Laboratory Detection of Respiratory Viruses by Automated Techniques

Mercedes Pérez-Ruiz*, Irene Pedrosa-Corral, Sara Sanbonmatsu-Gámez and José-María Navarro-Marí

Laboratorio de Referencia de Salud Pública para Enfermedades con Suspecha de Etiología Virica en Andalucía (Consejería de Salud), Servicio de Microbiología, Hospital Universitario Virgen de las Nieves, Avda, Fuerzas Armadas, 2, 18014 Granada, Spain

Abstract: Advances in clinical virology for detecting respiratory viruses have been focused on nucleic acids amplification techniques, which have converted in the reference method for the diagnosis of acute respiratory infections of viral aetiology. Improvements of current commercial molecular assays to reduce hands-on-time rely on two strategies, a stepwise automation (semi-automation) and the complete automation of the whole procedure. Contributions to the former strategy have been the use of automated nucleic acids extractors, multiplex PCR, real-time PCR and/or DNA arrays for detection of amplicons. Commercial fully-automated molecular systems are now available for the detection of respiratory viruses. Some of them could convert in point-of-care methods substituting antigen tests for detection of respiratory syncytial virus and influenza A and B viruses. This article describes laboratory methods for detection of respiratory viruses. A cost-effective and rational diagnostic algorithm is proposed, considering technical aspects of the available assays, infrastructure possibilities of each laboratory and clinic-epidemiologic factors of the infection.

Keywords: Acute respiratory infection, automation, influenza-like illness, microarray, real-time PCR, respiratory viruses.

1. INTRODUCTION

Acute respiratory infection (ARI) of viral aetiology is one of the main causes of morbidity and mortality worldwide [1-4]. Besides classical respiratory viruses (RV), i.e. influenza viruses (Flu), respiratory syncytial virus (RSV), human rhinovirus (HRV), parainfluenza virus (PIV) 1-4, adenovirus (ADV), enterovirus (EV) and human coronavirus (HCoV) OC43 and 229E [5-9], other “new” RV have currently been added to this list, such as human metapneumovirus (hMPV) [10], SARS-CoV [11], HCoV NL63 and HKU1 [12,13], human bocavirus (hBoV) [14] and others. Most RV cause benign and self-limited infections in immunocompetent adults. However, clinical impact of ARI increases when they affect children, elderly, immunocompromised individuals and/or subjects with chronic underlying diseases, in whom RV constitute an important cause of hospitalisations, mainly during the cold months of the year [15,16]. They involve important costs for health systems and constitute a frequent cause of school and work absenteeism. In certain circumstances, emerging RV cause a great impact in health systems worldwide, as occurred with SARS-CoV and Flu pandemics, the last one taking place in 2009.

Aetiological diagnosis of viral ARI is crucial in order to avoid unnecessary antibiotic use, to establish the appropriate use of antiviral drugs and a comprehensive cohort of hospitalised patients that minimizes the risk of nosocomial transmission, and to provide epidemiological information for an early release of recommendations for the prevention and treatment and, finally, to reduce the overall costs derived from patients’ management.

Several diagnostic methods have been described for RV. They can be classified in direct detection methods, which detect the virus or a part of it in the clinical sample, including viral culture, antigen detection methods and molecular techniques; and indirect methods, based on the detection of immune response to the virus by serologic assays. In clinical practice, serology is not useful for the diagnosis of viral infections, such as ARI, that cause frequent recurrences and are mainly restricted to the respiratory mucosa. Thus, in these cases, direct detection of the virus in the site of infection has to be a priority [16,17].

Advances in diagnostic virology for detecting RV have been focussed on nucleic acids amplification techniques (NAAT), since they are more sensitive than traditional methods such as viral culture or antigen detection assays. Moreover, NAAT are the only available method for most new RV because they do not grow or grow poorly in cell culture and for which no efficient antigen detection methods have been commercialised. NAAT offer additional advantages with respect to other classical methods as less turnaround time and less effect of virus viability on the efficiency of the assay [18].

