Co-administration of vaccines: a focus on tetravalent Measles-Mumps-Rubella-Varicella (MMRV) and meningococcal C conjugate vaccines

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ABSTRACT
Simultaneous administration of different vaccines is a strategy to increase the possibility to receive vaccines at appropriate age, safely and effectively, reducing the number of sessions and allowing a more acceptable integration of new vaccines into National Immunization Programs (NIPs). Co-administration can be performed when there are specific indications in the Summary of Product Characteristics (SmPC) of the vaccines; but, in absence of these indications, the practice is possible if there are no specific contraindications nor scientific evidence to discourage simultaneous administration.

The aim of this work is to review the safety and efficacy of co-administration of the tetravalent measles, mumps, rubella, and varicella (MMRV) and the meningococcal C (Men C) conjugate vaccines after 12 months of age.

Several studies demonstrated that MMRV and Men C conjugate vaccines can be administered concomitantly without a negative impact on the safety and immunogenicity of either vaccines, inducing highly immunogenic responses.

Introduction
The co-administration of vaccines is proposed to reduce the number of vaccination sessions, increase compliance, and ensure optimal coverage. In particular, the tetravalent measles, mumps, rubella, and varicella (MMRV) and the monovalent or tetravalent meningococcal C (Men C or Men ACWY) conjugate vaccines are administered worldwide between 13 and 15 months of life, a period during which children are immunized against different infectious diseases. Most of the scientific studies focus the analysis mainly on co-administration between MMRV and tetravalent meningococcal conjugate vaccines.1,2 The safety and efficacy of co-administration and to provide a focus on MMRV and Men C conjugate vaccine co-administration in this age group.

Background
The immunization programs of each country greatly vary depending on the local context (epidemiological, socio-economic, cultural) and, according to the World Health Organization (WHO) recommendations, on the most appropriate age for the administration of different vaccines.3

Vaccination strategies adopted in childhood can require a high number of sessions4 and the fulfillment of the entire vaccination schedule can be challenging for newborns, parents and health professionals.5

This phenomenon is even more evident with the development and introduction of new vaccines. This makes the vaccination schedule more complex due to the further increase in the number of injections and immunization sessions. The consequences of a complex schedule may be a low compliance to vaccination and a consequent reduction of vaccination coverage rates.1

Simultaneous administration of different antigens can be a valid practice to counteract the above-mentioned issues; this opportunity finds its rationale in the combination of different antigens as well as in the co-administration of several vaccines in the same session.

These practices increase the possibility to receive age-appropriate vaccines in a timely manner, reducing the number of sessions and allowing a more acceptable integration of new vaccines into schedules4,6,7

A combined vaccine can be defined as a vaccine consisting of two or more antigens, ready to use or to be mixed immediately before administration. The goal is to protect against multiple infectious diseases or multiple strains of infectious agents that cause the same disease.8,9

Co-administration of vaccines is defined as the administration of more than one vaccine on the same day. These administrations should be performed in different anatomic sites.9

The vaccine combination and co-administration provide an important means of vaccination strategies’ optimization in different countries.
The administration of combined vaccines and co-administration also brings with them some critical issues sometimes linked to a difficult acceptance by citizens. There are concerns about the possibility that too many vaccines overload the immune system, that the combined vaccines may be less effective and can cause a greater number of adverse reactions compared to the vaccines administered separately.\textsuperscript{10,11}

Combined products reduce the number of injections and pain compared to repeated injections; on the other hand, they also reduce cumulative exposure to preservatives and stabilizers that are erroneously linked by hesitant parents to the onset of adverse reactions.\textsuperscript{12,13}

Furthermore, co-administration reduces the number of visits and, therefore, decreases stress and discomfort for children and parents.\textsuperscript{5}

Combined vaccines are thoroughly tested before they are approved for marketing. The United States (US) Food and Drug Administration (FDA) and the European Medicines Agency (EMA) ensure that the new products are safe and effective. There is no evidence that combined vaccines increase the burden of the immune system, which instead is able to respond simultaneously to millions of antigens. In general, the combination of different antigens may or may not bring to an additive effect on reactogenicity but always within the limits required for the product to be approved.\textsuperscript{14}

