# **CO RONAVIRUS**

# **History of primary-series and booster vaccination and protection against Omicron reinfection**

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**Laboratory evidence suggests a possibility of immune imprinting for severe acute respiratory syndrome coro**navirus 2 (SARS-CoV-2) infection. We investigated the differences in the incidence of SARS-CoV-2 reinfection in a **cohort of persons who had a primary Omicron infection, but different vaccination histories using matched, national, retrospective, cohort studies. Adjusted hazard ratio for reinfection incidence, factoring adjustment for** differences in testing rate, was 0.43 [95% confidence interval (CI): 0.39 to 0.49] comparing history of two-dose vaccination to no vaccination, 1.47 (95% CI: 1.23 to 1.76) comparing history of three-dose vaccination to twodose vaccination, and 0.57 (95% CI: 0.48 to 0.68) comparing history of three-dose vaccination to no vaccination. **Divergence in cumulative incidence curves increased markedly when the incidence was dominated by BA.4/BA.5 and BA.2.75\* Omicron subvariants. The history of primary-series vaccination enhanced immune protection against Omicron reinfection, but history of booster vaccination compromised protection against Omicron reinfection. These findings do not undermine the public health utility of booster vaccination.**

**INTRODUCTION**

Three years into the coronavirus disease 2019 (COVID-19) pandemic, the global population carries heterogeneous immune histories derived from various exposures to infection, viral variants, and vaccination (*1*). Laboratory evidence suggests the possibility of immune imprinting, a negative impact of vaccination on subsequent protective immunity against severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) induced by vaccination or infection, or a combination of both (*1*–*4*). Epidemiological evidence for immune imprinting in immune histories related to infection was recently investigated, but no evidence was found for imprinting compromising protection against B.1.1.529 (Omicron) subvariants (*5*). A pre-Omicron infection followed by an Omicron reinfection enhanced protection against a second Omicron reinfection (*5*).

We investigated epidemiological evidence for imprinting in immune histories related to vaccination using matched,

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retrospective cohort studies conducted on the total population of Qatar from the onset of the Omicron wave on 19 December 2021 (*6*) through 15 September 2022. We compared the incidence of SARS-CoV-2 reinfection in the national cohort of individuals who had a primary documented Omicron infection after primary-series (two-dose) vaccination (designated as the two-dose cohort) to that in the national cohort of individuals with a documented primary Omicron infection, but no vaccination history (designated as the unvaccinated cohort). Analogously, we also compared reinfection incidence in those who had a documented primary Omicron infection after booster (third dose) vaccination (designated as the three-dose cohort) to each of the two-dose and unvaccinated cohorts.

These immune histories were investigated because of specific immunological scenarios observed in immunological laboratory data (*1*) because of their pervasiveness in the global population and because of their potential relevance to the protection of bivalent booster vaccination that is being scaled up in different countries.

A documented primary Omicron infection was defined as the first record of a SARS-CoV-2–positive polymerase chain reaction (PCR) or rapid antigen test after the onset of the Omicron wave in Qatar on 19 December 2021 (*6*) in an individual that had no record of a prior pre-Omicron infection. SARS-CoV-2 reinfection was defined, per the conventional definition in the literature, as a documented infection ≥90 days after an earlier infection, to avoid misclassifying prolonged SARS-CoV-2 positivity as reinfection if a shorter time interval is used (*6*–*8*). Matched pairs were followed from 90 days after the primary Omicron infection to record the incidence of SARS-CoV-2 reinfection.

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#### **RESULTS**

#### **Two-dose cohort versus unvaccinated cohort**

Figure S1 shows the study population selection process. Table 1 describes the baseline characteristics of the full and matched cohorts. Matched cohorts each included 56,802 individuals.

The median date of the second vaccine dose for the two-dose cohort was 9 June 2021. The median duration between the second dose and the start of follow-up was 312 days [interquartile range (IQR), 264 to 352 days]. The median duration of follow-up was 157 days (IQR, 140 to 164 days) for the two-dose cohort and 157 days (IQR, 139 to 164 days) for the unvaccinated cohort (Fig. 1A). There were 573 reinfections in the two-dose cohort and 1044 reinfections in the unvaccinated cohort during follow-up (fig. S1). None progressed to severe, critical, or fatal COVID-19.

The cumulative incidence of reinfection was 1.4% (95% CI: 1.2 to 1.5%) for the two-dose cohort and 2.4% (95% CI: 2.2 to 2.5%) for



**Fig. 1. Cumulative incidence of reinfection among those who had a primary infection with an Omicron subvariant.** Estimates of the cumulative incidence of reinfection using the Kaplan-Meier estimator are presented: after two-dose vaccination compared to no vaccination (**A**) and after three-dose vaccination compared to two-dose vaccination (**B**).

the unvaccinated cohort, after 165 days of follow-up (Fig. 1A). In the first 70 days of follow-up, the incidence was dominated by BA.2 (*9*–*11*). Subsequently, the incidence was dominated by BA.4/ BA.5 (*12*), and then by BA.2.75\* (*13*) (predominantly BA.2.75.2). Divergence between the cumulative incidence curves increased markedly when incidence was no longer dominated by BA.2.

