARS-CoV-2 and Influenza A virus
- Concurrent profiling of AMR marker expression for pathogen characterization in the same assay

This application note highlights a streamlined workflow for detecting and analyzing DNA- and RNA-based respiratory pathogens using Illumina RNA Prep with Enrichment combined with the Illumina Respiratory Pathogens ID/AMR Panel, rapid sequencing on the MiniSeq System, and simplified data analysis with the IDbyDNA Explify® Platform (Figure 1).

Methods

Sample preparation

Residual clinical samples were collected from the upper and lower respiratory tract positive for viral, bacterial, and fungal pathogens, as determined by conventional assays (Table 1). DNA and RNA were extracted from each sample and used as input to subsequent library preparation and enrichment. To mimic total nucleic acid extracted samples for spike-in studies, 10 ng of human DNA and 10 ng of RNA were pooled together. The total nucleic acid was carried through cDNA conversion and subsequent library preparation and was used to evaluate genomic coverage (Figure 3) and detection of AMR (Figure 5).

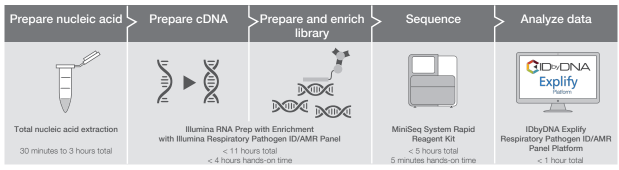


Figure 1: Enrichment workflow for respiratory pathogen detection—The streamlined NGS workflow for respiratory pathogen detection integrates sample preparation, library preparation, target enrichment, sequencing, and data analysis.

Table 1: Samples procured for analysis

		•	
Sample type	Number	Expected pathogens	Count
Dranahaaluaalar layaaa	6	Bacteria	7ª
Bronchoalveolar lavage		Fungi	1
Nasopharyngeal swabs	13	Viruses	14 ^b
Sputum	5	Bacteria	5
Tracheal aspirate	2	Bacteria	2
Total	26		29
a. One sample was positive forb. One sample was positive for			

Library preparation

Sequencing-ready libraries were prepared with Illumina RNA Prep with Enrichment (Illumina, Catalog no. 20040536) and IDT for Illumina DNA/RNA UD Indexes (Illumina, Catalog no. 20027213). Illumina RNA Prep with Enrichment uses On-Bead Tagmentation followed by a single hybridization step to generate enriched DNA and RNA libraries. The combined cDNA and DNA samples can be carried straight through into tagmentation with no quantification required.

After amplification, libraries were enriched as 1-plex or 3-plex reactions using the Illumina Respiratory Pathogen ID/AMR Panel. This panel provides targeted detection of 282 pathogens selected to cover > 95% of common and rare causes of respiratory infections using cutting-edge implementations of a combination of alignment-based, alignment-free, assembly-based, and machine learning-enhanced approaches. In addition, 1218 AMR markers are targeted to characterize resistance of 12 common bacterial pathogens to 16 widely used antibiotics (Table 2).

Sequencing

Libraries were denatured and diluted to a final loading concentration of 2 pM, according to the MiniSeq System Denature and Dilute Libraries Guide (Document no. 1000000002697 v07) and sequenced on the MiniSeq System at 1 \times 101 bp read length using either the MiniSeq High Output Reagent Kit (Illumina, Catalog no. FC-420-1002) or the MiniSeq Rapid Reagent Kit (Illumina, Catalog no. 20044338). Reads were trimmed to 1 \times 75 bp, unless otherwise noted. The read recommendation for this workflow is 1M reads per sample, but these numbers can vary, and this is only a recommended starting point.

Data analysis

FASTQ sequencing data files were input to the IDbyDNA Respiratory Pathogen ID/AMR Panel Platform, which can be accessed in BaseSpace™ Sequence Hub, for analysis. The IDbyDNA Explify Platform offers a paired and custom tailored data analysis solution for the Illumina Respiratory Pathogen ID/AMR Panel. It provides information on sample composition, including host and microbial abundance, and proportion of targeted vs. untargeted sequences. Users can select up to 35 individual samples or project folders containing any number of samples for analysis. Reports include a detailed text-based (JSON format) and PDF report.