According to recommendations of the Advisory Committee on Immunization Practices (ACIP), there are numerous vaccines that can be administered simultaneously in an effective and safe way.\textsuperscript{9} Co-administration of different vaccines results to be immunogenic and well-tolerated as described in several papers.\textsuperscript{15-17} To cite some examples, Nakashima et al.\textsuperscript{18} and Thompson et al.\textsuperscript{19} concluded that co-administration of the quadrivalent influenza vaccine and the 23-valent pneumococcal polysaccharide vaccine (PPV23) and with the 13-valent pneumococcal polysaccharide conjugate vaccine (PCV13) respectively, is a safe and effective practice and does not lead to an increase in adverse reactions.\textsuperscript{18,19} Several authors evaluated the safety and immunogenicity of the quadrivalent meningococcal conjugate vaccine (Men ACWY) co-administered with common childhood vaccines, confirming good tolerability, elevated immunogenicity, and safety.\textsuperscript{20-24} Finally, with regard to the concomitant administration of meningococcal C conjugated vaccines (as tetravalent or monovalent vaccines) and MMRV, which will be further investigated in the course of this review, in recent studies performed by Durando et al.\textsuperscript{5} and Klein et al.\textsuperscript{2} this co-administration appears to be well tolerated, without safety issues and with a good immune response for both vaccines.\textsuperscript{2,5}

Public health impact of co-administration

Combined and co-administered vaccines optimize vaccination coverage thanks to the simplification of vaccination programs.\textsuperscript{11,25,26}

These elements guarantee a significant impact in terms of public health, specifically with the increase in vaccination coverage and the more efficient management of services.\textsuperscript{9}

Considering the higher spending on the purchase of more expensive vaccines that could initially discourage their use, the combination of vaccination saves economic resources thanks to the more profitable management of vaccination services, which will benefit from an easier management of appointments and potentially reduce registration errors.\textsuperscript{27}

There are no studies aimed at evaluating the impact assessment of co-administration in terms of economic savings, but some studies have been developed for combined vaccines, which mainly share the same principles of co-administration. In 1999, Fendrick et al.\textsuperscript{28} conducted a study to investigate the potential impact in terms of public health offered by the combination of a vaccine against Haemophilus Influenzae type b (Hib) and Hepatitis B vaccine (HBV) compared to the previous strategy with single vaccines. The obtained results were very encouraging especially regarding the impact on HBV compared to single vaccinations, estimating a reduction of cases of acute hepatitis, chronic hepatitis, and deaths of 53%, and a saving of resources of 4 million dollars. The results showed an increase in costs related to the immunization with this combined vaccine equal to 11 million dollars, but almost exclusively due to the increased coverage achieved.\textsuperscript{28}

In 2014, Zhou et al.\textsuperscript{29} published a study to evaluate the direct and indirect cost savings offered by vaccinations through analysis of the data on the 2009 USA vaccine schedule and comparing it with a similar analysis dating back to 2001. Considering the increased use of combined vaccines in 2009, the researchers estimated an increase in the costs of immunization equal to about 4.7 billion US dollars. This estimation was due to higher costs related to combined vaccines in respect to single vaccines, and to the need to carry out more vaccination sessions in order to adhere to a more articulated schedule than the 2001 schedule.\textsuperscript{29}

However, the saving of direct and indirect costs related to the diseases and related to prevented complications has been estimated equal to almost 30 billion US dollars, mainly due to the increased vaccination coverage guaranteed by combination and co-administration.\textsuperscript{30}

Other advantages of combined vaccines are the simplification of the vaccination sessions and the enhanced compliance with the correct timing of the vaccination courses. Indeed, in 2017, Macartney et al.\textsuperscript{31} studied this phenomenon in relation to the introduction of the MMRV combined vaccine in the national schedule compared to the previous, already combined, measles, mumps, and rubella (MMR) vaccine. Taking into account the second dose of vaccine, MMRV allowed guaranteeing the correct timing in 13.5% more cases than the MMR vaccine. In addition, MMRV facilitated an increase by 4% coverage against varicella (first dose) in comparison to the monovalent vaccination.\textsuperscript{31}

Summary of product characteristics (SmPC): co-administration with or without specific indications

The SmPC is the most important regulatory document on a medicinal product in the European Union because it is part of the marketing authorization. It represents the basis of information for health-care professionals on how to use the medicinal product safely and effectively and it is frequently updated when new efficacy or safety data are available.

