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Effectiveness of mRNA-1273, BNT162b2, and BBIBP-CorV vaccines against infection and mortality in children in Argentina, during predominance of delta and omicron covid-19 variants: test negative, case-control study

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ABSTRACT

OBJECTIVE

To estimate the effectiveness of a two dose vaccine schedule (mRNA-1273, BNT162b2, and BBIBP-CorV) against SARS-CoV-2 infection and covid-19 related death and short term waning of immunity in children (3-11 years old) and adolescents (12-17 years old) during periods of delta and omicron variant predominance in Argentina.

DESIGN

Test negative, case-control study.

SETTING

Database of the National Surveillance System and the Nominalized Federal Vaccination Registry of Argentina.

PARTICIPANTS

844 460 χ hildren and adolescents without previous SARS-CoV-2 infection eligible to receive primary vaccination schedule who were tested for SARS-CoV-2 by polymerase chain reaction or rapid antigen test from September 2021 to April 2022. After matching with their corresponding controls, 139 321 (60.3%) of 231 181 cases remained for analysis.

EXPOSURES

Two dose mRNA-1273, BNT162b2, and BBIBP-CorV vaccination schedule.

MAIN OUTCOME MEASURES

SARS-CoV-2 infection and covid-19 related death. Conditional logistic regression was used to estimate

WHAT IS ALREADY KNOWN ON THIS TOPIC

mRNA and inactivated covid-19 vaccines are effective in preventing severe disease and infection in children and adolescents in the short term after vaccination, but data on mortality are lacking

Waning of protection against infection has been described, especially for mRNA vaccines, but evidence for inactivated vaccines in children is limited

WHAT THIS STUDY ADDS

Vaccine effectiveness remained high for preventing mortality in children and adolescents aged 3-17 years, regardless of the predominant circulating variant Waning of protection against infection was described and evaluated for both types of vaccines

Heterologous mRNA schedules showed comparable to superior vaccine effectiveness in comparison with homologous schedules

the odds of SARS-CoV-2 infection among two dose vaccinated and unvaccinated participants. Vaccine effectiveness was estimated as (1–odds ratio)×100%.

RESULTS

Estimated vaccine effectiveness against SARS-CoV-2 infection was 61.2% (95% confidence interval 56.4% to 65.5%) in children and 66.8% (63.9% to 69.5%) in adolescents during the delta dominant period and 15.9% (13.2% to 18.6%) and 26.0% (23.2% to 28.8%), respectively, when omicron was dominant. Vaccine effectiveness declined over time, especially during the omicron period, from 37.6% (34.2% to 40.8%) at 15-30 days after vaccination to 2.0% (1.8%to 5.6%) after ≥ 60 days in children and from 55.8%(52.4% to 59.0%) to 12.4% (8.6% to 16.1%) in adolescents.

Vaccine effectiveness against death related to SARS-CoV-2 infection during omicron predominance was 66.9% (6.4% to 89.8%) in children and 97.6% (81.0% to 99.7%) in adolescents.

CONCLUSIONS

Vaccine effectiveness in preventing mortality remained high in children and adolescents regardless of the circulating variant. Vaccine effectiveness in preventing SARS-CoV-2 infection in the short term after vaccination was lower during omicron predominance and decreasing sharply over time.

TRIAL REGISTRATION

National Registry of Health Research IS003720.

Introduction

The worldwide vaccination strategy against covid-19 changed the evolution of the pandemic. The reduction of morbidity and mortality in adults, who were the first prioritized population given their high fatality rate, was remarkable. Unlike this population, infection in childhood and adolescence is usually mild, although severe cases with sequelae (for example, multisystem inflammatory syndrome) and even fatal outcomes have also been reported, especially in those with comorbidities.¹ In Argentina, the paediatric population was tested with molecular methods (for example, reverse transcription polymerase chain reaction (RT-PCR)) and immunochromatographic assays (for example, rapid antigen test) (fig 1, top) and represented about 8% of the total number of confirmed cases, with a case fatality rate of 0.05%.² Comparisons of effectiveness by diagnostic method for this population are not available.

Notwithstanding that vaccination in this age group has generated multiple debates, it was recommended by international scientific societies such as the American Academy of Pediatrics in May 2021 and Argentina's National Immunization Technical Advisory Group (NITAG).^{3 4} Moreover, in 2022, the World Health Organization determined that, despite the fact that it is not a priority group,⁵ progress should be made in the paediatric population in countries with high vaccination coverage in adults. Vaccination should aim not only for individual benefit but also for a collective benefit, as it would serve to increase overall population immunity and to reduce global transmission.