On the contrary, some disadvantages have been attributed to NAAT: increased reagents’ costs, higher hands-on time, the need of knowing at least a part of the target genome to design appropriate primers and/or probes. Indeed, many RV are RNA viruses, subjected to a greater genetic variability that may affect the sensitivity and reliability of the molecular method.

Currently, many commercial molecular assays have been developed that allow partial or complete automation of the
procedure. Semi-automated or automated assays have solved in part one disadvantage of NAAT, which is the hands-on time. Many commercial NAAT require minimal management of the specimen, taking no more than a few minutes.

With the increasing availability of easy-to-perform and multiplex NAAT for detecting RV, cost-effective diagnostic algorithms should be used. Whether a more or less complete virological study should be carried out on a certain patient’s sample, depends not only on the resources and type of laboratory, but also on clinical-epidemiologic aspects such as patient’s age, severity of the disease, underlying conditions, etc.

This article describes the molecular methods used to detect respiratory viruses and the contribution that increasing automation, especially on molecular assays, has made to improve laboratory diagnosis. The possibilities of integration of these new assays on each laboratory portfolio are also discussed.

2. TRADITIONAL DETECTION METHODS

Traditional direct methods for RV are viral culture and antigen detection assays. Among indirect methods, many serologic assays, such as EIA, haemagglutination inhibition, complement fixation, etc. have been used for RV. However, serology is not useful in clinical practice since diagnosis must be achieved by demonstration of seroconversion or seroincrease of IgG antibodies, between acute-phase and convalescent-phase serum samples [16,18].

In past decades, viral culture was the gold standard for the aetiological diagnosis of RV. No automation has been reported for this technology. Rather, advances in this field of clinical virology have been focused on shortening detection times and on increasing sensitivity. Currently, this procedure is limited to reference laboratories that are provided with the appropriate infrastructure, with the main objective of recovering the viral strain for further genetic and/or antigenic characterisation.

Traditional viral culture requires at least 7 days for yielding a result. To solve this problem, the next step in the viral culture methodology was the introduction of the shell-vial assay, based on centrifugation-enhanced adsorption of the sample onto the cells monolayer reproduced on cover slips. Detection is commonly achieved by fluorescent assay with monoclonal antibodies on the cell monolayer. This method reduces the time to obtain a result to 24-48 h without a deleterious effect on sensitivity and specificity.

Although viral culture may represent an open system for detecting multiple viral pathogens, there is no single cell line that allows the growth of all RV, and there is no RV that may grow in any cell line. Thus, to detect the most prevalent RV, multiple cell lines must be used. The combination of shell-vial assay with the inclusion of different cell lines in the same vial has represented a subsequent step that has facilitated the recovery of RV isolates from cell cultures [19-21]. NAAT have recently contributed to increase the sensitivity of viral culture. PCR assays can be carried out on the cell culture supernatant to detect the isolates. This approach has demonstrated the highest sensitive reading procedure, and has allowed identifying viral isolates in cell monolayers with no cytopathic effect [22]. However, viral culture is not useful for detection of new RV, since they do not grow or grow poorly in cell culture. For this purpose, the best alternative is NAAT.

The availability of monoclonal antibodies has allowed the development of different technologies for detecting viral antigens from clinical samples. Fluorescent and chromatographic immunoassays are rapid, easy to perform and have proved their utility, mainly for RSV and Flu. The minimum (hands-on and hands-off) turnaround time that these techniques require for obtaining results have converted them as referral point-of-care diagnostic assays. However, sensitivity widely depends on the commercial method and on the type of patient and sample, ranging from 10% to 70% in the case of pandemic Flu [23]. Higher viral loads and thus, higher sensitivity of antigen detection methods for RSV and Flu have been observed in the paediatric population versus adults and in nasopharyngeal aspirates versus nasopharyngeal exudates [24,25].

Recently, the goal of automation in these type of diagnostic tools has been achieved by a new commercial assay, the mariPOC® respi test (ArcDia, Turku, Finland). This allows rapid detection and differentiation between viral and bacterial respiratory infections (Flu A, Flu B, RSV, hMPV, PIV 1, PIV2, PIV3, ADV and Streptococcus pneumoniae). The test is very simple to operate and hands-on-time is less than a minute per sample. Results for highly positive samples are obtained in 20 minutes whereas low positive and negative samples are reported in 2 hours.