The hazard ratio comparing the incidence of reinfection in the two-dose cohort to that in the unvaccinated cohort, adjusted for matching factors, was 0.59 (95% CI: 0.53 to 0.67; Table 2). The adjusted hazard ratio appeared stable by the month of follow-up (Fig. 2A). The proportion of individuals who had a test during follow-up was 48.9% for the two-dose cohort and 37.0% for the unvaccinated cohort. The testing frequency was 0.93 and 0.67 tests per person, respectively. Adjusting the hazard ratio additionally for differences in testing rate between cohorts yielded an adjusted hazard ratio of 0.43 (95% CI: 0.39 to 0.49).

#### **Three-dose cohort versus two-dose cohort**

Figure S2 shows the study population selection process. Table 1 describes the baseline characteristics of the full and matched cohorts. Matched cohorts each included 30,541 individuals.

Median dates of the second and third vaccine doses for the threedose cohort were 26 March 2021 and 6 December 2021, respectively. The median date of the second vaccine dose for the two-dose cohort was 11 May 2021. The median duration between the third dose and the start of follow-up was 124 days (IQR, 103 to 143 days), and between the second dose and the start of follow-up was 334 days (IQR, 286 to 371 days). The median duration of follow-up was 157 days (IQR, 135 to 164 days) in the three-dose cohort and 157 days (IQR, 137 to 164 days) in the two-dose cohort (Fig. 1B). There were 480 reinfections in the three-dose cohort and 248 reinfections in the two-dose cohort during follow-up (fig. S2). None progressed to severe, critical, or fatal COVID-19.

The cumulative incidence of reinfection was 2.1% (95% CI: 1.9 to 2.3%) for the three-dose cohort and 1.1% (95% CI: 1.0 to 1.3%) for the two-dose cohort, after 165 days of follow-up (Fig. 1B). In the first 70 days of follow-up, the incidence was dominated by BA.2 (*9*–*11*). Subsequently, the incidence was dominated by BA.4/BA.5 (*12*), and then by BA.2.75\* (*13*). The divergence between the cumulative incidence curves increased markedly when incidence was no longer dominated by BA.2.

The adjusted hazard ratio comparing the incidence of reinfection in the three-dose cohort to that in the two-dose cohort was 1.96 (95% CI: 1.64 to 2.34; Table 2). The adjusted hazard ratio appeared stable by the month of follow-up, but with wide 95% confidence intervals (CI) (Fig. 2B). The proportion of individuals who had a test during follow-up was 63.1% for the three-dose cohort and 49.0% for the two-dose cohort. The testing frequency was 1.39 and 0.98 tests per person, respectively. Adjusting the hazard ratio additionally for differences in testing rate between cohorts yielded an adjusted hazard ratio of 1.47 (95% CI: 1.23 to 1.76).

In the first sensitivity analysis with the cohorts being matched by the Charlson comorbidity index, instead of the number of chronic coexisting conditions, the adjusted hazard ratio, including the adjustment for the differences in testing rate, was 1.39 (95% CI: 1.16 to 1.67) (table S1).

In the second sensitivity analysis with the cohorts being matched additionally by primary-series vaccine type (two doses of BNT162b2 or two doses of mRNA-1273), the adjusted hazard



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\*\*The reason for testing of the SARS-CoV-2 test that ascertained the Omicron infection that made the person eligible for inclusion in the cohort.

**Table 2. Hazard ratios for the incidence of SARS-CoV-2 reinfection in studies investigating immune protection among those who had a primary infection with an Omicron subvariant, but different vaccination histories.** CI, confidence interval.



\*Cohorts were matched exactly one-to-one by sex, age, nationality, number of coexisting conditions, as well as the SARS-CoV-2 testing method, the reason for SARS-CoV-2 testing, and the calendar week of the SARS-CoV-2 test of the primary Omicron infection. + Cox regression analysis adjusted for sex, 10-year age groups, 10 nationality groups, number of coexisting conditions, as well as the SARS-CoV-2 testing method, the reason for SARS-CoV-2 testing, and the calendar week of the SARS-CoV-2 test of the primary Omicron infection.

ratio, including the adjustment for the differences in testing rate, was 1.43 (95% CI: 1.19 to 1.71) (table S1). In the subgroup analysis including only BNT162b2-vaccinated individuals, the adjusted hazard ratio was 1.39 (95% CI: 1.15 to 1.68). In the subgroup analysis including only mRNA-1273–vaccinated individuals, the adjusted hazard ratio was 1.83 (95% CI: 1.03 to 3.28).

#### **Three-dose cohort versus unvaccinated cohort**

Figure S3 shows the study population selection process. Table S2 describes the baseline characteristics of the full and matched cohorts. The cumulative incidence of reinfection is shown in fig. S4A.