In Argentina, the vaccination strategy against covid-19 aimed to vaccinate 100% of the prioritised population in a gradual and progressive manner. The prioritisation was established according to the risk of severity of the disease, risk of exposure, and social vulnerability. Following the recommendations of the NITAG, vaccination of adolescents aged 12 to 17 years began in the second half of 2021, with the mRNA-1273 (Spikevax, Moderna) and BNT162b2 (Comirnaty, Pfizer-BioNTech) vaccines, followed by children aged 3 to 11 years who were vaccinated with BBIBP-CorV (Sinopharm) (fig 1, middle).⁶⁻⁹

Although a systematic review found that lowmiddle income countries may have larger proportion of paediatric fatality due to covid-19 than high income countries,¹⁰ evidence on the effectiveness of vaccination, especially to prevent mortality in



children and adolescents, is scarce.¹¹⁻¹⁴ Some studies evaluated vaccine effectiveness for inactivated vaccines, but only one included BBIBP-CorV and only two reported effectiveness in preventing infection in children under 5 years old.¹⁵⁻¹⁸ The study that evaluated BBIBP-CorV vaccine effectiveness in children was conducted in Buenos Aires Province (Argentina) and estimated that vaccine effectiveness in preventing hospital admission in 3-11 year old children was 76.4% (95% confidence interval 62.9% to 84.5%). However, this study did not evaluate effectiveness for preventing infection or mortality or loss of effectiveness over time.¹⁷

Moreover, although several studies reported a decline in effectiveness over time with the BNT162b2 vaccine in children or adolescents, scarce evidence exists of changes in effectiveness with inactivated vaccines or of differences related to SARS-CoV-2 variants. Furthermore, despite the fact that studies on the effectiveness of covid-19 vaccines usually include cases diagnosed with both RT-PCR and rapid antigen tests, comparisons of effectiveness by diagnostic method for the paediatric population are not available.

Therefore, the aim of this study was to evaluate the effectiveness of two dose schedules of mRNA-1273, BNT162b2, and BBIBP-CorV vaccines in preventing SARS-CoV-2 infection and death related to covid-19 in children and adolescents, aged 3-17 years, during periods of delta and omicron BA.1 predominance between September 2021 and April 2022 in Argentina. Additionally, we aimed to assess the effect of the diagnostic method used for estimating vaccine effectiveness, the effectiveness of different combinations of mRNA vaccines, and short term waning of immunity in children and adolescents.

Methods

Study population, setting, and design

This study followed a test negative, case-control design. These designs attempt to provide control over confounding due to differential case ascertainment, access to care, and health seeking behaviour. They are considered powerful enough to estimate the effectiveness of vaccines and have been widely used for this purpose in SARS-CoV-2, influenza, and other viruses.^{19 20}

Argentina started vaccination of adolescents (aged 12-17 years) in August 2021 and of young children (aged 3-11 years) in October 2021. All participants aged 3 to 17 years who were tested for SARS-CoV-2 between 12 September 2021 and 23 April 2022 (considering the delta predominance period as until 25 December 2021 and the omicron predominance period as since then) (fig 1, bottom), without a previous positive test since 1 June 2021, were eligible for inclusion (fig 2). We restricted analyses to participants who sought RT-PCR or antigen testing in healthcare facilities or testing centres. We included participants only once in the study period, whether as a case or as a control. The first positive test in the study period was considered

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inclusion as a case in the primary analysis, regardless of the number of previous negative tests. Controls were those participants who tested only negative in the study period and were selected on the date of their first test. To assess vaccine effectiveness against covid-19 associated mortality, we included as cases only those participants who tested positive in the short time before death. Additionally, the National Epidemiology Department evaluated each case with the jurisdictional epidemiology department to corroborate the cause of death.