While the SmPC is a document intended primarily for health professionals (doctors, pharmacists, nurses), the Package Leaflet
(PL) is a document intended for the patient/user and contains all the useful information for a safer and more correct use of the drug expressed in a clear and easy language.

The preparation of the SmPC and the PL (Figure 1) is performed by referring to the document Corporate Core Data Sheet (CCDS), prepared by the marketing authorization holder and containing information concerning the safety in the narrow sense (Corporate Core Safety Information, CCSI), information on the indications, posology, pharmacology and all other information concerning the product.32

SmPC is a document with legal value and the contents of the PL and SmPC are established by law.33

In the SmPC, information is presented following a predefined structure. One of the most relevant parts of SmPC is Section 4.1, that defines the pathological conditions for which the drug is intended. Moreover, it defines the use of the drug (therapeutic, preventive, diagnostic) and the limits of age for the drug use, according to the results of clinical trials.

From a medical-legal point of view, therapeutic indications established by SmPC should not debar the use of the drug/vaccine in other subjects. For example, hexavalent vaccines are administered “since 6 weeks of life”; therefore, they have no upper limit of use and this indication does not preclude its use in older age groups.32

Moreover, the indication of use in the SmPC should not be confused with the term “recommended” which is applied to the NIPs. In fact, the term “recommended” is used on the basis of epidemiological evidence, suggesting a lower risk of disease in other age groups, but it does not relate to indication for use of the vaccine. For example, the monovalent Hib vaccine is not recommended above 5 years of age in the current NIPs; however, it is widely used also in adults with particular risk factors (i.e. asplenic subjects), for which it can represent a real life-saving intervention. There is no evidence that the administration of one, or more doses, of the Hib vaccine over 5 years entails risks of increased frequency and severity of adverse reactions.34

Section 4.3 of the SmPC defines “the situations where the medicine must not be used for safety reasons”32 and the patient populations who must not take the medicine.

The SmPC of a vaccine provides specific indications on co-administration, but for a number of reasons, when the SmPC is produced, it may not define all possible vaccine co-administration. Firstly, additional information, such as trials and post-marketing data on vaccine effectiveness, as well as safety data generated after large-scale use of the vaccine may be available after vaccine authorization. Secondly, decisions regarding vaccine use should also take into consideration a wide range of parameters including the age-specific pre-vaccine disease burden, public health needs, other health interventions, costs and cost-effectiveness studies, programmatic issues (such as existing schedules, impact on equity and policies) that are specific to the country or region, independently from the strict indications included in the SmPC. Thus, public health recommendations may differ across countries and WHO regions.35

In addition, co-administration can be useful when there are no specific contraindications to co-administration and even if there are no specific indications in the SmPC. Moreover, sometimes there are scientific pieces of evidence to support the co-administration, which have not yet been included in the SmPC. Actually, international recommendations indicate that vaccines can be co-administered even in the absence of specific studies, unless there are specific pieces of evidence to avoid co-administration: the same situation is applied in the co-administration of all drugs.36

For example, Meningococcus B and Rotavirus vaccines are co-administered in the absence of explicit specification in the SmPC but this practice is supported in the literature.37
Co-administration of vaccines does not lead to an overload of the immune system; instead, it allows to obtain an antibody response for different diseases in less time.

As a matter of fact, since August 2017 the Joint Committee on Vaccination and Immunization in United Kingdom (UK) recommends during the first vaccination session (starting from the eighth week of life) the co-administration of Meningococcal B vaccine with Hexavalent, PCV-13 and Rotavirus vaccines.38,39

Also in Italy, the Ministry of Health recommends the co-administration of the Hexavalent, PCV-13 and Rotavirus vaccines to all newborns starting from the first vaccination session.40