The adjusted hazard ratio comparing the incidence of reinfection in the three-dose cohort to that in the unvaccinated cohort was 1.10 (95% CI: 0.92 to 1.31; Table 2). The adjusted hazard ratio appeared stable by the month of follow-up, but with wide 95% CIs (fig. S4B). The proportion of individuals who had a test during follow-up was 66.4% for the three-dose cohort and 36.8% for the unvaccinated cohort. The testing frequency was 1.46 and 0.70 tests per person, respectively. Adjusting the hazard ratio additionally for differences in testing rate between cohorts yielded an adjusted hazard ratio of 0.57 (95% CI: 0.48 to 0.68).

The results of this additional study confirm the relative differences in the incidence of reinfection observed in the first two studies, with incidence being the lowest among the two-dose cohort and the highest among the unvaccinated cohort.

#### **DISCUSSION**

Primary-series vaccination followed by a primary Omicron infection was associated with enhanced immune protection against Omicron reinfection compared to primary Omicron infection with no prior vaccination. This result is notable because the start of follow-up in this study was  $\sim$ 1 year after the two-dose primary series. Protection of the primary series against Omicron infection that is mediated by neutralizing antibodies should have fully waned by this time, considering how rapidly vaccine protection wanes against Omicron subvariants (*10*, *14*). This finding suggests that the primary Omicron infection may have stimulated other components of the immune system, specifically immune memory of the earlier primary-series immune response in a manner that enhanced protection against a subsequent Omicron reinfection, particularly against BA.4/BA.5 and BA.2.75\*.

Similar effects and effect sizes were observed recently in an analogous study (*5*). The incidence of reinfection among unvaccinated persons who had contracted an Omicron infection following an earlier pre-Omicron infection was lower than the incidence of reinfection among unvaccinated persons who had only an Omicron infection and no prior pre-Omicron infection (*5*). mRNA vaccines used in Qatar are based on index-virus design (*15*, *16*). The median duration between the first and second vaccine doses was <1 month (*17*). Given this short duration between doses, twodose vaccination counts perhaps as a single pre-Omicron immunological event. This may explain the similarity in both effect and effect size in these two studies, since in essence, both investigate immune protection elicited by a pre-Omicron immunological event followed by an Omicron immunological event, compared to protection of only a single Omicron event.





Fig. 2. Adjusted hazard ratio by the month of follow-up for SARS-CoV-2 reinfection among those who had a primary infection with an Omicron subvariant. Estimates of adjusted hazard ratios for reinfection that compare two-dose vaccination to no vaccination (**A**) and three-dose vaccination to two-dose vaccination (**B**). Analyses were performed on 56,802 and 30,541 matched pairs, respectively. Error bars indicate 95% CIs.

While two-dose vaccination was associated with enhanced protection against subsequent Omicron reinfection, three-dose vaccination was associated with reduced protection compared to that of two-dose vaccination. While this finding remainsto be explained, it is consistent with the immune response against the primary Omicron infection being compromised by differential immune imprinting in those who received a third booster dose, also consistent with laboratory data (*1*–*4*) and emerging epidemiologic data (*18*– *21*) on imprinting effects. The booster dose, a pre-Omicron immunological event, that occurred several months after the primaryseries vaccination, another pre-Omicron immunological event, may have trained the immune response to expect a specific narrow pre-Omicron challenge; thus, the response was inferior when the actual challenge was an immune-evasive Omicron subvariant. Repeat immunological events of the same kind (here pre-Omicron challenge) may be associated with compromised protection against a different kind of immunological event (here Omicron challenge). It is important to emphasize that the findings of this study should be regarded as preliminary and hypothesis-generating. While the observed effects are consistent with the concept of immune imprinting, they do not directly provide evidence for immune imprinting. Therefore, additional research, comprehensive understanding, and confirmation of this phenomenon are essential and warranted.

This imprinting effect appears related to the memory component of the immune response, perhaps explaining why the effect was observed only after the waning of the antibody-mediated short-term booster protection, as supported also by another study on the same population of the long-term effectiveness of booster vaccination (*18*). Those with a booster may have had their immune memory geared and narrowed down toward expecting a specific pre-Omicron challenge (*22*). The imprinting effect seems to arise from the mismatch between such specific immune memory and the actual substantially different immune challenge (*22*). The size of the imprinting effect appeared also to be larger for mRNA-1273-vaccinated persons than for BNT162b2-vaccinated persons, possibly because of the larger dose of the mRNA-1273 vaccine (*17*), and perhaps suggesting a dose-response relationship for the imprinting effect.

We investigated two immune histories with different observed effects. Primary-series vaccination followed by a primary Omicron infection enhanced immune protection against Omicron reinfection. Booster vaccination followed by a primary Omicron infection reduced immune protection against Omicron reinfection. This highlights the complexity of the immunity landscape at this stage of the pandemic, in which people have different immune histories. These findings, however, do not undermine the utility of booster vaccination, at least in the short term. The lower protection was observed only after the waning of the antibody-mediated short-term booster protection, as follow-up commenced >4 months after the booster, at a time when booster effectiveness is expected to be marginal (*10*, *14*, *18*). There is no question that the booster dose reduced infection incidence in the first 6 months after its administration, based on evidence from this same population (*9*, *10*, *18*, *23*). Nonetheless, the findings suggest that the shortterm effects of boosters may differ from their long-term effects.

