# THE LANCET **Infectious Diseases**

# **Supplementary appendix**

**This online publication has been corrected. The corrected version first appeared at thelancet.com/infection on July 13, 2022.** 

This appendix formed part of the original submission. We post it as supplied by the authors.

Supplement to: Parker EPK, Desai S, Marti M, et al. Emerging evidence on heterologous COVID-19 vaccine schedules—To mix or not to mix? *Lancet Infect Dis* 2022; published online March 9. https://doi.org/10.1016/S1473-3099(22)00178-5.

# **Appendix**

The material below is adapted with permission from the WHO's *Interim recommendations for heterologous COVID-19 vaccine schedules* (published under a CC BY-NC-SA 3.0 IGO license).1

# **Review methods**

*Search strategy for review of COVID-19 vaccine response following heterologous primary or boosting schedules*

A search in MEDLINE was performed to identify articles published between 1 July 2020 and 19 November 2021 using the search terms listed below.



Preprints in *medRxiv* were identified using the search strategy given below, implemented on 22 November 2021 with the package *medrxivr* using the programming language R.



The search was extended by scanning the reference lists of included articles and by consulting experts in the field. Studies published after the formal searches described above were included, up to a final cut-off of 6 December 2021.

### *Scope and eligibility criteria*

Duplicates were removed, and titles and abstracts were screened to identify articles reporting on the safety, immunogenicity, and/or vaccine effectiveness (VE) of a heterologous COVID-19 vaccine schedule involving a combination of WHO Emergency Use Listing (EUL) COVID-19 vaccines from multiple platforms (e.g. a vectored vaccine followed by an mRNA vaccines). At the time of the search, the list of WHO EUL products included Ad26.COV2.S (Janssen), BIBP (Sinopharm), BNT162b2 (Pfizer/BioNTech), ChAdOx1-S (AstraZeneca), CoronaVac (Sinovac), Covaxin (Bharat), and mRNA-1273 (Moderna). Articles reporting heterologous vaccine outcomes for fewer than 10 non-immunocompromised individuals were excluded. Multiple reports on the same population were combined under a single study ID during data extraction.

#### *Data extraction*

Data were extracted on study location, design, size, vaccine schedules, and dosing interval across all included studies. For studies of VE, we extracted data on: start and end dates; duration of follow-up (average and range); variant profile; and adjusted VE estimates, stratified by vaccine schedule and outcome (infection, symptomatic disease, hospitalisation, and/or intensive care unit admission).

For studies reporting on antibody response, we extracted data on: timing of sample collection after the final dose; binding antibody assay endpoint (target and assay); neutralizing antibody assay endpoint (measurement and target); and average binding and neutralizing antibody levels, stratified by vaccine schedule. We used Plot Digitizer<sup>3</sup> software to extract average (mean/median) antibody concentrations where these were not reported directly. We prioritised RBD-specific binding antibody concentrations and wild-type-specific neutralizing antibody concentrations where multiple endpoints were reported. Where antibody responses were reported at multiple timepoints, samples obtained 4 weeks after vaccination (or the nearest available timepoint) were selected. Formal analysis of reactogenicity data was beyond the scope of this review, but key trends are summarised in the main text. Cellular immunity endpoints were beyond the scope of this review.

Studies were included in the quantitative synthesis of antibody response rates if they reported post-vaccination antibody response data (binding or neutralizing antibody concentrations) for comparable heterologous and homologous schedules. These were used to calculate the ratio of antibody concentrations between heterologous and homologous vaccine groups, whereby a ratio of >1 indicates higher antibody levels among heterologous vaccine recipients. Given the multiplicity of vaccine products and schedules involved, we did not seek to combine these ratios into pooled estimates. Comparisons were excluded if they involved a different number of doses overall (e.g. Ad26.COV2.S–BNT162b2 vs BNT162b2–BNT162b2–BNT162b2).