Table 2: Targets on the Respiratory Pathogen ID/AMR Panel

Viruses: 42 targets				
Bacteria: 187 targets				
Fungi: 53 targets				
AMR: 1218 markers				
Curated antibiotics				
Amoxicillin	Gentamicin			
Amoxicillin-Clavulanate	Levofloxacin			
Cefazolin	Meropenem			
Cefepime	Oxacillin			
Ceftriaxone	Sulfamethoxazole			
Clindamycin	Tetracycline			
Colistin	Trimethoprim			
Erythromycin	Vanomycin			
Curated bact	terial pathogens			
Acinetobacter baumannii	Mycobacterium abscessus			
Enterobacter cloacae	Mycobacterium tuberculosis			
Enterococcus faecium	Pseudomonas aeruginosa			
Enterococcus faecalis	Staphylococcus aureus			
Escherichia coli	Stenotrophomonas maltophilia			
Klebsiella pneumoniae	Streptococcus pneumoniae			

Results

Improved respiratory pathogen detection with enrichment

Detection of various respiratory pathogens in clinical sample libraries was significantly improved with enrichment using the Respiratory Pathogen ID/AMR Panel, as compared to aliquots of the same libraries pre-enrichment (Table 3). Indeed, normalized pathogen reads were increased by a median of 330-fold in enriched samples (Figure 2). Enrichment was most pronounced in samples with lower pathogen load, ie, fewer identified pathogens in the pre-enriched libraries.

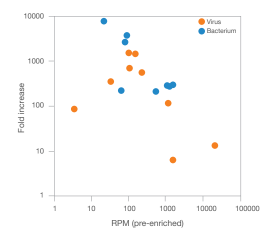


Figure 2: Enrichment increases normalized pathogen reads—Normalized pathogen reads were increased by a median of 330-fold in enriched viral (orange) and bacterial (blue) samples.

Table 3: Improved detection of pathogens with enrichment using the Respiratory Pathogen ID/AMR Panel

Pathogen		Detected without enrichment	Detected only with enrichment
Viruses	Human adenovirus Ba	1	1
	Human adenovirus C	2	2
	Influenza A virus (H1N1)	0	1
	Human metapneumovirus	1	2
	Influenza B virus	0	1
	Human parainfluenza virus 1	1	2
	Human parainfluenza virus 3	2	2
	Respiratory syncytial virus B	1	1
	SARS-CoV-2	1	2
Bacteria	Enterobacter cloacae complex	0	1
	Escherichia coli	1	2
	Haemophilus influenzae	2	2
	Klebsiella pneumoniae	1	2
	Legionella pneumophila	0	1
	Proteus mirabilis	0	1
	Pseudomonas aeruginosa	1	1
	Serratia marcescens	0	1
	Staphylococcus aureus	2	2
	Streptococcus pneumoniae	1	1
Fungi	Pneumocystis jirovecii	0	1
Total		17/29	29/29
a. Reported	d adenovirus B/E by conventional test.		

Comprehensive genomic coverage of critical viral pathogens

Full genome coverage of SARS-CoV-2 and Influenza A virus was achieved at $\geq 100,\!000$ viral copies (Figure 3). These results are significant, as they demonstrate that the Respiratory Pathogen ID/AMR Panel provides sufficient coverage of critical viral pathogens for analysis of viral evolution or viral surveillance.

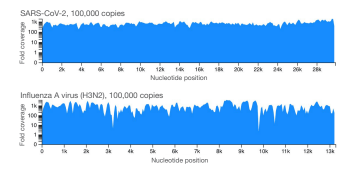


Figure 3: Comprehensive genomic coverage of critical viral pathogens — The Respiratory Pathogen ID/AMR Panel enables comprehensive coverage of viral genomes, including SARS-CoV-2 and Influenza A virus (H3N2), at 100,000 viral copies. Sequencing was run on the MiniSeq System at 1 \times 75 bp.

Pathogen detection using the MiniSeq Rapid Kit

To assess performance with the MiniSeq Rapid Kit, one pool of enriched libraries (n=19, including 17 libraries from clinical samples) were sequenced separately with the MiniSeq High Output Reagent Kit (150-cycles) and the MiniSeq Rapid Reagent Kit (100 cycles), both run with 1 ×101 cycles. Sequencing metrics were equivalent with the MiniSeq High Output Kit and MiniSeq Rapid Kit (Table 4).

Table 4: Comparison of sequencing metrics by reagent kit

Metric	MiniSeq High Output Kit (1 × 101 cycles)	MiniSeq Rapid Kit (1 × 101 cycles)
Total run yield	4.3 Gb	3.2 Gb
% bases ≥ Q30	94%	94%
Cluster density	241 K/mm²	210 K/mm ²
% clusters passing filter	87%	87%
PhiX error rate	0.50%	0.30%
% tile pass	100%	100%

In this experiment, clinical samples were positive for four viruses, eight bacteria, and one fungus. All respiratory pathogens detected using the MiniSeq High Output Kit were also detected using the MiniSeq Rapid Kit (Figure 4). These results demonstrate the highly consistent data and equivalent performance of the two reagent kits.