Although we planned to investigate effectiveness against severe COVID-19, no reinfection in any cohort of the three studies progressed to severe, critical, or fatal COVID-19. Although some of the patients with COVID-19 were hospitalized, none reached the World Health Organization (WHO) classification of severe or critical COVID-19, and none ended up with COVID-19 death following the longitudinal review of their individual charts. This outcome is not unexpected given the lower severity of Omicron infections (*24*–*26*) and the strong protection of natural infection against severe COVID-19 at reinfection, estimated at 97% in this same population (*27*), as well as the long-term effectiveness of primary series and boosters against severe COVID-19 (*9*, *10*, *14*, *18*, *28*, *29*). While we were unable to quantify the same effects on COVID-19 severity, the results do not suggest imprinting compromising protection against severe COVID-19. This has also been supported by other analyses of the same population (*18*, *30*).

The central analysis in this study compares the incidence of infection among boosted persons versus those with only a primary series, both groups of which had an Omicron primary infection after vaccination. However, these two groups may not be immunologically comparable with respect to their ability to produce a strong immune response following vaccination and Omicron infection, and there could be other nonobserved differences between these cohorts. The three-dose group consists of individuals with three vaccine doses and a primary infection shortly after the third dose. By contrast, the two-dose group consists of individuals with only two vaccine doses and a primary infection long after their second dose. It is possible that the shorter duration between dose and Omicron infection in the three-dose group versus the two-dose group may have contributed to inferior immunological response to the Omicron infection, perhaps explaining the higher incidence among boosted persons thereafter. However, the negative imprinting effect observed in this study has now been also observed among groups who are immunologically comparable with respect to their ability to produce a strong immune response following vaccination and/or infection and in different study designs (*18*, *30*) arguing against this explanation of the study results.

Following the preprint of this article (*31*) it has been suggested that the conditioning on having infection may introduce bias that explains the higher incidence among boosted persons (*32*). Since the groups have different immune histories before primary infection, with one history more protective than the other one, the conditioning on having the infection may implicitly select persons with more propensity for infection in the group that had the more protective immune history before the primary infection. Persons in the three-dose group may have chosen to receive a third vaccine dose because they are aware that they have high levels of exposure, thereby also explaining the higher infection incidence among boosted persons.

However, if this bias existed, then its effect needs to be consistent throughout the time of follow-up, not only in one part of it as opposed to another. The results of the analyses presented here, and the earlier analysis for natural immunity (*5*), are not consistent with such a bias effect. There were no differences in incidence between the groups when incidence was due to BA.1/BA.2. The differences between the groups were observed only after the incidence was dominated by BA.4/BA.5, consistent with an immune imprinting effect rather than a bias effect.

Moreover, in the analysis comparing the history of pre-Omicron infection to no pre-Omicron infection (*5*), and in the analysis comparing the history of primary-series vaccination to no vaccination, a strong positive imprinting effect was found, opposite in direction to effect of this potential bias. If bias existed, then the already strong positive imprinting effect is substantially underestimated, an outcome that does not seem plausible given how strong the effect was already in the opposite direction. The effect size was also similar for both of these analyses, despite the differences in immune history, further supporting immune imprinting as an explanation of the study outcomes. Last, rigorous matching was implemented to balance infection exposure risk across the groups, and this may have minimized the effect of bias.

This study has limitations. We investigated the incidence of documented reinfections, but undocumented reinfections may have occurred. Unvaccinated individuals are a minority in Qatar and may not be truly immune-naïve due to undocumented prior infections or undocumented vaccinations, perhaps outside the country, especially now that we are 3 years into this pandemic. Bias due to unequal depletion of the unvaccinated versus vaccinated susceptible population may underestimate vaccine protection (*33*). Effects were observed long after vaccination, but long-term effects are more likely to be affected by bias than short-term effects. With Qatar's young population, our findings may not be generalizable to older individuals or to other countries where elderly citizens constitute a large proportion of the total population.

Testing rates differed between cohorts suggesting the possibility of bias due to differential outcome ascertainment. Receiving a booster dose could be correlated with health-seeking behavior that would result in more frequent testing. Different travel testing guidelines for vaccinated and unvaccinated individuals affect also the testing rate. This bias due to testing differences may affect the estimated effects and may explain the higher infection incidence among boosted persons. However, the adjustment for the differences in testing rate showed overall similar findings to the main-analysis findings. While the adjustment quantitatively affected the estimated hazard ratios, the adjusted analyses confirmed the finding of higher incidence among boosted persons compared to those with only a primary series. Note also that the study matched observable confounders across cohorts to control for potential effects of differentials in testing across confounder values. The ratio of testing frequency in the matched cohorts was also overall stable over time of follow-up suggesting the absence of substantial differential changes in behavior over time (fig. S5). Therefore, bias due to differences in testing may not explain the negative imprinting effect observed in this study.