The mariPOC® system uses polystyrene microparticles as solid phase reaction carriers for the immunocomplex formation. Viral antigens are captured by specific antibodies fixed onto the microsphere and bind to a fluorescently-labelled conjugate. The amount of immunocomplexes formed on the microsphere is proportional to the analyte concentration. Detection of the fluorescence signal is carried out by separation-free two-photon excitation fluorescence detection assay, the ArcDia™ TPX (TPX). In contrast to other antigen detection methods, TPX assay avoids the need for multiple washes, and readout is achieved by on-line monitoring of reaction kinetics [26]. Preliminary studies have reported a sensitivity and specificity of this method comparable to ultrasensitive fluorescent immunoassays (DELPHIA® method) [27,28] and offers the advantage of automated result readout, allowing connections to laboratory information systems.

3. NAAT FOR DETECTION OF RESPIRATORY VIRUSES: FROM THE SIMPLER TO THE MOST COMPLEX TARGETING APPROACH

NAAT, particularly the polymerase chain reaction (PCR), have expanded the range of pathogens that can be identified in clinical laboratories. They have shown their utility in clinical diagnosis of viral respiratory infections. Molecular assays have greater sensitivity than conventional methods as viral culture or antigen detection and indeed, they are able to detect new viral agents. Another advantage of NAAT compared with viral culture is that they do not require viable viruses. Thus, they have converted in the gold standard for detection of RV. Thus, most advances in clinical diagnostic virology have been made on NAAT.
NAAT are especially useful for detecting “low-positive” patients. Yearly, during periods of RSV and Flu epidemics, the availability of NAAT eases the management of hospitalised patients and the control of outbreaks, since antigen detection tests are not sensitive enough for certain population groups, such as immunosuppressed individuals.

Flu, HRV, PIV 1, PIV 3, PIV 4, ADV and hCoV (except SARS-CoV) usually cause upper respiratory tract infections (URTI), whereas RSV, hMPV and PIV 2 have been frequently associated to lower respiratory tract infections (LRTI) [18].

Most RV are RNA viruses except ADV. Two alternatives are feasible for reverse transcription (RT) of viral RNA into its appropriate cDNA PCR target. One approach is the use of random and/or oligo(dT) primers in the RT step. Subsequently, PCR can be directed against any RV with specific primers. The main disadvantage of this procedure is the need for opening the RT tube to carry out the PCR. One-step RT-PCR protocols have been reported elsewhere for most RNA viruses, and solve the problem the former protocol has. These are based on the inclusion in the reaction mix of reverse transcriptases and DNA polymerases for both the RT and the PCR steps, or of enzymes with reverse transcriptase and DNA polymerase activities.

Monoplex PCRs have been published and commercial kits have been developed for detecting the most prevalent RV [18]. Whether a simpler or a more complete virological study must be carried out depends on many factors: severity of the disease, type of patient, capacity of the laboratory, availability of validated and optimized trade-marked commercial kits, number of samples that the laboratory can process, etc.

Nevertheless, in many cases, clinical signs and symptoms of ARI are not specific of a single RV, and mild and severe infections have been described for all viruses mentioned above. Currently, monoplex PCR is not useful in a clinical laboratory equipped enough to conduct an extended aetiological diagnosis. However, the advent of the 2009 pandemic allowed many laboratories for equipping with the infrastructure necessary to carry out PCR assays for detection of pandemic Flu. In these circumstances, when a rapid answer is encouraging, monoplex PCRs provide an efficient response.

