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Emerging Molecular Assays for Detection and Characterization of Respiratory Viruses

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Rapid detection and identification of viral pathogens causing respiratory tract infections is critical for initiating antiviral therapy, avoiding unnecessary antimicrobial therapy, preventing nosocomial spread, decreasing the duration of hospitalization, and reducing management costs. Molecular assays, which provide high sensitivity and specificity, short test turnaround time, and automatic, high-throughput batch processing, have played critical roles in rapid detection, screening, and identification of emerging respiratory viral pathogens, such as severe acute respiratory syndrome coronavirus (SARS-CoV) and novel A/H1N1 influenza (Flu) virus.\textsuperscript{1–3} The superiority of polymerase chain reaction (PCR), reverse transcription-PCR (RT-PCR), and other in vitro nucleic acid amplification assays over conventional methods for the diagnosis of respiratory viral infections has already been established.\textsuperscript{4,5} This article describes several emerging molecular assays that have potential applications in the diagnosis and monitoring of respiratory viral infections.

DIRECT NUCLEIC ACID DETECTION BY QUANTUM DOTS BIOSENSORS

Biosensors offer the possibility of real-time monitoring, and the deployment of these devices in the field would provide a means for prompt etiologic diagnosis. All
biosensors are essentially composed of a biologic recognition element or bioreceptor, which interacts with the analyte and responds in some manner that can be registered by a transducer. The bioreceptor is a crucial component, and its function is to impart selectivity so that the sensor responds only to a particular analyte or biomolecule of interest, hence avoiding interference from other substances. The transducer converts the microbial biorecognition event into an electrical signal detected using electrochemical, optical, or piezoelectric platforms.\textsuperscript{6,7} A biosensor specifically targeting nucleic acids through hybridization is called a genosensor. Genosensors have been used to for direct, on-demand, and real-time detection and discrimination of microbial pathogens in clinical specimens. Malamud and colleagues\textsuperscript{8} developed a group of genosensor-based assays to detect microbial pathogens in oral specimens for use in the diagnosis of multiple infectious diseases. A piezoelectric DNA biosensor to directly detect hepatitis B virus was developed based on the mass-transducing function of a quartz crystal microbalance and nucleic acid hybridization\textsuperscript{9}, another hybridization-based amperometric biosensor, using osmium as an electrochemical indicator, was used for the detection and confirmation of virus-specific PCR products.\textsuperscript{10} A generic semidisposable fluorescence biosensor was developed to directly detect dengue virus RNA.\textsuperscript{11} A hybridization-based genosensor on gold film coupled with enzymatic electrochemical detection was designed to detect SARS-CoV RNA.\textsuperscript{12}

Fluorescent semiconductor nanocrystals, known as quantum dots (Qdots), are colloidal particles consisting of a semiconductor core, a high band gap material shell, and typically an outer coating layer. The core-size–dependent photoluminescence with narrow emission bandwidths that span the visible spectrum and the broad adsorption spectra allow simultaneous excitation of mixed Qdot populations at a single wavelength. Qdots also exhibit several unique features: high quantum yield, high resistance to photodegradation, and better near-infrared emission.\textsuperscript{13,14} The new generation of Qdots has far-reaching potential for the study of intracellular processes in broad fields, including diagnostics.\textsuperscript{14} High-sensitivity bacterial detection using biotin-tagged phage and quantum-dot nanocomplexes has been described, which provides specific limits of detection at 10 bacterial cells/mL in 1 hour.\textsuperscript{15} A bead-based microfluidic device was developed to achieve an ELISA with Qdots as the labeling fluorophore for virus detection.\textsuperscript{16} Three groups have reported the use of Qdots conjugated to specific monoclonal antibodies to detect and identify the presence of respiratory syncytial virus (RSV) in a real-time manner, implying that Qdots may provide a method for early, rapid detection of RSV infections.\textsuperscript{17–19} In addition to microbial pathogen antigen detection, positively charged compact Qdot-DNA complexes were described that can detect H5N1 Flu-A virus nucleic acids presented at concentrations as low as 200 nmol.\textsuperscript{20} Simultaneous excitation of several emission-tunable Qdot populations can be combined with a pool of differentially labeled probes for multiplex target analysis.\textsuperscript{21,22} Qdot-based techniques are under development to detect a panel of respiratory viruses, producing more efficient assays that require smaller quantities of target nucleic acids.