Home-based rapid antigen testing is not documented in Qatar and is not factored in these analyses. However, there is no reason to believe that home-based testing could have differentially affected the following cohorts to alter study estimates. Matching was done while factoring key sociodemographic characteristics of the population (*34*–*38*), such as nationality, age, and sex, and this may also have controlled or reduced differences in home-based testing between cohorts. Nationality, age, and sex provide a powerful proxy for socioeconomic status in Qatar (*34*–*38*). Nationality is also strongly associated with occupation (*34*, *36*–*38*).

We did not have access to other demographic data or the complete medical records of individuals, which limited our ability to control for other potential confounders such as variations in treatments or medications or the duration of a participant's interaction with the medical system. Comorbidities were ascertained and classified on the basis of the ICD-10 codes for chronic conditions as recorded in a compilation of the electronic health record encounters

of each individual in the Cerner-system national database that includes all citizens and residents registered in the national and universal public health care system. Individuals who have comorbidities but never sought care in the public health care system, or seek care exclusively in private health care facilities, were classified as individuals with no comorbidity due to the absence of recorded encounters for them. This misclassification bias and these aspects are not likely to have affected the study results considering that the proportion of persons with serious coexisting conditions is very small. The study was conducted on a young and healthy population, using large national samples. In Qatar, the population consists mainly of individuals of working age who are healthy by recruitment (*39*), with a very small proportion having severe or multiple chronic conditions (*34*, *40*). The national list of vaccine prioritization included only 19,800 individuals of all age groups with serious comorbid conditions to be prioritized in the first phase of the vaccine rollout (*28*). Note that the results were invariable by matching the Charlson comorbidity index instead of the number of coexisting conditions.

As an observational study, investigated cohorts were neither blinded nor randomized, so unmeasured or uncontrolled confounding cannot be excluded. Although matching covered key factors affecting infection exposure (*34*–*38*), it was not possible for other factors such as geography or occupation, for which data were unavailable. However, Qatar is essentially a city-state, and infection incidence was broadly distributed across neighborhoods. Nearly 90% of Qatar's population are expatriates from more than 150 countries, who come here for employment (*34*). Nationality, age, and sex provide a powerful proxy for socioeconomic status and occupation in this country (*34*–*38*).

The matching prescription used in this study was investigated in previous studies of different epidemiologic designs and using control groups to test for null effects (*17*, *28*, *29*, *41*, *42*). These control groups included unvaccinated cohorts versus vaccinated cohorts within 2 weeks of the first dose (*28*, *29*, *41*, *42*), when vaccine protection is negligible (*15*, *16*), and mRNA-1273– versus BNT162b2-vaccinated cohorts, also in the first 2 weeks after the first dose (*17*). These studies showed repeatedly and at different times during the pandemic that this prescription provides adequate control of differences in infection exposure (*17*, *28*, *29*, *41*, *42*), suggesting that the used matching may also have controlled for differencesin infection exposure in the present analyses. All analyses were implemented on Qatar's total population, perhaps minimizing the likelihood of bias.

In conclusion, primary-series vaccination followed by a primary Omicron infection enhanced immune protection against Omicron reinfection. However, booster vaccination followed by a primary Omicron infection had lower protection against Omicron reinfection than primary-series vaccination followed by a primary Omicron infection. These findings do not undermine the utility of booster vaccination in the short term but may point to potential complexities in designing boosters with optimal effects.

#### **MATERIALS AND METHODS**

# **Study population and data sources**

This study was conducted in the population of Qatar from the onset of the Omicron wave on 19 December 2021 (*6*) through 15 September 2022. It analyzed the national, federated databases for COVID-

19 laboratory testing, vaccination, hospitalization, and death, retrieved from the integrated, nationwide, digital health information platform. Databases include all SARS-CoV-2–related data with no missing information since the pandemic onset, such as all PCR tests, and from 5 January 2022 onward, all rapid antigen tests conducted at health care facilities. SARS-CoV-2 testing in the health care system in Qatar is done at a mass scale, and mostly for routine reasons, where about 5% of the population is tested every week (*9*, *28*). About 75% of those diagnosed are diagnosed not because of the appearance of symptoms but because of routine testing (*9*, *28*). Every PCR test and an increasing proportion of the facility-based rapid antigen tests conducted in Qatar, regardless of location or setting, are classified on the basis of symptoms and the reason for testing (clinical symptoms, contact tracing, surveys or random testing campaigns, individual requests, routine health care testing, pretravel, at the port of entry, or other). All facilitybased testing done during follow-up in the present study was factored in the analyses of this study.