**Off-label use of vaccines**

Off-label use of vaccines or medicines has no standard definition. EMA defines off-label use as the situation where a medicine is intentionally used for a medical purpose not in accordance with the authorized product information.41 Off-label use (called unlabeled indication) is defined by the FDA under the perspective of the health-care provider only (i.e. when a marketed drug is prescribed to treat a patient for an unlabeled indication).42 For the American Academy of Pediatrics Committee on Drugs, off-label use of an approved drug refers to a use that is not included in the approved label.43 On the Medical Dictionary for Regulatory Activities (MEDRA) “off-label use” is defined as a practice of prescribing pharmaceuticals outside the scope of the drug’s approved label, most often concerning the drug’s indication.44 For the Canadian Agency for Drugs and Technologies in Health, “off-label” or “unlabelled” drug use occurs when a drug is used in a treatment regimen or patient population that is not included in the Notice of Compliance (NOC), and a drug is used for an indication other than those specifically included in the NOC.45 Many off-label drug uses are effective, well documented in the peer-reviewed literature, and widely used.35,46–48 In the chapter of the Practical Law of Company’s multi-jurisdictional guide to life sciences, L’Ecluse et al.49 define off-label use the prescription or administration of an authorized medicinal product outside any of the terms of the marketing authorization, as reflected in the SmPC. This might include use for a different indication, at a different dosage (or dosage frequency) or in a different patient group (for example, children or pregnant women).49 Therefore, off-label use can be defined as a prescription for a different indication, posology, age or population group from that included on the label.50 The Italian legislation that regulates the off-label use of medicinal products50 indicates that the physician, when prescribing a drug, must follow the therapeutic indications and the ways of administration foreseen by the marketing authorization of the drug. However, the law allows using the drug outside the authorized indications if there are pieces of evidence documented in the literature and in the absence of better therapeutic alternatives. The off-label use of drugs exposes the patient to potential risks, considering that their efficacy and safety have been evaluated in different populations. From a medical/legal perspective, in 2017 the law n.24 “Provisions on professional liability of health personnel” was issued in Italy. This law shows that there is no medical liability when following established guidelines, such as the National Immunization Plan or the recommendations of scientific societies.51

**Immunogenicity and safety of co-administration of MMRV and meningococcal C conjugate vaccines**

MMRV vaccines available in Europe are live attenuated tetravalent vaccines for the prevention of measles, mumps, rubella, and varicella (chickenpox) and they are licensed since 2005.52,53

In the scientific literature, the efficacy of individual components of the MMRV vaccine has been established previously.10,54–58 MMRV vaccine had similar immunogenicity and overall safety profiles as MMR vaccine administered with or without varicella vaccine.59,60

The vaccines available in Europe for the immunization of children toward Meningococcus C are conjugate vaccines, as monovalent and tetravalent ACYW,61–67

The safety and the immunogenicity of meningococcal conjugate vaccines had been clearly evaluated in the literature.22,23,68–72

In most European countries, both the meningococcal conjugate and MMRV (or MMR+V) vaccines are recommended at around 13–15 months of age (Table 1).38

Some international studies demonstrated the advantages of vaccinations administered in the same visit, such as an increase in vaccination coverage, an increase in compliance with the entire vaccination course and the respect of the correct timing.9,11,25,26

Kurosky et al.73 show how the combination of vaccines has considerably increased adherence to the complete vaccination course compared to what happens with the administration of monovalent vaccines. For example, as regards MMR vaccination, compliance with the timing of the vaccination course in those who received the combined vaccine was significantly higher (81.4%) compared to those who received the single vaccines (63.4%).73

The considerations that emerge from these studies can also be made for the co-administration of MMRV and meningococcal vaccines, although at present there are no specific studies on the subject in the literature. To our knowledge, there are few studies on the co-administration of monovalent Men C conjugate vaccines and MMRV vaccines, while in the international scientific literature, some studies have evaluated co-administration of Men ACYW conjugate vaccine with MMRV vaccine.2,23

A study by Vesikari et al.1 showed that the quadrivalent Men ACYW conjugate vaccine can be co-administered with the MMRV vaccine between 12 and 23 months of age without affecting the immunogenicity or safety profiles of either vaccine. The occurrence of fever in the groups of children vaccinated with Men ACYW conjugate and MMRV versus MMR alone was within the same range.1

Actually, the co-administration of meningococcal conjugate vaccines does not affect the fever pattern, which is probably induced by the MMRV vaccine. Indeed, the prevalence of fever in both groups peaked between 4 and 10 days after
vaccination with MMRV, in line with the known timing of fever in association with MMRV vaccines.47

Similarly, Klein et al.17 evaluated the safety in subjects who received Men ACWY conjugate vaccine and MMRV vaccine compared with those who received the MMRV vaccine alone. The study enrolled three groups of healthy children aged 7–9 months and 12 months. The first two groups included children aged from 7 to 9 months who received Men ACWY-CRM at 7–9 months either alone (group 2) or with MMRV at 12 months (group 1). A third group (group 3) was simultaneously enrolled at 12 months, receiving only MMRV.