**AMPLIFICATION METHODS AND PLATFORMS**

**Loop-Mediated Isothermal Amplification**

First described by Notomi and colleagues\textsuperscript{23} in 2000, loop-mediated isothermal amplification (LAMP) is a simple, rapid, and specific nucleic acid amplification method, which is characterized by the use of multiple primers specifically designed to recognize several distinct regions on the target gene. Amplification and detection of target genes can be completed in a single step, by incubating the mixture of samples, primers, DNA polymerase with strand displacement activity and substrates at
a constant temperature. Because amplification is isothermal, LAMP does not require special reagents or sophisticated temperature control devices. Because the increase in turbidity of the reaction mixture according to the production of precipitate correlates with the amount of DNA synthesized, real-time monitoring of the LAMP reaction can be achieved by turbidity measurement. With a detection limit of about one to two copies, LAMP is capable of detecting the presence of pathogenic agents earlier than PCR if the gene copy number is low.

LAMP has successfully been applied to the rapid and real-time detection of several emerging and reemerging human pathogens, including West Nile virus, dengue virus, Japanese encephalitis virus, monkeypox virus, Rift Valley virus, SARS-CoV, Chikungunya virus, and noroviruses. Poon and colleagues described the use of an RT-LAMP to detect Flu-A viruses covering H1 to H3. Another similar RT-LAMP assay was described more recently that detects Flu-A virus H1 and H3 subtype strains and Flu-B virus strains. At a limit of detection of 10 focus-forming units per mL, both assays can be completed within 3 hours, providing rapid and sensitive detection. Two one-step RT-LAMP assays with analytical sensitivities of 0.01 to 0.1 plaque-forming units (pfu) per reaction were developed specifically for detection of highly pathogenic avian Flu-A (H5N1) viruses and validated using H5N1 viral strains isolated over the past 10 years and clinical specimens. An RT-LAMP assay was reported to specifically detect the H9 subtype of avian Flu virus with a detection limit of 10 copies per reaction, 10-fold lower than that of RT-PCR. In Japan, the LAMP assay was used to rapidly subtype Flu-A virus and confirm two cases of influenza in patients who had returned from Thailand.

In addition to the detection and typing of Flu viruses, a subgroup-A/B–specific RT-LAMP assay was developed to amplify RSV to improve current diagnostic methods for RSV infections. The assay was validated using nasopharyngeal aspirates from children who had respiratory tract infections, and the results indicated that the RT-LAMP is more sensitive than viral isolation and antigen testing for RSV detection. Several LAMP-based assays were reported for rapid detection of SARS-CoV with the advantages of rapid amplification, simple operation, and ease of detection. LAMP-based assays have also been used to detect other respiratory viral pathogens, such as mumps, measles, and adenoviruses. In comparison to conventional RT-PCR, RT-LAMP assays demonstrated 10- to 100-fold enhanced sensitivity, with a detection limit of 0.01 to 10 pfu of virus in most cases.

**Multiplex Ligation-Dependent Probe Amplification**

Recently established in The Netherlands, multiplex ligation-dependent probe amplification (MLPA) makes use of both ligation and PCR. Inventively modified from previously described ligation-dependent PCR assays, the MLPA platform features greatly reduced probe concentrations and longer hybridization periods to generate conditions compatible with multiplex analysis. Each MLPA probe consists of a pair of oligonucleotides subject to ligation when hybridized to a target sequence, analogous to a padlock probe (see later discussion). One oligonucleotide consists of a 5’ fluorescent label, a universal forward primer binding site, and a target-specific recognition sequence at the 3’ end, whereas the other oligonucleotide consists of a target-specific recognition sequence at the 5’ end, a nonspecific stretch of DNA of defined length (“stuffer” sequence), and a universal reverse primer binding site at the 3’ end. Each MLPA assay is divided into three basic steps: (1) annealing of probes to their target sequences, (2) ligation of the probes, and (3) PCR amplification of ligated probes using universal primers. Multiplexing is achieved by varying the length of stuffer sequence for each unique set of probes used in the assay. Amplification
products are detected using high-resolution electrophoretic techniques, such as
capillary electrophoresis, and it is claimed that this approach allows relative
quantification.42