Rapid antigen test kits are available for purchase in pharmacies in Qatar, but the outcome of home-based testing is not reported nor documented in the national databases. Since SARS-CoV-2 test outcomes are linked to specific public health measures, restrictions, and privileges, testing policy and guidelines stress facility-based testing as the core testing mechanism in the population. While facility-based testing is provided free of charge or at low subsidized costs, depending on the reason for testing, home-based rapid antigen testing is de-emphasized and not supported as part of national policy. There is no reason to believe that home-based testing could have differentially affected the following matched cohorts to affect our results.

The infection detection rate is defined as the cumulative number of documented infections, that is, diagnosed and laboratory-confirmed infections, over the cumulative number of documented and undocumented infections. Serological surveys and other analyses suggest that a substantial proportion of infections in Qatar and elsewhere are undocumented (*35*–*38*, *43*–*45*). With the absence of recent serological surveys in Qatar, it is difficult to estimate the current or recent infection detection rate, but mathematical modeling analyses and their recent updates suggest that, at present, no less than 50% of infections are never documented (*35*, *46*).

Differences in testing rate during follow-up may introduce differential ascertainment of infection across the cohorts if routine testing varies by cohort. There was evidence for differences in the testing rate across the cohorts. These differences could result in different rates of undocumented infection before and during followup. To address these differences, analyses were conducted by further adjusting the hazard ratios in the Cox regressions for the differences in testing rate (please note below).

Qatar has unusually young, diverse demographics, in that only 9% of its residents are  $\geq$  50 years of age, and 89% are expatriates from over 150 countries (*34*, *40*). Qatar launched its COVID-19 vaccination program in December of 2020 using the BNT162b2 and mRNA-1273 vaccines (*17*). Detailed descriptions of Qatar's population and of the national databases have been reported previously (*9*, *23*, *28*, *34*, *47*).

# **Study design and cohorts**

Matched, retrospective, observational cohort studies were conducted to investigate epidemiological evidence for immune imprinting

In the first study, we compared the incidence of reinfection in the national cohort of individuals who had a primary Omicron infection after primary-series (two-dose) vaccination (designated as the two-dose cohort) to that in the national cohort of individuals who had a primary Omicron infection, but no vaccination history (designated as the unvaccinated cohort).

In the second study, we compared the incidence of reinfection in the national cohort of individuals who had a primary Omicron infection after booster (third dose) vaccination (designated as the three-dose cohort) to that in the two-dose cohort. In a third study, to confirm and complement the results of the first two studies, we compared the incidence of reinfection in the threedose cohort to that in the unvaccinated cohort. The majority of primary Omicron infections in these three studies involved the BA.2 subvariant (*9*–*11*).

SARS-CoV-2 reinfection was defined as a documented infection ≥90 days after an earlier infection, to avoid misclassifying prolonged positivity as reinfection (*6*–*8*). Children vaccinated with the pediatric dose of BNT162b2 and adults who received different vaccines were excluded. Classification of infection severity followed WHO guidelines for COVID-19 case severity (acute care hospitalizations) (*48*), criticality (intensive care unit hospitalizations) (*48*), and fatality (*49*).

# **Cohort matching and follow-up**

Cohorts were matched exactly one-to-one by sex, 10-year age group, nationality, and number of chronic coexisting conditions (none, one, two, three, or more comorbid conditions) to balance observed confounders between exposure groups that are related to infection risk in Qatar (*34*–*38*). Individuals who were first diagnosed with SARS-CoV-2 in a specific week in one cohort were matched to individuals who were first diagnosed with SARS-CoV-2 in that same calendar week in the comparator cohort, to ensure that matched pairs were exposed to the same Omicron subvariants and had a presence in Qatar at the same time. Cohorts were also matched exactly by testing method (PCR versus rapid antigen testing) and by reason for testing for the primary Omicron infection to control for potential differences in testing modalities between cohorts.

Matching was performed iteratively such that individuals in the comparator cohort were alive, had not been reinfected, and had maintained the same vaccination status at the start of follow-up. Each matched pair was followed from 90 days after the primary Omicron infection of the individual in the two-dose cohort for the study comparing the incidence of reinfection in that cohort with the unvaccinated cohort. Follow-up was from 90 days after the primary Omicron infection of the individual in the three-dose cohort for studies comparing the incidence of reinfection in that cohort to that in each of the two-dose and unvaccinated cohorts.

For exchangeability (*23*, *50*), both members of each matched pair were censored as soon as one of them received a new vaccine dose (change in vaccination status, that is, at earliest occurrence of an unvaccinated individual in the matched pair receiving the first dose, or the individual with two-dose vaccination receiving a third dose, or the individual with three-dose vaccination receiving a fourth dose). Accordingly, individuals were followed up until the first of any of the following events: a documented SARS-CoV-2 reinfection (defined as the first PCR-positive or rapid antigen–positive test after the start of follow-up, regardless of symptoms), a change in vaccination status (with matched-pair censoring), or death, or end of study censoring (15 September 2022).