Multiplex PCR assays have been the most widely used molecular methods for RV detection in laboratories of clinical virology capable of developing in-house and/or robust published protocols. The advantage with respect to monoplex PCR is the cost savings compared to multiple monoplex PCRs and an easier processing of runs. Indeed, multiplex PCR extends the portfolio of virological diagnostic methods. Although advantages are strong enough to compensate for any disadvantage, multiplex PCR have intrinsic inconveniences: as the number of targets increase, the possibility of losing sensitivity and specificity increases as well, due to competition among primers and viral targets that consequently yields unspecific PCR products. Indeed, multiplex PCR has to be optimized to ensure the adequate amplification of each viral genome. Multiplex PCR assays have demonstrated that co-detections of RV in a sample are a common event [29]. Whether these co-detections express true infections by all viruses and/or prolonged shedding of certain agents is difficult to demonstrate, although some authors have pointed out the role of co-detections of RV in the severity of the disease [30]. Table 1 shows the most important advantages and disadvantages of different methods for detection of RV.

4. AUTOMATION POSSIBILITIES FOR DETECTING RESPIRATORY VIRUSES BY NAAT

Automation in molecular assays, aiming at a continuous reduction of hands-on time, yields higher reproducibility and offers possibility of detecting several pathogens with a minimum processing.

The availability of automated procedures for detecting RV is increasing day by day. This obliges the virologists to choose the best alternative taking into account laboratory throughput and epidemiologic, clinical and cost-effectiveness parameters since, in many instances, each commercial assay requires its own apparatus. Indeed, commercial diagnostic kits are less versatile and often, more expensive than well-optimized in-house PCR.

Molecular procedures for achieving automation goals for RV detection could be divided in two main straightforward phases: procedures and protocols for a stepwise automation (semi-automation) and fully-automated NAAT.

4.1. Stepwise Automation (Semi-Automation)

From nucleic acids extraction to the readout of PCR results, many efficient automated systems covering all steps of the whole procedure have emerged in the last 10-15 years. By chronological order, automated extractors, real-time PCR instruments, DNA arrays (and other systems) for detection of PCR products, and recent platforms for nucleic acids purification and preparation of the PCR master mix, are the main contributions to a semi-automation of molecular methods.

4.1.1. Automated Nucleic Acids Extraction Methods

Several commercial assays offer automated solutions for nucleic acids extraction, isolation and purification that range from low to high throughput. The high quality of nucleic acids obtained with these systems are suitable for a broad range of applications in nucleic acid research, including gene expression analysis (using real-time PCR or conventional PCR on standard block cyclers), microarray analysis, genotyping reactions, and many other technologies.

These instrument purify viral DNA and RNA from a broad range of respiratory samples (nasopharyngeal aspirates and swabs, sputum, bronchoalveolar lavages, ...). Contaminations are reduced since the system is closed, which avoids splashing and generation of aerosols and drops. Many of these systems include ready-to-run, prefilled cartridges with all reagents required for the purification procedure. Setup is rapid and saves time. Commercial assays for the simultaneous processing of a maximum of 8 up to 96 samples are available.

Specific studies on the efficiency of these methods in recovery of DNA and RNA from RV have been carried out [31-34]. Besides its higher reproducibility, the amount of nucleic acids recovered from automated extraction methods is

greater than the amount achieved with manual methods [31]. Overall, a good recovery of viral RNA and DNA, without significant differences among automated extractors, has been found. No evidence of the presence of inhibitors is observed as well. However, some studies have assessed that, whereas most instruments yield the same render in recovering DNA from respiratory pathogens, they differ in reproducibility, recovery and linearity in recovering viral RNA [34].

Other parameters have to be considered for the purification of RV genomes. The extractor must be versatile so that it allows extraction from a wide range of respiratory samples and from variable sample volume. Indeed, lysis buffer must be provided independently by the manufacturer to inactivate respiratory specimens in biosafety cabinets if samples are fed in open containers into the instrument.

### 4.1.2. Real-Time Assays

Currently, real-time PCR assays are widely used in clinical laboratories. The principle of real-time PCR is that amplification of the nucleic acid target is combined with simultaneous detection of amplicons in a single step. There is no need for manipulation of amplified products, which minimises contaminations due to amplicon carry-over and avoids false positive reactions.

The monitoring of accumulating amplicons in real time is achieved by labelling of primers, oligonucleotide probes or amplicons with fluorophores. These emit fluorescent signals that accumulate during the exponential phase of the amplification step, which is recorded by the instrument. The signal is related to the amount of amplicon present during each cycle and will increase as the amount of specific amplicon increases. Various chemistries have been used for real-time PCR assays, i.e. Taqman® probes, FRET probes, intercalating dyes as Sybr Green I, molecular beacons, scorpions oligonucleotides, etc [35].