MLPA-based techniques have proved sufficiently sensitive, reproducible, and
sequence specific for use in screening human DNA. Recent studies have use of the
MLPA assay for the detection and identification of several pathogenic microorgan-
isms, including rapid characterization of Mycobacterium tuberculosis,45 and relative
quantification of targeted bacterial species in oral microbiota.46 Reijans and
colleagues47 described an MLPA technology–based RespiFinder assay to detect 15
respiratory viruses simultaneously in one reaction. In this case, the MLPA reaction
was preceded by a preamplification step that ensured detection of both RNA and
DNA viruses with the same specificity and sensitivity as individual monoplex real-
time RT-PCR assays. The RespiFinder assay showed satisfactory specificity and
perfect sensitivity for adenovirus, human metapneumovirus (hMPV), Flu-A, parain-
fluenza virus (PIV) types 1 and 3, rhinovirus (RhV), and RSV. Use of the RespiFinder
assay resulted in a 24.5% increase in the diagnostic yield compared with cell culture.
This assay is being extended to cover four additional bacterial pathogens that cause
respiratory tract infections: Mycoplasma pneumoniae, Chlamydophila pneumoniae,
Legionella pneumophila, and Bordetella pertussis.

Polymerase Chain Reaction Amplification Using Arbitrary Primers

PCR amplification techniques using arbitrary primers, including arbitrarily primed (AP)
PCR,48 sequence-independent single-primer amplification (SISPA),49 and randomly
amplified polymorphic DNA (RAPD),50 are generally based on the PCR amplification
of random DNA segments with short primers (usually a single 1 of 10 nucleotides) con-
taining arbitrary nucleotide sequences. RAPD-based assays have increasingly been
used to type microorganisms, especially during clinical outbreaks.51 The RAPD-
PCR technique seems to be practical and efficient for routine use in high-resolution
viral diversity studies by providing assemblage comparisons through fingerprinting,
probing, or sequence information.52 Similar techniques have been used to charac-
terize the polymerase gene and genomic termini of Nipah virus53 and avian Flu virus
genome sequences.54

On the other hand, AP-PCR and SISPA-based assays have mainly been used for the
discovery and characterization of novel and noncultivatable viruses.55 Because viral
pathogens do not possess conserved, universal genes, such as 16S rRNA genes, SIS-
PA was used in the early 1990s as a random PCR amplification strategy to amplify
known and unknown viral genes, including those of hepatitis C virus, rotavirus, and
norovirus.56–58 The AP-PCR technique was used successfully to obtain sequence
information on a novel hMPV after the virus was cultured.59 Wang and colleagues60,61
used a similar random amplification technique in conjunction with a long oligonucleo-
tide pan-viral microarray to simultaneously screen and detect hundreds of viral path-
ogens. This system has successfully been used for the detection of a human PIV-4
strain associated with respiratory failure,62 for identification of a novel gammaretrovi-
rus in a patient who had prostate tumors,63 for the diagnosis of a critical respiratory
illness caused by hMPV,64 and for the identification of cardioviruses related to Theiler
murine encephalomyelitis virus in human infections.65 Quan and colleagues66 recently
reported the use of a similar random amplification process followed by comprehensive
microarray analysis (GreeneChipResp) to detect diverse respiratory viral pathogens
and subtype Flu-A viruses.