Detection of AMR markers

A titration experiment was conducted in which diminishing amounts of *Klebsiella oxytoca* was spiked into human background RNA and taken though library preparation and enrichment with the Respiratory Pathogen ID/AMR Panel. Analysis of sequencing data showed that all AMR genes expressed by *Klebsiella oxytoca* were detected as expected for this pathogen (Table 5).

Effect of enrichment plexity on pathogen detection

To evaluate the effect of plexity, the number of pre-enriched libraries that are pooled together in a single enrichment reaction, on pathogen detection, libraries from 21 clinical samples were prepared and enriched with the Respiratory Pathogen ID/AMR Panel either at 1-plex or 3-plex. Clinical samples were positive for 11 viruses, 13 bacteria, and one fungus. While libraries enriched in the 3-plex format resulted in more variable read counts than 1-plex enrichment (data not shown), pathogen detection was highly consistent, with all respiratory pathogens detected in all libraries at both plexities (Figure 5).

Summary

The identification and characterization of respiratory pathogens is central to improving public health. NGS is a powerful method for simultaneous, broad-range detection of multiple infectious agents. Combining Illumina RNA Prep with Enrichment and the Respiratory Pathogen ID/AMR Panel with MiniSeq Rapid sequencing and the IDbyDNA Explify Platform enables detection of hundreds of DNA and RNA respiratory pathogens in a single assay. By including AMR markers, the panel analyzes the expression of resistance genes to widely used antibiotics by common bacterial pathogens. Together, this sample-to-results workflow provides rapid, accurate, and broad-spectrum respiratory pathogen detection and characterization.

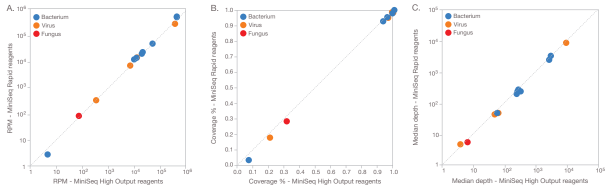


Figure 4: Consistent detection of respiratory pathogens using MiniSeq High Output and Rapid reagents—All respiratory pathogens detected using MiniSeq High Output reagents were also detected using MiniSeq Rapid reagents. Performance metrics, including (A) reads per million (RPM), (B) coverage %, and (C) median depth were highly consistent between reagent kits.

Table 5: Detection of Klebsiella oxytoca and expressed AMR genes with the Respiratory Pathogen ID/AMR Panel

			5	3			
Genome copy number	K. oxytoca	AAC(6')-lb-cra	aadA	ANT(3")-IIa	dfrA1	SHV-5	sul1
100,000	3/3	3/3	3/3	3/3	3/3	3/3	3/3
10,000	3/3	3/3	3/3	3/3	3/3	3/3	3/3
1000	3/3	2/3	1/3	3/3	2/3	3/3	3/3
100	1/3	0/3	0/3	0/3	0/3	0/3	0/3
10	0/3	0/3	0/3	0/3	0/3	0/3	0/3

a. The best hit for each resistance gene was used.

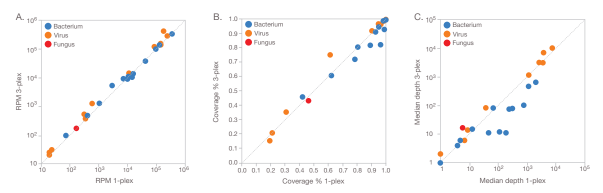


Figure 5: Consistent detection of respiratory pathogens at different plexities—All respiratory pathogens detected at 1-plex were also detected at 3-plex enrichment. Performance metrics, including (A) RPM, (B) coverage %, and (C) median depth were highly consistent between plexities.

Learn more

Learn more about using NGS for infectious disease detection at www.illumina.com/areas-of-interest/microbiology/infectious-disease-surveillance.html

Learn more about target enrichment detection of respiratory viruses with the MiniSeq Rapid Reagent Kit at www.illumina.com/content/dam/illumina/gcs/assembled-assets/marketing-literature/miniseqrapid-covid-app-note-470-2020-008/miniseq-rapid-covid-app-note-470-2020-008.pdf

Learn more about the IDbyDNA Explify Platform at www.idbydna.com/explify-platform/

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