Commercial assays (Prodesse®, Gen-probe), based on real-time PCR for detection of several RV, have received the FDA clearance. Kits for Flu (types and subtypes), hMPV, ADV, RSV and PIV detection are available.

Real-time PCR using dual labelled probes such as Taqman® probes is widely used by virologists in their “in house” protocols, since it is easy to optimize, and despite the higher costs compared to conventional PCR protocols, the reduction in hands-on-time and the high efficiency compensates favourably the use of this approach. Common PCR parameters of 45 cycles of denaturation at 95°C for 30 s and annealing and extension at 60°C for 60 s, can be applied to any real-time assay [36]. This protocol is carried out in our laboratory for any RNA virus, including Flu, RSV, HRV, hMPV, hBoV and HCoV, by using the same instrument and the same protocol (personal communication).

### Table 1. Methods for Detection of Respiratory Viruses from Clinical Samples. Advantages and Disadvantages

<table>
<thead>
<tr>
<th>Laboratory Method</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>ASSAY (Turnaround Time)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Viral Culture</strong></td>
<td>Recovering of the viral strain for characterisation Open system for detecting multiple viral pathogens</td>
<td>Labour-intensive and time-consuming; lower sensitivity than NAAT; multiple cell lines required for the most prevalent RVs; new RVs do not grow or grow poorly in cell culture; expertise and appropriate infrastructure needed</td>
<td>Traditional viral culture (7-10 days) Shell-vial assay (24-48 h)</td>
</tr>
<tr>
<td><strong>Antigen Detection</strong></td>
<td>Rapid and easy to perform Point-of-care assay Especially useful for detection of RSV, Flu A and Flu B in paediatric patients</td>
<td>Variable sensitivity depending on the commercial assay, type of patient and/or sample, viral target; lower sensitivity for detection of RV in adults</td>
<td>Immunofluorescence (1-2 h) Immunochromatography (15-30 min)</td>
</tr>
<tr>
<td><strong>NAAT</strong></td>
<td>Greater sensitivity than viral culture and antigen detection methods Method of choice for detection of new RV</td>
<td>At least a part of the viral genome must be known in order to design adequate primers and probes</td>
<td>Conventional end-point PCR (1-4 h): commercial &amp; “in-house”; monoplex &amp; multiplex Real time PCR (45-90 min): commercial &amp; “in-house”; monoplex &amp; multiplex</td>
</tr>
</tbody>
</table>

#### Two-by-Two Comparison of Different NAAT Assays

<table>
<thead>
<tr>
<th>Assay</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Real time PCR vs conventional PCR</td>
<td>Reduced cross-contamination; less handling and turnaround time</td>
<td>Real-time instruments and fluorescent probes increase overall costs; the limited number of fluorophores with different wavelengths hampers highly multiplexing protocols</td>
</tr>
<tr>
<td>Commercial vs “in house” PCR</td>
<td>Validated and controlled reagents for in vitro diagnosis</td>
<td>Only available for the most prevalent viruses; more expensive</td>
</tr>
<tr>
<td>Multiplex vs multiple monoplex PCR</td>
<td>Cost saving; easier and faster processing of runs</td>
<td>More difficult to design and optimize; sensitivity and specificity is usually inversely proportional to the number of viral targets</td>
</tr>
<tr>
<td>RT + PCR vs one-step RT-PCR</td>
<td>Versatility; any viral target can be amplified from the cDNA product</td>
<td>Higher risk of cross contamination</td>
</tr>
</tbody>
</table>
The technical disadvantage of real-time PCR is the limited number of fluorophores that can be included in a single assay, thus, limiting the multiplexing capabilities. In the best case, 3 targets can be detected in a single tube. A published protocol is able to detect 12 RV by a multiplex real-time PCR, including 4 tubes per sample and 3 viral targets per tube [37].