A modified SISPA incorporating DNAse treatment has recently been used to
discover, identify, and characterize several novel bovine and human viral pathogens

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directly from clinical samples. The same technology has been used for the character-
ization of common epitopes in enterovirus (EnV), identification of a novel human
coronavirus, detection of TT virus in stool samples collected during a gastroenteritis
outbreak, and discovery of novel unculturable viruses in specimens collected from
patients presenting with fever of unknown origin. Although PCR amplification
using arbitrary primers has been an extremely powerful approach for screening and
discovery of new or noncultivable viral pathogens directly from clinical specimens,
subsequent identification and confirmation steps are hindered by a background of
nonspecific random amplification products. Further development is thus required to
optimize this technology for routine diagnostic use in molecular microbiology
laboratories.

**Target-Enriched Multiplexing Amplification**

Multiplex PCR was developed to use numerous primers within a single reaction tube to
amplify nucleic acid fragments from different targets. Multiple sets of high-concentra-
tion primers in the conventional multiplex reaction often favor primer-dimer formation,
however, resulting in nonspecific amplification. To meet the challenges of conventional
multiplex PCR, Han and colleagues developed target-enriched multiplexing (TEM)-
PCR technology, which uses nested gene-specific primers at extremely low concentra-
tions to enrich specific targets during early PCR cycles and relies on universal forward
and reverse “superprimers” at high, but unequal, concentrations to achieve exponen-
tial asymmetric target amplification. TEM-PCR amplification has been reported for the
detection, typing, and semiquantification of 25 human papillomaviruses, detection
and differentiation of a panel of respiratory bacterial pathogens, detection and
and differentiation of 24 antituberculosis drug resistance-related mutations, determina-
tion of antibiotic resistance and detection of toxin-encoding genes in Staphylococcus
aureus, screening and differentiation of methicillin-resistant *S. aureus* and vancomy-
cin-resistant enterococci, and characterization and typing of Flu-A, including H5N1.

Using TEM technology, the ResPlex II assay was developed to detect Flu-A, Flu-B,
PIV-1, PIV-2, PIV-3, PIV-4, RSV, hMPV, RhV, EnV, and SARS-CoV in a single reac-
tion. When monoplex RT PCR is used for pathogen detection, the clinician often
does not consider the possible presence of other pathogens when given a positive
result. The multiplex approach offered by the ResPlex II system enhances diagnosis
through detection of respiratory viral etiologic agents in cases in which their presence
was unsuspected and an appropriate test consequently was not ordered by the clini-
cian. A recent study by Brunstein and colleagues revealed that, using the ResPlex
II kit covering 12 viral pathogens, 2.5% of specimens were coinfected with two or
three different viruses. (A low level of cross-reactivity between PIV-1 and PIV-3 was
noticed using this assay.) These coinfections are medically relevant, and effective
treatment of severe respiratory tract infections will increasingly require diagnosis of
all involved pathogens, as opposed to single-pathogen reporting. The original Re-
spLex II system detects only RNA viruses, but adenoviruses, bocavirus, and four coro-
naviruses have been added to a recently released new version of ResPlex II. Preliminary data indicate that the overall sensitivity and specificity of ResPlex II v2.0
is comparable to that of the ResPlex II panel. A notable number of previously negative
samples were found to be positive for one of the newly added bocavirus or corona-
virus targets (John Brunstein, 2009; personal communication). A factor that could
diminish the analytical and clinical performance of ResPlex II and ResPlex II v2.0 is
the potential for false-positive results caused by carryover of PCR products using
the Luminex platform.
Direct amplicon sequencing provides simple, rapid, and accurate means of detection and identification of amplification products. The need for robust, high-throughput methods to replace the elegant Sanger method, which was described more than 30 years ago, has led to the development of several new principles. Ronaghi and colleagues described in 1998 a pyrosequencing technique, a non–gel-based real-time approach to sequencing DNA by monitoring DNA polymerase activity. Pyrosequencing is based on enzymatic inorganic pyrophosphate release by DNA polymerase. This reaction is stoichiometric; the amount of light produced is proportional to the number of pyrophosphate molecules generated and, hence, the number of incorporated nucleotides. Unincorporated nucleotides are degraded with apyrase before the next nucleotide is added. In this way, sequence information on an interrogated region is generated quantitatively in real time. Although basic approaches to performing pyrosequencing remain the same, numerous commercial systems have been used widely to rapidly identify infectious agents and screen for antimicrobial drug resistance. Multiplexed pyrosequencing involving the simultaneous extension of several primers hybridized to one or more target DNA templates has gained broad acceptance in the fields of cytogenetics, pharmacogenetics, and medical genetics.

