Syndromic panels or the panels’ syndrome? A perspective through the lens of respiratory tract infections - Author’s reply

Marie-Céline Zanella1,2,3, Pascal Meylan4, Laurent Kaiser1,2

1 Laboratory of Virology, Division of Laboratory Medicine and Division of Infectious Diseases, Geneva University Hospitals, Geneva, Switzerland
2 University of Geneva Medical School, Geneva, Switzerland
3 Laboratory of Bacteriology, Division of Laboratory Medicine and Division of Infectious Diseases, Geneva University Hospitals, Geneva, Switzerland
4 University of Lausanne Faculty of Biology and Medicine, Lausanne, Switzerland

Submitted to: Clinical Microbiology & Infection (Letter to the Editor - Author’s reply)

Word count: 799

Corresponding author:

Marie-Céline Zanella
Laboratory of Virology, Division of Laboratory Medicine and Division of Infectious Diseases
Geneva University Hospitals
4 Rue Gabrielle Perret-Gentil
1211 Geneva 14, Switzerland
Tel: +41 79 553 5523; fax: +41 22 372 4097
E-mail: marie-celine.zanella@hcuge.ch
To the Editor,

We thank Brendish et al [1] for their interest in our commentary [2] and they rightly point out that it was not a systematic review. Our main intention was to provide food for thought and discussion regarding the use of panel assays in the light of some relevant publications. In particular, we aimed at discussing the limitations of their analytical aspects and clinical validation. We respectfully disagree with the statement that “The authors suggest that the increasing use of rapid, automated, syndromic molecular panels for respiratory viruses (RVs) should be abandoned in favor of more limited PCR testing for RVs”. As microbiologists and clinicians, we rather suggest that we have the responsibility to promote diagnostic stewardship in order to integrate these new technologies in clinical management, while considering their strengths and limitations. We also highlight the value of a multiple-step approach of testing that does not necessarily preclude their use.

We agree with the implementation of rapid diagnostic assays for RVs given that several studies have shown a clinical impact with a more appropriate use of oseltamivir, reduced length of stay and fewer chest X-rays, as summarized in the meta-analysis of Vos et al [3], which included the ResPOC study [4]. However, it should be highlighted that there is not enough evidence so far for a reduction in antibiotic prescription and duration or hospital admission due to rapid molecular tests compared to conventional molecular ones [3]. The ResPOC study concluded that a point-of-care (POC) assay was associated with a more appropriate use of antiviral treatment, brief courses of antibiotics and shorter length of stay, but no difference was observed regarding antibiotics’ mean duration [4]. In addition, only 45% of patients in the control group were tested and the conclusions may have some limitations.

We agree that one of the important aspects of testing is that the results be obtained in a meaningful timeframe for clinicians. In particular, short turnaround times have an impact on
clinical management when testing for influenza virus and RSV. Among paediatric patients, the use of rapid tests demonstrated a decrease in emergency department length of stay, a reduction of further diagnostic tests, and an increase in the appropriate use of antibiotics and antivirals [5]. Similarly, a more appropriate use of oseltamivir, shorter time to isolation and a reduction in length of stay were also shown among adults [6]. Nevertheless, considering the methodological limitations, we believe that the conclusions driven by the post hoc analysis of the ResPOC study regarding the impact of turnaround time on antibiotic use and length of stay should be interpreted with caution [7]. The impact of short turnaround times on patient clinical management has been well summarized in the commentary of Kuypers [8]. We have already implemented a POC test for influenza virus and RSV at our institution. However, we continue to perform conventional molecular testing for RV with in-house panels that encompass diverse viral targets according to clinical presentation and epidemiology.

Regarding influenza virus infection specifically, we remind clinicians that antiviral treatment should be started as soon as possible for patients with a suspected or documented influenza infection and treatment and isolation measures should not be deferred while waiting for any assay result [9].

Clinicians should be also aware that clinical validation studies have assessed the performance of some respiratory panels with suboptimal approaches and reported cumulative performance results or positive and negative percent agreement as surrogates of sensitivity and specificity [10]. Notably, certain panels encompass bacterial targets that have been validated with fewer than 10 positive samples and performance has sometimes been reported to be lower than those of viral targets [10]. Thus, clinicians should be particularly attentive when interpreting cumulative performance results or percent agreement values, which may represent an important and underappreciated limitation.
Overall, we consider that respiratory panels can be integrated in clinical management, but we encourage the use of smaller, more targeted panels for pathogens that actually impact on clinical management as part of a multiple-step approach of testing. Indeed, systematic first-line testing for a broad range of pathogens that are neither clinically nor epidemiologically suspected and with no demonstrated impact on clinical endpoints and cost-effectiveness is contrary to appropriate diagnostic stewardship and evidence-based decisions for diagnostic and therapeutic strategies. Importantly, new technologies should serve the clinician’s needs and not be driven by marketing strategies. Furthermore, in the era of the SARS-CoV-2 epidemic, targeted testing is needed more than ever. Finally, we agree with Kuypers [8] that the decision to implement a rapid molecular assay for respiratory pathogens requires consideration of clinical and economic factors unique to each healthcare facility. Well-designed multicentre studies are needed using methods and standards to allow others to use the results in evidence-based laboratory practice guidelines, as well as to increase evidence regarding their impact in clinical management and cost-effectiveness.

**Transparency declaration:** The authors have no conflicts of interest to declare. No funding was received for this letter.

**Author’s contributions:** MCZ and LK contributed to the conceptualization of the letter. MCZ contributed to the writing of the original draft, reviewing and editing of the letter. LK contributed to the supervision, reviewing and editing of the letter. PM contributed to the reviewing and editing.
References