Ad26.COV2.S can be given as a one-dose or two-dose primary series. For the purposes of this review, a two-dose heterologous series involving Ad26.COV2.S alongside another COVID-19 vaccine was considered a heterologous primary series.



**Supplementary Figure 1. Flow diagram.** \*Includes studies published after 19 November 2021; † identified via bibliographies and expert recommendation; § includes studies reporting on SARS-CoV-2 infection rates by vaccine group without vaccine effectiveness estimates. FIGW diagram. Includes studies published and 17 November 2021,



**Supplementary Figure 2. Comparative immunogenicity of COVID-19 vaccine schedules involving heterologous versus homologous platforms.** Post-vaccination antibody ratios are presented relative to homologous inactivated vaccine schedules (top row), homologous vectored vaccine schedules (middle row), and homologous mRNA vaccine schedules (bottom row). Study-level data including citations details are provided in appendix p 5 and pp 9–14. The number of estimates available for each comparison is indicated in italics. Ab, antibody; INA, inactivated; VEC, vectored.



**Supplementary Figure 3. Comparative immunogenicity of COVID-19 vaccine schedules involving heterologous versus homologous platforms.** Post-vaccination antibody ratios are presented relative to (A) homologous inactivated vaccine schedules, (B) homologous vectored vaccine schedules, and (C) homologous mRNA vaccine schedules. Citations details are provided in appendix pp 9–14. For studies in which the number of participants differed between binding and neutralizing antibody measurements, we quote the higher number here. Some study arms are included in multiple comparisons (e.g. one homologous group used as a reference for multiple heterologous schedules). Ab, antibody; AZ, AstraZeneca (ChAdOx1-S); BH, Bharat (Covaxin); CT, clinical trial; INA, inactivated; OBS, observational; SP, Sinopharm (BIBP); SV, Sinovac (CoronaVac); VEC, vectored.

#### **Supplementary Table 1. Evidence of COVID-19 vaccine effectiveness for schedules involving heterologous platforms.**

The table below includes studies reporting estimates of VE following heterologous vaccine series involving a combination of WHO EUL COVID-19 vaccine. VE estimates for RNA vaccines (i.e. BNT162b2 and mRNA-1273) were combined where possible. Vaccination groups without a relevant heterologous comparator (e.g. those involving a single dose) were excluded.







AZ: AstraZeneca (ChAdOx1-S); BNT: Pfizer-BioNTech (BNT162b2); CI: confidence interval; d: days; ICU: intensive care unit; MOD: Moderna (mRNA-1273); n.r.: not recorded; RNA: mRNA vaccines (BNT162b2) or mRNA-1273); SP: Sinopharm (BIBP) SV: Sinovac (CoronaVac); VE: vaccine effectiveness.

<sup>a</sup> Mean/median interval rounded to nearest week, or range if average interval not specified.

b VE estimate were adjusted for age, sex, heritage, hospital admission, and comorbidity.

 $\epsilon$  Median follow-up of 58 days for controls (range 0–265). Interval between doses varied from 3 to  $\geq$ 16 weeks, with >70% in the range of 7–11 weeks. Data for British Columbia are displayed. Similar findings were reported for Quebec. Over 99% of heterologous vaccine recipients received ChAdOx1-S first. VE estimates were adjusted for age, sex, epidemiological week, and region. Combined rather than vaccine-specific VE are reported for mRNA vaccines.

<sup>d</sup> VE estimates were adjusted for age, sex, baseline vaccination date, home-maker service, place of birth, education, and diagnoses at baseline.

<sup>e</sup> VE estimates were adjusted for age, sex, country of birth, and crowded living conditions.

<sup>f</sup> VE estimates were adjusted for age group, sex, chronic conditions, contact setting, month, and vaccination status of index case.