Most applications of pyrosequencing in the identification and characterization of respiratory viruses have focused on Flu-A. Based on pyrosequencing technology, a rapid and highly informative diagnostic assay was reported for the detection of H5N1 Flu viruses; sequencing of critical regions within the H5 virus was developed as a screening method during high volumes of H5N1 activity. A real-time RT-PCR pyrosequencing assay was developed that combines restriction enzyme digestion and direct sequencing to screen and verify H5 Flu infections in humans. Another RT-PCR assay with subsequent pyrosequencing analysis allows for a rapid, high-throughput, and cost-effective screening of subtype A/H1N1, A/H3N2, and A/H5N1 viruses and can clearly discriminate wild-type from a mutant viruses. A study reported by Bright and colleagues showed an alarming increase in the incidence of amantadine- and rimantadine-resistant H3N2 Flu-A viruses worldwide when the pyrosequencing technique was configured to cover a 44-base pair region of the M2 protein-encoding gene. Pyrosequencing assay capabilities were expanded to screen for 52 amino acid changes defined as avian or human specific, and pyrosequencing-based assays recently were designed for detection and surveillance of the most commonly reported mutations associated with resistance to neuraminidase inhibitors and the adamantanes. The latter detects mutations associated with resistance directly in clinical specimens, thus reducing the time required for testing and avoiding selection of novel sequence variants by cell culture. In addition, pyrosequencing-based assays have been reported for the characterization, quantification, typing, subtyping, and drug-resistance profiling of other viruses.

One unique feature of pyrosequencing is its theoretical adaptability to the analysis of any genetic marker, which allows for the detection of multiple known and unknown mutations in a single pyrosequencing reaction. Integration of high-throughput pyrosequencing with the Roche/454 instrument has become a powerful tool for whole genome sequencing without the need for additional equipment or molecular techniques other than standard PCR, Genome Sequencer FLX sample preparation, and the sequencing pipeline. Pyrosequencing generates sequence content quantitatively, which has made pyrosequencing a primary choice for quantifying specific mutations (eg,
detection of drug resistance–associated signatures) in mixed genomic populations. Because pyrosequencing byproducts inhibit the sequencing reaction, pyrosequencing read lengths are limited to less than 100 base pairs. Another drawback of pyrosequencing-based techniques includes secondary structure formation, which affects quality of the results, particularly with GC-rich targets. Additionally, it may be difficult to determine the precise number of nucleotides in a homopolymeric region based on peak heights.87 It is expected that pyrosequencing-based diagnostic devices will soon become available for rapid characterization and typing of viral pathogens.

**Padlock Probes**

Padlock probes, originated by Nilsson and colleagues114 in 1994, are linear oligonucleotides designed so that the two end segments, connected by a linker region, are both complementary to a target sequence. On hybridization to a target sequence, the two probe ends become juxtaposed and can be joined by a DNA ligase. Reacted probes can be detected by way of reporter molecules attached to the linker.115 Alternatively, an amplified signal can be obtained from the circularized probes by rolling circle amplification. Padlock probes provide a means for detection and quantification of large numbers of DNA or RNA sequences and for highly multiplexed genetic studies.116 The application of padlock probes for the detection of microbial pathogens is a recent trend in molecular diagnoses.117

The unique padlock probe design provides the benefit of speed and sensitivity derived from using a nucleic acid–based method, and the amount of information is greatly increased by extensive multiplexing. Indeed, this method was used to simultaneously detect and type 16 HA and 9 NA subtypes of avian Flu virus. The analysis is completed within approximately 4 hours and performed in a single reaction tube, which helps to decrease the risk for contamination, with just a few sequential additions of reagents before the readout is performed using an oligonucleotide array.118 Padlock probes combined with back-end microarray technology have been developed to detect foot-and-mouth disease, vesicular stomatitis, and swine vesicular disease viruses.119 Besides viral pathogens, padlock probe–based techniques have been rapidly extended in recent years to the identification and characterization of bacterial and fungal pathogens.120–124 In addition to the applicability of padlock probes for direct target detection, a universal primer binding site can be introduced into the probe and used for MLPA (see previous discussion).