# **Comorbidity classification**

Comorbidities were ascertained and classified on the basis of the ICD-10 codes for chronic conditions as recorded in the electronic health record encounters of each individual in the Cerner-system national database that includes all citizens and residents registered in the national and universal public health care system. The public health care system provides health care to the entire resident population of Qatar free of charge or at heavily subsidized costs, including prescription drugs. With the mass expansion of this sector in recent years, facilities have been built to cater to the specific needs of subpopulations. For example, tens of facilities have been built, including clinics and hospitals, in localities with a high density of craft and manual workers (*37*).

All encounters for each individual were analyzed to determine the comorbidity classification for that individual, including all laboratory data, as part of a recent national analysisto assess health care needs and resource allocation. The Cerner-system national database includes encounters starting from 2013, after this system was launched in Qatar. As long as each individual had at least one encounter with a specific comorbidity diagnosis based on clinical and laboratory data since 2013, this person was classified with this comorbidity.

Individuals who have comorbidities but never sought care in the public health care system, or seek care exclusively in private health care facilities, were classified as individuals with no comorbidity due to the absence of recorded encounters for them.

We did not have access to the complete medical records of individuals, which prevented us from assessing variations in treatments, medications, or the duration of participants' interaction with the medical system. However, the study was conducted on a young and healthy population, using large national samples. In Qatar, the population consists mainly of individuals of working age who are healthy by recruitment (*39*), with a very small proportion having severe or multiple chronic conditions (*34*, *40*). The national vaccine prioritization list included only 19,800 individuals across all age groups with serious comorbid conditions, who were given priority in the initial phase of the vaccine rollout, while the total population of Qatar is approximately 3 million people (*28*). These factors suggest that variations in treatments, medications, or the duration of participants' interaction with the medical system are unlikely to have substantially affected our results.

# **Laboratory methods and variant ascertainment** *Real-time reverse transcription polymerase chain reaction testing*

Nasopharyngeal and/or oropharyngeal swabs were collected for PCR testing and placed in a universal transport medium (UTM). Aliquots of UTM were as follows: (i) extracted on KingFisher Flex (Thermo Fisher Scientific, USA), MGISP-960 (MGI, China), or ExiPrep 96 Lite (Bioneer, South Korea) followed by testing with

real-time reverse transcription quantitative PCR (RT-qPCR) using TaqPath COVID-19 Combo Kits (Thermo Fisher Scientific, USA) on an ABI 7500 FAST (Thermo Fisher Scientific, USA); (ii) tested directly on the Cepheid GeneXpert system using the Xpert Xpress SARS-CoV-2 (Cepheid, USA); or (iii) loaded directly into a Roche cobas 6800 system and assayed with the cobas SARS-CoV-2 Test (Roche, Switzerland). The first assay targets the viral S, N, and ORF1ab gene regions. The second targets the viral N and E gene regions, and the third targets the ORF1ab and E gene regions. All PCR testing was conducted at the Hamad Medical Corporation Central Laboratory or Sidra Medicine Laboratory, following standardized protocols.

# *Rapid antigen testing*

SARS-CoV-2 antigen tests were performed on nasopharyngeal swabs using one of the following lateral flow antigen tests: Panbio COVID-19 Ag Rapid Test Device (Abbott, USA). SARS-CoV-2 Rapid Antigen Test (Roche, Switzerland), Standard Q COVID-19 Antigen Test (SD Biosensor, Korea), or CareStart COVID-19 Antigen Test (Access Bio, USA). All antigen tests were performed point-of-care according to each manufacturer's instructions at public or private hospitals and clinics throughout Qatar with prior authorization and training by the Ministry of Public Health (MOPH). Antigen test results were electronically reported to the MOPH in real time using the Antigen Test Management System which is integrated with the national COVID-19 database.

# *Classification of infections by variant type*

Surveillance for SARS-CoV-2 variants in Qatar is based on viral genome sequencing and multiplex real-time RT-qPCR variant screening (*51*) of random positive clinical samples (*28*, *42*, *52*– *55*), complemented by deep sequencing of wastewater samples (*53*, *56*, *57*). Further details on viral genome sequencing and multiplex RT-qPCR variant screening throughout the SARS-CoV-2 waves in Qatar can be found in previous publications (*6*, *9*, *10*, *12*, *23*, *28*, *42*, *52*–*55*, *58*–*60*).

# **COVID-19 severity, criticality, and fatality classification**

Classification of COVID-19 case severity (acute care hospitalizations) (*48*), criticality (intensive care unit hospitalizations) (*48*), and fatality (*49*) followed the WHO guidelines. Assessments were made by trained medical personnel independent of study investigators and using individual chart reviews, as part of a national protocol applied to every hospitalized patient with COVID-19. Each hospitalized patient with COVID-19 underwent an infection severity assessment every 3 days until discharge or death. We classified individuals who progressed to severe, critical, or fatal COVID-19 between the time of the documented infection and the end of the study based on their worst outcome, starting with death (*49*), followed by critical disease (*48*), and then severe disease (*48*).