<sup>g</sup> Vaccine groups were matched by age, sex, region, and date of second dose.

h Population comprised adults over 50 years of age. Absolute VE relative to unvaccinated individuals is reported. Relative VE for AZ-AZ-BNT vs AZ-AZ (140+ days since second dose) was 87% (95% CI: 85–89%); relative VE for BNT-BNT vs BNT-BNT (140+ days since second dose) was 84% (95% CI: 82–86%). VE estimates were adjusted for age, sex, deprivation, ethnic group, care home residence status, region, calendar week of onset, health and social care worker status, clinical risk group, extreme clinical vulnerability, immunosuppression status, and prior positive testing.

<sup>i</sup> VE estimates were adjusted for age and sex, among others (full list not specified in available data).

#### **Supplementary Table 2. Relative binding and neutralizing antibody concentrations for heterologous versus homologous vaccine schedules involving a combination of WHO EUL COVID-19 vaccines. Data are displayed for (A) inactivated vaccines; (B) vectored vaccines; and (C) mRNA vaccines.**

Comparisons were included if antibody concentrations were reported for comparable heterologous and homologous schedules. Comparisons were excluded if they involved a different number of doses overall (e.g. JNJ-MOD vs MOD-MOD-MOD). Where antibody responses were reported at multiple timepoints, samples obtained 4 weeks after vaccination (or the nearest available timepoint) were selected. RBD-specific binding antibody concentrations and wild-type-specific neutralizing antibody concentrations were prioritised where multiple endpoints were reported. Some study arms are included in multiple comparisons (e.g. one homologous group used as a reference for multiple heterologous schedules).



**(A) WHO EUL inactivated vaccines: heterologous vs homologous schedules**



# **(B) WHO EUL vectored vaccines: heterologous vs homologous schedules**







#### **(C) WHO EUL RNA vaccines: heterologous vs homologous schedules**





† approximate values extracted using Plot Digitizer software; § excluded from summary plot as metric is not a titre or concentration and is therefore not comparable to other ratios.

AU: arbitrary units; AZ: AstraZeneca (ChAdOx1-S); BH: Bharat (Covaxin); BNT: Pfizer-BioNTech (BNT162b2); CI: confidence interval; CT: clinical trial (randomized or non-randomized); EUL: Emergency Use Listing (WHO); GMC: geometric mean concentration; Ig: binding antibody; IQR: interquartile range; IU: international units; JNJ: Janssen (Ad26.COV2.S); MOD: Moderna (mRNA-1273); *N*: number; NAb: neutralizing antibody; NT: neutralization test; NT50: 50% neutralizing antibody titre; OBS: observational study; PRNT: plaque reduction neutralization test; RBD: receptor binding domain; RNA: mRNA vaccines (BNT162b2 or mRNA-1273); SP: Sinopharm (BIBP); SV: Sinovac (CoronaVac); sVNT: surrogate virus neutralization test; WT: wild-type.

Vaccine platform colour coding: green = inactivated; blue = mRNA; orange = vectored. The colour represents the heterologous component that differentiates the schedule from the homologous schedule being used as a reference.

<sup>a</sup> Average interval rounded to nearest week, or range if average interval not specified.

b The administration of SP to individuals who had previously received two doses of SV was considered a homologous boost for the purposes of this comparison. Separate groups received BNT doses at 30 μg (full  $\frac{1}{2}$ dose) and 15 μg (fractional dose); the 30 μg recipients were included here, reflecting the formulation in the WHO EUL.

c Shared with SAGE on 5 October 2021.

d Comparisons involving different numbers of doses overall were excluded (e.g. JNJ-BNT vs BNT-BNT-BNT).

e Data for individuals aged >55 years (as opposed to <55 years) included given the presence of homologous RNA and homologous vectored comparators.

<sup>f</sup>Trial involved 18 study sites that were split into three groups, with multiple EUL and non-EUL COVID-19 vaccine products per group. Comparisons involving non-EUL vaccine products or fractional doses of EUL products were excluded. Some comparisons involved participants from separate study groups.

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