**Microarrays**

Applications of microarrays to detect and characterize respiratory viruses began with solid arrays. The first respiratory pan-viral microarray system was described in 2002, which incorporated 1600 unique 70-mer oligonucleotide probes covering approximately 140 viral genome sequences.60–65 Resequencing microarrays were developed to use short oligonucleotides for the simultaneous identification of respiratory pathogens at both the species and strain level.125–127 Another comprehensive and panmicrobial microarray, the GreeneChipResp system, was developed for the detection of respiratory viruses and subtype identification of Flu-A viruses.66 Other recently developed solid microarray systems for detection and identification of a panel of respiratory viruses include the Infiniti analyzer, an integrated molecular diagnostic device incorporating microarray hybridization128; the electronic microarray-based Nano-chip85,129, the TaqMan Low Density Array cards, which use real-time PCR assays for 13 viruses and 8 bacteria known to cause pneumonia (Dean Erdman, 2009; personal communication); and the FilmArray, which detects and differentiates 17 viral
4 bacterial etiologies of respiratory tract infections (Mark Poritz, 2009; personal communication).

Suspension bead-based liquid xMAP microarrays have been developed by Luminex Corp, which are essentially three-dimensional arrays based on the use of microscopic polystyrene beads as the solid support and flow cytometry for bead and target detection.

Robust multiplexing detection is accomplished using different bead sets based on fluorescence. The system enables multiplexing of up to 100 analytes in a single reaction using small sample volumes. Numerous studies have described the use of xMAP technology for the detection and differentiation of nucleic acid sequences of microbial pathogens, including enteric bacteria, viruses, mycobacteria, fungi, and protozoa. A molecular typing method incorporating the suspension array was reported to characterize and type Flu-A viruses, including H5N1. The Luminex suspension array has been incorporated into several commercial devices as the detection platform to support the laboratory differential diagnosis of common respiratory viral pathogens. These include the xTAG Respiratory Viral Panel from Luminex Molecular Diagnostics, the ResPlex II assay from Qiagen, and the MultiCode-PLx RVP assay from EraGen Biosciences. The suspension array system exhibits rapid hybridization kinetics, flexibility in assay design and format, and relatively low costs, which have made it the most practical microarray platform for clinical diagnostic applications. Users should carefully determine the positive fluorescence threshold for each viral target in multiplexed, user-defined assays during validation.

Mass Spectrometry

Matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF) mass spectrometry (MS) is widely used as a powerful proteomic tool. Its rapidity and high resolution provide another powerful platform for the detection and characterization of nucleic acid amplification products. The technology is premised on the capacity of MALDI-TOF MS to discriminate individual PCR products contained in complex amplification mixtures according to nucleotide base composition. The deconvolution algorithm allows base composition of PCR products to be deduced from mass spectrometrically measured molecular weights and the complementary nature of DNA, leading to organism identification. Early studies successfully used this technique to directly detect amplification products from PCR and ligase chain reaction (LCR). Soon after, the MALDI-TOF MS platform was linked to PCR amplification for genotypic analysis of hepatitis C virus and human papillomavirus. Detection of human herpesviruses from clinical specimens was performed using MALDI-TOF MS following multiplex PCR amplification. A MALDI-TOF MS-based genotyping assay has been described that monitors development of hepatitis B virus polymerase YMDD mutant genotypes during lamivudine treatment.