Severe COVID-19 diseasewas defined per WHO classification as a SARS-CoV-2–infected person with "oxygen saturation of <90% on room air, and/or respiratory rate of >30 breaths/minute in adults and children >5 years old (or  $\geq$ 60 breaths/minute in children <2 months old or ≥50 breaths/minute in children 2–11 months old or ≥40 breaths/minute in children 1–5 years old), and/or signs of severe respiratory distress (accessory muscle use and inability to complete full sentences, and, in children, very severe chest wall indrawing, grunting, central cyanosis, or presence of any other general danger signs)" (*48*). Detailed WHO criteria for classifying SARS-

CoV-2 infection severity can be found in the WHO technical report (*48*).

Critical COVID-19 disease was defined per WHO classification as a SARS-CoV-2–infected person with "acute respiratory distress syndrome, sepsis, septic shock, or other conditions that would normally require the provision of life sustaining therapies such as mechanical ventilation (invasive or non-invasive) or vasopressor therapy" (*48*). Detailed WHO criteria for classifying SARS-CoV-2 infection criticality can be found in the WHO technical report (*48*).

COVID-19 death was defined per WHO classification as"a death resulting from a clinically compatible illness, in a probable or confirmed COVID-19 case, unless there is a clear alternative cause of death that cannot be related to COVID-19 disease (e.g., trauma). There should be no period of complete recovery from COVID-19 between illness and death. A death due to COVID-19 may not be attributed to another disease (e.g., cancer) and should be counted independently of preexisting conditions that are suspected of triggering a severe course of COVID-19." Detailed WHO criteria for classifying COVID-19 death can be found in the WHO technical report (*49*).

# **Oversight**

The Institutional Review Boards at Hamad Medical Corporation and Weill Cornell Medicine-Qatar approved this retrospective study with a waiver of informed consent. The study was reported according to the Strengthening the Reporting of Observational Studies in Epidemiology guidelines (table S3). The authors vouch for the accuracy and completeness of the data and for the fidelity of the study to the protocol. Data used in this study are the property of the Ministry of Public Health of Qatar and were provided to the researchers through a restricted-access agreement for the preservation of the confidentiality of patient data. The funders had no role in the study design; the collection, analysis, or interpretation of the data; or the writing of the manuscript.

# **Statistical analysis**

Eligible and matched cohorts were drawn from independent samples and described using frequency distributions and measures of central tendency and were compared using standardized mean differences (SMDs). An SMD of  $\leq 0.1$  indicated adequate matching (*61*). The cumulative incidence of reinfection (defined as the proportion of individuals at risk, whose primary endpoint during follow-up was reinfection) was estimated using the Kaplan-Meier estimator method (*62*). The incidence rate of reinfection in each cohort, defined as the number of identified reinfections divided by the number of person-weeks contributed by all individuals in the cohort, was estimated, with the corresponding 95% CI using a Poisson log-likelihood regression model with the Stata 17.0 *stptime* command.

Hazard ratios, comparing the incidence of reinfection in the cohorts and corresponding 95% CIs, were calculated using Cox regression, adjusted for the matching factors with the Stata 17.0 *stcox* command. The overall hazard ratio and the month-by-month hazard ratios in the Cox regression were additionally adjusted for differences in testing rate (low testers, intermediate testers, and high testers defined as persons having  $\leq$ 2, 3 to 6, and  $\geq$ 7 tests per person-year during follow-up, respectively). This additional adjustment was conducted because most SARS-CoV-2 testing in Qatar is done for routine reasons and not because of symptoms (*9*, *28*). About 75% of those diagnosed with the infection are diagnosed not because of the appearance of symptoms, but because of routine testing (*9*, *28*). Testing guidelines also differed by vaccination status (such as for travel-related testing) (*28*). Any differences in testing rate can potentially introduce differential ascertainment of infection across the cohorts if routine testing varies by cohort.

Sensitivity analyses were also conducted for the central analysis comparing the incidence of reinfection in the three-dose cohort to the two-dose cohort. In the first analysis, the cohorts were matched by the Charlson comorbidity index instead of the number of coexisting conditions. In the second analysis, the cohorts were matched additionally by primary-series vaccine type (two doses of BNT162b2 or two doses of mRNA-1273). Subgroup analyses were also conducted for the latter sensitivity analysis where the hazard ratios were calculated separately for each of the BNT162b2- and mRNA-1273–vaccinated individuals.

Schoenfeld residuals and log-log plots for survival curves were used to test the proportional hazards assumption. CIs were not adjusted for multiplicity; thus, they should not be used to infer definitive differences between groups. Interactions were not considered. Statistical analyses were conducted using Stata/SE version 17.0 (Stata Corporation, College Station, TX, USA).

# **Supplementary Materials**

**This PDF file includes:** Figs. S1 to S5 Tables S1 to S3

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confidentiality of patient data. Access can be obtained through a direct application for data accessto Her Excellency the Minister of Public. The raw data are protected and are not available due to data privacy laws. Data were available to authors through .csv files where information was downloaded from the CERNER database system (no links/accession codes were available to authors).

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