One of the advantages of real-time PCR compared to other newly described protocols is the possibility of quantitation. The role of high viral loads in ARI have provided in many circumstances valuable information for the optimal management and treatment of infected individuals [38].

Pre-prepared, frozen real-time PCR reagents reduce turnaround times and increase reproducibility. Frozen aliquots of primers-probes mixes and, on the other hand, of generic PCR reagents (buffers, dNTPs, enzymes, etc) are a good alternative to reduce hands-on-time of the PCR preparation step [36].

Other emerging amplification procedures have been described for RV detection. Examples of these methods are: loop-mediated isothermal amplification, multiplex ligation-dependent probe amplification (MLPA) and padlock probes [39].

4.1.3. DNA Arrays and Other Systems for Detection of PCR Products

The combination of PCR and hybridisation using DNA arrays is the principle of recently developed commercial assays for RV detection. Depending on the commercial system, it uses generic or multiplex PCR and specific detection relies on the array.

Examples of this technology are: Luminex xTAG® Respiratory Virus Panel (RVP)(Luminex Molecular Diagnostics) and Resplex II (Qiagen).

Luminex xTAG® RVP was the first panel that received the FDA approval. The version RVP Fast has shortened the initial time of 8-10 h required to accomplish the whole procedure. The RVP Fast assay incorporates multiplex RT-PCR using primers with universal tags. The amplified product is hybridized to a bead array conjugated to specific probes. Detection is performed using a streptavidin-R-phycocerythrin conjugate. Each bead population detects a unique viral target by hybridisation to a specific anti-tag. A signal (median fluorescence intensity) is generated for each unique viral target by hybridisation to a specific anti-tag. A phycoerythrin conjugate. Each bead population detects a product is hybridized to a bead array conjugated to specific PCR using primers with universal tags. The amplified procedure. The RVP Fast assay incorporates multiplex RT-PCR followed a MLPA reaction, which comprises a probe hybridization step, a probe ligation step and a probe amplification step. Only two primers are used in the final PCR. Every target results in a PCR fragment of a specific length [45]. Seeplex® RV15 uses dual-priming oligonucleotides that allow multiplex PCR of high specificity and minimizes the possibility of unspecific DNA products. Detection of amplicons is accomplished by capillary electrophoresis with both assays [46]. Although these systems do not avoid the potential of contamination with amplicons, they are able to detect a wide range of RV at a lower cost. However, specific instruments for capillary electrophoresis are available only in high-throughput laboratories.

4.1.4. Platforms for Nucleic Acids Purification and Preparation of the PCR Master Mix

In recent years, several automated platforms for nucleic acids purification also enable the preparation of the PCR reaction mix in 96-well microplates ready for amplification-detection steps. VERSANT kPCR Sample Prep System (Siemens), QIAsymphony (Qiagen), m2000sp (Abbott) and MagnaPure 96 (Roche Diagnostics) are examples of these platforms. This improvement reduces even more the hands-on-time by abolishing the manual step associated with preparation of PCR mixes. Evaluations with some of these systems have demonstrated good and comparable performances [33,47].

4.2. Fully-Automated NAAT Platforms

The availability of fully-automated NAAT for detecting more or less number of RV is continuously increasing. The main advantages associated to the use of these tools are: ease of use, minimisation of hands-on-time, enclosed pouch to perform all reaction steps, minimum contamination risk, high-order multiplex testing, high sensitivity, specificity and reproducibility.
Most of these platforms have focussed on the automated detection of RSV and/or Flu, probably in an attempt to constitute in future a better alternative to antigen detection methods as point-of-care diagnosis.

Some available commercial fully-automated assays that detect Flu A, Flu B and/or RSV are: Xpert Flu (Cepheid) that detect Flu A and Flu B and differentiates A(H1N1)pdm09 [48] and Simplex Flu A/B & RSV kit (Focus Diagnostics) for detection and differentiation of Flu A, Flu B and RSV (http://www.focusdx.com/pdfs/pi/US/MOL2600.pdf). The advantage of the first one is that independent modules of the instrument allow continuous workflow, although each added module constitutes a new apparatus and increases costs. The later uses an 8-well cycler (3M® Integrated Cycler), allowing the simultaneous processing of 8 respiratory samples within the same apparatus.