An integrated system, the Ibis T5000 Biosensor, has been developed to couple broad-range nucleic acid amplification to high-performance electrospray ionization MS and base-composition analysis. The system enables the identification and quantification of a broad set of pathogens, including all known bacteria, all major groups of pathogenic fungi, and the major families of viruses that cause disease in humans and animals, along with the detection of virulence factors and antibiotic resistance markers. The system has been used for rapid identification and strain typing of respiratory bacterial pathogens for epidemic surveillance, identification and genotyping of Acinetobacter baumannii strains in an outbreak associated with war trauma, determination of quinolone resistance in Acinetobacter species, genotyping of Campylobacter species, and rapid genotyping and clonal complex assignment of Staphylococcus aureus isolates. We have used this system
| System                                      | Company                      | Viruses/Genotypes Detected                                                                 | Amplification Platform | Detection Platform         | Characteristics                                                                 |
|---------------------------------------------|------------------------------|---------------------------------------------------------------------------------------------|------------------------|---------------------------|--------------------------------------------------------------------------------|
| FimArray respiratory pathogen panel         | Idaho Technology Inc (Salt Lake City, UT) | AdV, bocavirus, 4 CoV, Flu-A, Flu-B, hMPV, PIV-1, PIV-2, PIV-3, PIV-4, RSV, and RhV        | Nested multiplex RT-PCR | Solid array analyzer      | Integrated and closed system. Also covers 4 bacterial pathogens               |
| Infiniti respiratory viral panel            | AutoGenomics, Inc (Carlsbad, CA) | Flu-A, Flu-B, PIV-1, PIV-2, PIV-3, PIV-4, RSV-A, RSV-B, hMPV-A, hMPV-B, RhV-A, RhV-B, EnV, CoV, and AdV | Multiplex PCR and RT-PCR | Infiniti solid array analyzer | Detection step by the Infiniti analyzer is completely automatic               |
| Jaguar system                               | HandyLab, Inc (Detroit, MI)   | Flu-A, Flu-B, and RSV A/B                                                                  | Multiplex real-time RT-PCR | Melting temperature analysis | Completely closed and automatic. Universal system compatible with detection of other pathogens. Throughput of 1–24 specimens/run |
| MultiCode-PLx respiratory virus panel       | EraGen Biosciences (Madison, WI) | Flu-A, Flu-B, PIV-1, PIV-2, PIV-3, PIV-4, RSV, hMPV, RhV, AdV, and CoV                    | Multiplex PCR and RT-PCR | Luminex suspension array   | Universal beads used for detection use EraCode sequences                     |
| NGEN Respiratory Virus (RVA) Analyte-specific reagent | Nanogen (San Diego, CA) | Flu-A, Flu-B, PIV-1, PIV-2, PIV-3, and RSV | Multiplex RT-PCR | NanoChip (solid chip) | Discontinued in 2008. Probe labeling, target capture, and detection accomplished using electronic microarray technology |
| ProFLU+, ProPARAFLU +<sup>161,162</sup> | Prodesse, Inc (Waukesha, WI) | Flu-A, Flu-B, and RSV (ProFLU+); PIV-1, PIV-2, PIV-3, and PIV-4 (ProPARAFLU+) | Multiplex real-time RT-PCR | Melting temperature analysis | ProFLU+ FDA cleared. Limited multiplex formats (triplex) |
|---|---|---|---|---|---|
| ResPlex II<sup>78,84,85</sup> | Qiagen (Valencia, CA) | Flu-A, Flu-B, PIV-1, PIV-2, PIV-3, PIV-4, RSV-A, RSV-B, hMPV, RhV, EnV, and SARS-CoV | TEM-RT-PCR | Luminex suspension array | Unique Tem-PCR permits multiple target screening in single reaction without significant loss in sensitivity |
| Seeplex respiratory virus detection assay<sup>163</sup> | Seegene, Inc (Seoul, Korea) | Adv, hMPV, 2 CoV, PIV-1, PIV-2, PIV-3, Flu-A, Flu-B, RSV-A, RSV-B, and RhV | Two sets of multiplex RT-PCR | Gel electrophoresis | Dual priming oligonucleotide system |
| xTAG respiratory viral panel (RVP)<sup>137–139</sup> | Luminex Molecular Diagnostics (Toronto, Canada) | Flu-A, Flu-B, PIV-1, PIV-2, PIV-3, PIV-4, RSV-A, RSV-B, hMPV, Adv, EnV, CoV, and RhV | Multiplex PCR and RT-PCR | Luminex suspension array | FDA cleared. Target-specific primer extension used in combination with universal detection beads |