Local reactogenicity was similar between the different groups. As to systemic reactions, children vaccinated with conjugate Men ACWY and MMRV compared with subjects who received Men ACWY conjugate vaccine alone experienced slightly higher rates of severe reactions during the 7 days of the period of follow-up, including fever; however, they had similar or lower rates of severe systemic reactions than those who received MMRV vaccine alone.

Furthermore, it was noted that within 7 days after vaccination, in the subjects who received conjugate Men ACWY and MMRV vaccines concomitantly, there were fewer febrile episodes compared to the group that received MMRV alone.2

Instead, Durando et al.5 reported that the occurrence of fever and rash in the Men C + MMRV and MMRV groups was within the same range.5

Also, Vesikari et al.1 reported no significant differences were found in the presentation of adverse events between the Men ACWY + MMRV and MMRV groups alone.2

These studies demonstrated that Men ACWY conjugate vaccine can be administered in co-administration with MMRV vaccine at about 12 months of age, without a negative impact on the safety of both vaccines, and induced highly immunogenic responses against all four serogroups.2,23

In the same way, in the study by Scott et al.54 co-administration of MMRV vaccine with a tetravalent Men ACWY conjugate vaccine in infants aged 12 months was associated with robust immune responses to all strains of both vaccines.54

In an open-label multicenter trial, 1,014 children vaccinated with MMRV and conjugated Men ACWY vaccines and 616 children vaccinated with MMRV alone were analyzed. This analysis showed that the simultaneous administration of Men ACWY-CRM with MMRV at 12 months is well tolerated and without safety problems. The antibody responses to all vaccine components met the non-inferiority criteria following the administration of the meningococcal conjugate vaccine administered alone or co-administered with MMRV.2

To date, the study of Durando et al.5 conducted in Italy is the only one to prove the immunogenicity and safety of co-administration of a MMRV vaccine with a monovalent Men C conjugate vaccine.

The vaccines used in the study were a single dose (0.5 ml) of MMRV vaccine, Priorix-TetraTM (GSK, Belgium) and a single dose (0.5 ml) of Men C CRM-197 conjugated vaccine, Meningitec (Nuron Biotech). In the study, 716 children between 13 and 15 months of age were enrolled. The bactericidal activity (rSBA- Men C: anti-menignococcal serogroup activity, cutoff: P ≥ 1.8) in groups of toddlers vaccinated with MMRV and Men C conjugate vaccines and Men C conjugate vaccine alone was 98.3% (95% CI 96–99.4 CI) and 99.3% (95% CI 96.2–100 CI), respectively. Antibody titers against single antigens of the MMRV vaccine, measured using enzyme-linked immunosorbent assays (ELISA), had the same protective values when co-administered with Men C conjugate vaccine or when given separately. The immune responses elicited by co-administered MMRV and Men C conjugate vaccines were high in all groups and non-inferior to those elicited by either MMRV or Men C conjugate vaccines alone. Regarding the safety profile, the percentages of toddlers who developed fever during the post-vaccination period (i.e. 43 days) were similar and likely due to the MMRV vaccine (64.9% in the group of subjects vaccinated with MMRV and conjugate Men C, 64.4% in subjects vaccinated with MMRV alone, and 37.4% in the group vaccinated with conjugate Men C alone). Indeed, the characteristic peak of fever at 5–12 days post MMRV vaccination was observed in both the MMRV/conjugate Men C and MMRV groups.5 In addition to the positive data reported in the literature on co-administration of MMRV and quadrivalent meningococcus conjugate vaccines, also the co-administration of MMRV and Men C conjugate vaccines is possible and effective and does not affect the safety and immunogenicity of the respective vaccines.

Table 1. Recommended vaccinations in Europe for meningococcal C, Measles-Mumps-Rubella and Varicella vaccines (Adapted from European Center for Disease Prevention and Control (ECDC)-Vaccine Scheduler: https://vaccine-schedule.ecdc.europa.eu/).
Guidelines

The international guidelines on good practices to be implemented in the vaccination area give a consensus on the co-administration of vaccines. Regarding the co-administration of MMRV and meningococcal conjugate vaccines, there are specific indications in the SmPC for one of the two MMRV vaccines. The most accepted opinion regarding co-administration is summarized in the Pink Book where the following indications of the ACIP are reported: "simultaneously routine administration of all age-appropriate doses of vaccines is recommended for children for whom no specific contraindications exist at the time of the visit".