Nanosphere Verigene® Respiratory Virus Nucleic Acids Test (Nanosphere) is a recently commercialised fully-automated assay similar to previous ones, with increased diagnostic capability since it detects and differentiates Flu A, seasonal H1N1, A(H1N1)pdm09 and H3N2 subtypes, Flu B and RSV A and B viruses. Indeed, an added innovation of this system is the detection of the H275Y mutation within the neuraminidase of A(H1N1)pdm09 that confers resistance to oseltamivir. Studies evaluating this commercial test have shown optimal analytical and clinical performances [49,50].

The FilmArray® Respiratory Panel (Idaho Technology Inc.) is commercialised in Europe to fully automate the detection and identification of 21 respiratory pathogens from a single sample in about one hour, including 18 viruses and 3 bacteria. An unprocessed clinical sample is subjected to nucleic acid purification, RT, a high-order nested multiplex PCR and amplicon melting curve analysis in a closed disposable pouch [51]. The larger spectrum of this panel that allows the additional detection of other respiratory pathogens may be an advantage for diagnosing ARI in some population groups [52]. However, detection of more pathogens leads to a more expensive system. This assay is probably cost-effective for rapid detection of respiratory agents from certain clinically severe patients. Table 2 displays a list of commercial fully-automated assays for detection and identification of RV.

A recent revision of molecular assays for RV carried out by Ginocchio records other available automated and semi-automated commercial tests [43].

### Table 2. Fully-Automated Assays for Detection and Identification of Respiratory Viruses

<table>
<thead>
<tr>
<th>Assay</th>
<th>Manufacturer</th>
<th>Method</th>
<th>Virus Detected</th>
<th>Hands–On/ Turnaround Times (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MariPOC® respi test</td>
<td>ArcDia</td>
<td>Antigen detection</td>
<td>Flu A, Flu B, RSV, hMPV, PIV1-3, ADV</td>
<td>1 / 20-120</td>
</tr>
<tr>
<td>Xpert® Flu</td>
<td>Cepheid</td>
<td>Nucleic acid extraction</td>
<td>Flu A, Flu B, 2009 H1N1</td>
<td>1 / 60</td>
</tr>
<tr>
<td>Simplex® Flu A/B &amp; RSV Direct kit</td>
<td>Focus Diagnostics</td>
<td>Nucleic acid extraction</td>
<td>Flu A, Flu B, RSV</td>
<td>1 / 60</td>
</tr>
<tr>
<td>Verigene® Respiratory virus XP test</td>
<td>Nanosphere</td>
<td>Nucleic acid extraction</td>
<td>FluA, FluB, RSV A, RSV B, 2009 H1N1, Flu A H1, Flu A H3, oseltamivir-resistant 2009 H1N1 (H275Y mutation), PIV 1-4, ADV, hMPV, HRV, HEV</td>
<td>5 / 90</td>
</tr>
<tr>
<td>Filmarray® Respiratory Panel</td>
<td>Idaho technology Inc.</td>
<td>Nucleic acid extraction</td>
<td>ADV, hBoV, HCoV (HKU1, NL63, 229E, OC43), Flu A H1, 2009 H1N1, Flu A H3, Flu B, hMPV, PIV1-4, ADV, RSV, HRV/HEV</td>
<td>2 / 60</td>
</tr>
</tbody>
</table>

5. COST-EFFECTIVENESS OF LABORATORY PROCEDURES: INTEGRATION OF AUTOMATED NAAT TO RATIONAL DIAGNOSTIC ALGORITHMS

The optimal use of the diagnostic tools for RV detection depends on the correct choice of the method in combination with other clinic-epidemiologic criteria. The origin of the sample, patient’s age, predisposing factors for severe disease and the prevalence of a specific RV in a certain area and within a study period must be considered for an appropriate diagnostic algorithm. The overwhelming demand for RV detection in clinical samples has been evident and continuous since the advent of the 2009 influenza pandemic, when clinicians and health institutions have realised that, in many circumstances, an aetiological diagnosis of ARI is desirable. Thus, clinical laboratories must take into account all these points and the availability of infrastructure and resources, to organise their workflow and portfolio related to the diagnosis of viral ARI.