Abbreviations: Adv, adenoviruses; CoV, coronaviruses; EnV, enteroviruses; Flu, influenza virus; hMPV, human metapneumovirus; PIV, parainfluenza virus; RhV, rhinoviruses; RSV, respiratory syncytial virus; TEM, target enriched multiplex.
to detect *Ehrlichia*, *Anaplasma*, and *Rickettsia* pathogens directly from blood specimens for diagnosis of tick-borne sepsis (manuscript in preparation). In the field of diagnostic virology, this strategy successfully led to the inclusion of SARS-CoV in the coronavirus family.\textsuperscript{158} Furthermore, the Ibis T5000 Biosensor system has been used as a rapid and inexpensive tool for global surveillance of emerging Flu virus genotypes\textsuperscript{159} and rapid detection and molecular serotyping of adenoviruses.\textsuperscript{160} The system was able to detect and type all available Flu A genotypes, including recently emerged novel A/H1N1 (David Ecker, 2009; personal communication). The main advantages are high resolution, speed, and substantial degree of automation. The main disadvantages include the engineering difficulty of MS device miniaturization and need for continuous enrichment of databases with new genomic sequences.

**MULTIPLEXING AMPLIFICATION AND HIGH-THROUGHPUT DETECTION SYSTEMS**

Respiratory infections caused by a many bacterial, viral, and fungal pathogens often present with overlapping signs and symptoms nearly indistinguishable by clinical diagnosis. Molecular screening of at-risk populations for a group of possible viral pathogens is an exciting area of development in molecular microbiology. Several multiplexing amplification and high-throughput detection systems are commercially available for the detection and differentiation of a panel of respiratory viral pathogens. Examples include the FilmArray platform from Idaho Technology Inc; the Infiniti Respiratory Viral Panel from AutoGenomics, Inc.\textsuperscript{128}; the Jaguar system from HandyLab, Inc.; the Multi-Code-PLx respiratory virus panel from EraGen Biosciences\textsuperscript{140,141}; the NGEN Respiratory Virus ASR from Nanogen\textsuperscript{85,129}, the proFLU+ and the proPARA-FLU+ from Prodesse, Inc.\textsuperscript{161,162}; the ResPlex II assay from Qiagen\textsuperscript{78,84,85}; the Seeplex respiratory virus detection assay from Seegene, Inc.\textsuperscript{163}; and the xTAG Respiratory Viral Panel from Luminex Molecular Diagnostics\textsuperscript{137–139}. Some of these systems cover all varieties of Flu A genotypes including recently emerged novel A/H1N1.\textsuperscript{164}

A comparative summary of these devices is presented in Table 1. Relative simplicity, powerful multiplexing capabilities, and affordability for high-throughput detection make these platforms most attractive for screening and detection of a panel of respiratory viruses in clinical infectious disease diagnostics. Although not essential, the availability of Food and Drug Administration–cleared products is a critical step in getting these systems into less-experienced diagnostic microbiology laboratories. Opening of postamplification tubes and subsequent pipetting steps in the workflow of suspension arrays increases the risk for intra- and inter-run contamination for some assays. Careful attention should be paid to contamination control measures and the re-establishment of dedicated postamplification laboratory space in the real-time PCR era. Simultaneous testing for all possible pathogens is an efficient means to obtain a conclusive result and improves etiologic diagnosis.\textsuperscript{81,137,165} In addition, assaying for all potential pathogens may yield crucial information regarding coinfections or secondary infections.\textsuperscript{84,166,167} One study from the Netherlands indicated that implementation of multiple molecular assays for the etiologic diagnosis of lower respiratory tract infections increased the diagnostic yield considerably, yet did not reduce antibiotic use or costs.\textsuperscript{168} Clinical relevance and cost effectiveness of simultaneous multipathogen detection and identification strategies merit further investigation.

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