The same ACIP guidelines, in the “Vaccine Administration” section, recommend performing vaccinations in the same session using separate anatomical sites, documenting for each patient in which place a specific vaccine has been injected. As a good clinical practice, it is also suggested to create an anatomical reference map for professionals in order to know where to administer a specific vaccine. If it is necessary to simultaneously administer more than two vaccines in the same site, it is a good practice to distance the injections by at least 1 inch (about 2.5 cm) to be able to identify and distinguish any local reactions.

The Immunization Book of the New Zealand Ministry of Health, revised in March 2018, encourages the simultaneous administration of vaccinations in order to reduce immunization sessions. In particular, this practice is suggested in the 15th month of life when the New Zealand vaccination schedule foresees the administration of four vaccinations (MMR, Varicella, PCV, Hib) that, according to the same guidelines, should be performed simultaneously in four distinct anatomical sites.

In addition, the guidelines of the Public Health Agency of Canada, updated in May 2017, suggest to implement co-administration, every time multiple vaccinations are provided for the schedule. This practice is considered always possible using different inactivated vaccines, as well as administering inactivated vaccines with live attenuated viral vaccines, as in the case of meningococcal conjugated vaccines and MMRV.

In the Green Book, the guidelines of the National Institute for Health and Care Excellence (NICE) of the UK are reported, updated in August 2017, which recommend the simultaneous administration between inactivated vaccines or inactivated vaccines with live attenuated ones as always possible. In Italy, the 5th edition of the "Guide to contraindications to vaccinations" published in March 2018 by the Ministry of Health and the High Health Council states that vaccinations can be safely administered in the same immunization session, maintaining a high immunogenicity against each antigen and helping to increase the successful completion of the vaccination courses. Co-administration of MMRV and meningococcal conjugated vaccines is therefore always possible, except for specific contraindications related to individual vaccinations. The only co-administration linked to MMRV, which should be avoided is the one with yellow fever vaccine because a lower immune response to mumps, rubella and yellow fever has been documented.

In Italy, the last edition of the "Vaccination Calendar for Life" (2019 edition), endorsed by four Scientific Societies (Italian Society of Hygiene, Preventive Medicine and Public Health – SItI, Italian Federation of Family Pediatricians – FIMP, Italian Federation of General Practitioners – FIMMG, and Italian Society of Pediatrics – SIP), supports co-administration of MMRV with monovalent meningococcal C conjugate vaccine. In the notes on the co-administrations of MMRV, it is suggested: "Administration of MMR or MMRV is possible in association with conjugate meningococcus C, meningococcus quadrivalent (ACWY) or meningococcus B vaccines. The same recommendation is reported in the Italian NIP 2017–2019, in order to provide a rapid protection against many antigens simultaneously.

Conclusions

Co-administration, and the use of combined vaccines, can be considered as one of the best approaches to reach high vaccination coverage rates, and to guarantee the objectives of the immunization program, ensuring immunogenicity and safety. Furthermore, co-administration allows to obtain a greater compliance, a reduction of costs, a simplification of operational procedures, and the possibility of adopting new vaccines. As a matter of fact, the addition of varicella to MMR vaccines has had a documented positive carry-over effect on varicella coverage. Based on clinical trials, co-administration of MMRV vaccine with other routine childhood vaccinations does not impact significantly on either the immunogenicity of co-administered vaccines or the safety profile of MMRV. In particular, the co-administration of MMRV and meningococcal C conjugate vaccines is not contraindicated, it is supported in the literature and recommended in the guidelines of international health authorities. Therefore, its application is desirable in a public health context in order to increase the current immunization coverage rates. The achievement of vaccination coverages to meet individual protection and public health prevention goals for both MMRV and meningococcal C conjugate vaccines is considered very important for infants >1 year. In conclusion, according to the collected evidence, the lack of specific indications for co-administration of these two types of vaccines on the SmPC should not represent a barrier and should be recognized in all countries in order to support health-care workers to use it as a good and safe practice.

Disclosure of potential conflicts of interest

No conflict of interest to declare

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