

Vaccination of older adults against RSV: the final pieces of the puzzle

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With an estimated 186,000-614,000 older adults hospitalised annually as a result of respiratory syncytial virus (RSV) infection and with no effective treatment, there is an unmet need to prevent these infections, many of which lead to death [1]. Fortunately, we are in a golden age of RSV vaccine development with five formulations targeting older adults currently emerging from phase III trials [2]. With favourable trial results appearing, policy makers are now asking crucial questions about how we will use these new pharmaceuticals to improve the health of our elderly population: which vaccine product should we use? should our vaccine campaign be seasonal? do we need annual boosters? which age group should be eligible?

Answers to these questions will rely in part on using mathematical models to project the population impact of each vaccine. Consequently, mathematical models have become an indispensable part of the policy maker's evidence toolkit. Indeed, one of the benefits of mathematical models is that they can capture the considerable uncertainty in both the epidemiology of RSV disease and the protection afforded by potential vaccines.

Van Effelterre et al. [3] use a mathematical model of RSV transmission to capture this uncertainty to predict the impact of an RSV vaccination programme targeted at adults 60 years and over in the United States. The study uses a Bayesian framework to incorporate existing knowledge on the epidemiology of RSV. The mathematical model itself is an adaptation of a previously published model that captures the uptick in older adult RSV incidence by assuming an elevated risk of disease

and clinical outcomes [4]. The van Effelterre et al. study uses scenario analyses to predict the impact of potential vaccination programmes with varied assumptions around vaccine protection. Specifically, the authors assume vaccine efficacy against acute respiratory infection (ARI) varies between 50–70%, that vaccine efficacy against onward transmission varies between 0–50%, and that the duration of vaccine protection is either 3 or 5 years. Here we look at whether the uncertainty in these three components of vaccine protection reflect true gaps in our knowledge about the current suite of RSV vaccines, and, importantly, does this uncertainty matter when considering whether to introduce a vaccination programme.

First, and surprisingly, the efficacies for each of the four vaccines aimed at older adults are remarkably consistent against trial endpoints, including ARI and the more severe lower respiratory tract infection (LRTI) [5–10]. Specifically, the midpoint estimates range between 62–71% against RSV-associated ARI and 80–86% for RSV-associated LRTI, consistent with what was assumed in van Effelterre et al. Notwithstanding the rather wide confidence intervals around these estimates, these trials offer the first promise of reducing the considerable burden of severe respiratory disease in the elderly population. What is less known is the vaccine efficacy against very severe disease and death, with only one trial suggesting that efficacy against severe disease is higher at 94% (95% CI 62–100) [5]. Thus, for studies such as van Effelterre et al. that want to predict the impact of a vaccine programme on hospitalisations and death, we may have to wait for more information on the full range of clinical benefits from these vaccines. However, if these early indications are correct and consistent across all vaccines, the predicted reduction in hospitalisations and deaths by van Effelterre et al. would underestimate the vaccine impact, all else being equal.

Second, the completed trials do not give us any indication on how vaccines prevent onward transmission, instead measuring disease endpoints rather than infection or infectiousness (although see [11]). However, consistent with many studies describing vaccination of the elderly population, van Effelterre et al.'s work concludes that assumptions about the infectiousness of vaccinated individuals who become infected matter little to the impact of any vaccination programme. This invariance arises because the model assumes few opportunities for pathogen transmission between older adults and other individuals, consistent with studies in the US and elsewhere [12,13]. Consequently, despite vaccine efficacy against infection or infectiousness being unknown, this is unlikely to significantly influence vaccine impact and the cost-effectiveness of implementation.

Finally, there is an important knowledge gap around the duration of protection of these vaccines, because all completed trials that report results are powered to evaluate efficacy up to one year after vaccination. While van Effelterre et al. simply assume booster vaccines are given prior to any vaccine waning, thus maintaining the considerable reduction in disease burden, the implications of this uncertainty are of crucial importance to the efficiency and affordability of widespread vaccine roll-out. Van Effelterre et al. calculate the 'number needed to vaccinate' which is a measure of efficiency of the vaccine programme, and equivalent to the number of administered doses necessary to prevent one RSV-associated ARI. When the model assumed that vaccine duration dropped from 5 years to 3 years, there was, unsurprisingly, a proportional increase in the number needed to vaccinate, from 6–12 to 10–20. Although van Effelterre et al. did not calculate this explicitly, to achieve the same clinical impact as the base case predictions if a booster were to be needed every year – as it is the case with influenza – the number needed to vaccinate would increase five-fold, as would the total cost of the vaccine programme. Consequently, van Effelterre et al. implicitly highlights the importance of evaluating multi-season vaccine efficacy before any decisions about widespread roll-out are made.