A rational algorithm for viral diagnosis of ARI could be
divided in four levels of increasing use of diagnostic tools depending on the type of patient [53].

Overall, viral diagnosis in outpatients with mild ARI is not probably cost-effective, and should not be routinely performed, except for studies from surveillance programs or of epidemiological interest.

Antigen detection methods for RSV and/or Flu may be enough for patients attended at emergency units that do not fulfill severity clinical criteria for hospitalisation. Some individuals should constitute an exception of this protocol as pregnant women, patients with severe underlying conditions and other special situations (immunosuppression, obesity, residents in chronic-care facilities, etc) for which a more extended aetiological study and/or the use of NAAT for the most prevalent RV is encouraging [54-57].

Accurate and timely diagnosis of viral ARI has additional benefits as preventing unnecessary use of antibiotics, thus, decreasing the overall costs of treatments and minimizing the selection of resistant organisms. Moreover, specific antiviral therapy is recommended for treating certain risk groups with Flu and RSV infection. Neuroaminidase inhibitors are recommended in patients with suspected or confirmed Flu who require hospitalisation and/or in high-risk groups. Early treatment with neuroaminidase inhibitors has demonstrated to decrease the risk of complications, to reduce the duration of the disease and to improve outcome [54]. Ribavirin is approved for the treatment of hospitalised infants, young children and immunocompromised adults with severe lower respiratory tract infections due to RSV, despite the limited evidence of benefit [58,59]. New automated assays may improve point-of-care diagnosis of RV since they include a wider range of pathogens within the same kit. In future, automated NAAT may be the alternative of antigen detection methods if costs become comparable and a reduction of turnaround time is achieved.

Samples from patients requiring hospitalisation with negative RSV or Flu antigen results should be subjected to extended virological study that includes viral culture assays and/or NAAT targeted at more or less RV depending on the severity of the disease and the need for a rapid result. With this premise, NAAT for detection of RSV and Flu (at the type and subtype level) in patients with less severe ARI, that do not require admission at intensive care units (ICU), should be carried out first of all. Otherwise, other molecular multiplexed methods of low complexity could be performed in certain patients that included hMPV, HRV and PIV detection, depending on patient’s age and/or the existence of underlying diseases. More complex NAAT that detect other less frequent RV should be reserved to certain situations such as more severe cases (ICU patients) in which previous NAAT for the most prevalent viruses have yielded negative results [60-62]. At this level, automated and semi-automated NAAT may provide higher throughput than in-house protocols.

In the future, the feasibility of introducing fully-automated NAAT in a wider range of health institutions can constitute an improvement in viral diagnosis of ARI. Basic and first-level hospitals may provide, as well, with an efficient point-of-care assay to ensure the appropriate management of their in-patients.

6. CONCLUSIONS

Automation in the field of clinical virology for diagnosis of ARI relies on molecular assays. New approaches have yielded point-of-care methods, optimal for a rapid detection of several RV.

Cost-effective laboratory methods must take into account the benefits of aetiological diagnosis depending on technical characteristics of the assays, on the availability of the appropriate resources and on clinic-epidemiologic criteria. Automated NAAT are expensive but probably the most cost-effective tools for severely ill hospitalised patients. Many automated commercial assays require their own instrument; thus, each laboratory should make its forecast for future to provide with the most feasible automation strategies. Since no expertise is required to carry out automated molecular assays, extended aetiological diagnosis of ARI can be achieved if first-level hospitals implement these techniques in their portfolio.

CONFLICT OF INTEREST

The authors declare that they have no competing interests.

ACKNOWLEDGEMENTS

Declared none.

REFERENCES

The Open Virology Journal, 2012, Volume 6

Pérez-Ruiz et al.

158


32. Clinical Microbiology, 4-7 September, Madrid 2012. p. 105.
adults is similar to that of 2009 H1N1 influenza. J Clin Microbiol 2012; 50: 1061-3.