While van Effelterre et al. strengthen the consensus that vaccination of older adults against RSV has the potential to significantly reduce RSV seasonal burden, it simply highlights, but does not fill, the pressing knowledge gaps that need resolution. With a large and growing older adult population, these vaccines will, rightly, come under intense scrutiny about their affordability and cost-effectiveness. And with little to distinguish each vaccine product's efficacy against acute respiratory infection, the decision to implement a vaccine programme, and which vaccine to choose, will likely rest on the vaccine's duration of disease protection, its protection against very severe outcomes and death and, ultimately, its price.

1. Shi T, Denouel A, Tietjen AK, et al. Global Disease Burden Estimates of Respiratory Syncytial Virus-Associated Acute Respiratory Infection in Older Adults in 2015: A Systematic Review and Meta-Analysis. *J Infect Dis* **2020**; 222:S577–S583. Available at: <http://dx.doi.org/10.1093/infdis/jiz059>.
2. RSV Vaccine and mAb Snapshot. Available at: https://media.path.org/documents/RSV-Snapshot_03JAN2023_HighResolution.pdf. Accessed 3 September 2023.

3. Van Effelterre T, Hens H, White LJ, Gravenstein S, Bastian AR, Buyukkaramikli N, Cheng C-Y, Hartnett J, Krishnarajah G, Weber K, Hernandez Pastor L. Modeling Respiratory Syncytial Virus Adult Vaccination in the United States with a Dynamic Transmission Model. *Clin Infect Dis*
4. Hodgson D, Pebody R, Panovska-Griffiths J, Baguelin M, Atkins KE. Evaluating the next generation of RSV intervention strategies: a mathematical modelling study and cost-effectiveness analysis. *BMC Med* **2020**; 18:348. Available at: <http://dx.doi.org/10.1186/s12916-020-01802-8>.
5. GSK's older adult respiratory syncytial virus (RSV) vaccine candidate shows 94.1% reduction in severe RSV disease and overall vaccine efficacy of 82.6% in pivotal trial. 2022. Available at: <https://www.gsk.com/en-gb/media/press-releases/gsk-s-older-adult-respiratory-syncytial-virus-rsv-vaccine-candidate/>. Accessed 9 March 2023.
6. Pfizer Announces Positive Top-Line Data from Phase 3 Trial of Older Adults for its Bivalent Respiratory Syncytial Virus (RSV) Vaccine Candidate. Available at: <https://www.pfizer.com/news/press-release/press-release-detail/pfizer-announces-positive-top-line-data-phase-3-trial-older>. Accessed 9 March 2023.
7. Janssen Announces Respiratory Syncytial Virus (RSV) Adult Vaccine Candidate Maintains High Efficacy Regardless of Lower Respiratory Tract Disease Severity. Available at: <https://www.janssen.com/janssen-announces-respiratory-syncytial-virus-rsv-adult-vaccine-candidate-maintains-high-efficacy>. Accessed 9 March 2023.
8. Falsey AR, Williams K, Gymnopoulos E, et al. Efficacy and Safety of an Ad26.RSV.preF-RSV preF Protein Vaccine in Older Adults. *N Engl J Med* **2023**; 388:609–620. Available at: <http://dx.doi.org/10.1056/NEJMoa2207566>.
9. Moderna Announces mRNA-1345, an Investigational Respiratory Syncytial Virus (RSV) Vaccine, Has Met Primary Efficacy Endpoints in Phase 3 Trial in Older Adults. Available at: <https://investors.modernatx.com/news/news-details/2023/Moderna-Announces-mRNA-1345-an-Investigational-Respiratory-Syncytial-Virus-RSV-Vaccine-Has-Met-Primary-Efficacy-Endpoints-in-Phase-3-Trial-in-Older-Adults/default.aspx>. Accessed 9 March 2023.
10. Papi A, Ison MG, Langley JM, et al. Respiratory Syncytial Virus Prefusion F Protein Vaccine in Older Adults. *N Engl J Med* **2023**; 388:595–608. Available at: <http://dx.doi.org/10.1056/NEJMoa2209604>.
11. Sadoff J, De Paepe E, DeVincenzo J, et al. Prevention of Respiratory Syncytial Virus Infection in Healthy Adults by a Single Immunization of Ad26.RSV.preF in a Human Challenge Study. *J Infect Dis*

2022; 226:396–406. Available at: <https://academic.oup.com/jid/article-pdf/226/3/396/45583790/jiab003.pdf>. Accessed 10 March 2023.

12. Prem K, Cook AR, Jit M. Projecting social contact matrices in 152 countries using contact surveys and demographic data. *PLoS Comput Biol* **2017**; 13:e1005697. Available at: <http://dx.doi.org/10.1371/journal.pcbi.1005697>.
13. Zagheni E, Billari FC, Manfredi P, Melegaro A, Mossong J, Edmunds WJ. Using time-use data to parameterize models for the spread of close-contact infectious diseases. *Am J Epidemiol* **2008**; 168:1082–1090. Available at: <http://dx.doi.org/10.1093/aje/kwn220>.