Data sources and definitions

The study used epidemiological surveillance data from the National Surveillance System (SNVS 2.0), preserving the confidentiality of the participants according to the Helsinki declaration and local regulations. The notification of suspected cases of covid-19 and their confirmation and outcomes were done by certified users (professionals, technicians, administrative and health authorities of the 24 districts) of the public healthcare, private healthcare, and social security subsectors. For the death registry, each district systematically reviewed and verified data from other death records, such as bureaus of vital records (death certificate data), hospitals, and funeral companies. These data were incorporated into SNVS 2.0. Vaccine information was reported in the Nominalized Federal Vaccination Registry (NOMIVAC), including all vaccinated participants' date of vaccination, number of doses, type of vaccine, and vaccination centre. We followed the STROBE checklist.²¹

We considered 3-11 year old children to be fully vaccinated if they had received two doses of BBIBP-CorV and had received the second dose a minimum of 14 days before testing. We considered 12-17 year old adolescents to be fully vaccinated if they had received two doses of a homologous or heterologous schedule with BNT162b2 and/or mRNA-1273 and had received the second dose a minimum of 14 days before testing. Individuals who did not meet these criteria were included in the analyses in this article in three additional categories: those who had received only one dose a minimum of 14 days before testing (partially vaccinated), those who received three doses with the same minimum interval, and others. The third category includes those who were vaccinated in the two weeks before their test, those who received another vaccine, and those with inconsistent or incomplete vaccination data. Information about variants could not be established at an individual level, and we used the prevalent variant from surveillance data when analysing our results.

Outcomes

The primary outcome of this study was SARS-CoV-2 infection, confirmed by antigen or RT-PCR testing. The secondary outcome was death related to SARS-CoV-2 infection.

Statistical analyses

We matched cases and controls one to one on province of residence, week of testing, type of test (antigen or genomic), presence of comorbidities, sex, and age (±1 year). Ties during matching were automatically and randomly resolved by the algorithm. Matching improves statistical efficiency in case-control studies and yields unbiased estimates when a model accounting for matching, such as conditional logistic regression, is used. The effect of the matching factors on the outcome cannot be estimated in matched sets and thus is a considerable disadvantage of matching. To compare characteristics between cases and test negative controls, we did descriptive analyses for the





unmatched and matched cohorts. P values shown refer to Student's *t* test, Mann-Whitney U test, or Kruskal-Wallis rank sum test for differences in numerical variables or χ^2 tests for proportions.

We used conditional logistic regression to compare the odds of SARS-CoV-2 infection as a function of vaccination status, taking the matching pairs into consideration. We estimated vaccine effectiveness as (1–odds ratio)×100% and presented it with its corresponding 95% confidence interval by using the profile likelihood method. We estimated the same intervals by using bootstrapping as a robustness analysis. We created 1000 resampled datasets of matched case-control pairs. The total number of participants in each sample was equal to that of the original dataset (or the subset, for subgroup analysis), but differences within datasets were due to allowing replacement when sampling. For each of these datasets, we fitted a conditional logistic regression model and calculated vaccine effectiveness as in the main analysis. We calculated bootstrapping based vaccine effectiveness by taking the mean of the 1000 resampled datasets and defined its confidence interval limits as the empirical centiles 0.25 and 97.5.

We classified vaccination status in different levels depending on the subanalysis. The unvaccinated group was the selected reference group in all the analyses. Other levels of the vaccination status variable refer to participants who were partially vaccinated (at least 14 days between first dose and test), those who received three doses, and others (details in Data sources and definitions). The level of interest of this study is the fully vaccinated group with at least 14 days between their second dose and the diagnostic test.

		Test result			Vaccination status				
Characteristic	Overall (n=278 642)	Negative (n=139321)	Positive (n=139321)	P value	Unvaccinated (n=60 205)	Partially vaccinated (n=37941)	Fully vaccinated (n=150 966)	Boosted (n=3755)	Other* (n=25 775)
Any comorbidity:				1					
No	265 422 (95.3)	132711 (95.3)	132711 (95.3)		56162 (93.3)	36040 (95.0)	145 172 (96.2)	3706 (98.7)	24342 (94.4)
Yes	13220 (4.7)	6610 (4.7)	6610 (4.7)		4043 (6.7)	1901 (5.0)	5794 (3.8)	49 (1.3)	1433 (5.6)
Sex:				1					
Female	142 988 (51.3)	71494 (51.3)	71494 (51.3)		29 447 (48.9)	19022 (50.1)	79594 (52.7)	2078 (55.3)	12847 (49.8)
Male	135 654 (48.7)	67 827 (48.7)	67 827 (48.7)		30758 (51.1)	18919 (49.9)	71372 (47.3)	1677 (44.7)	12928 (50.2)
Type of test:		·		1					
RT-PCR	120 420 (43.2)	60210 (43.2)	60210 (43.2)		30 2 18 (50.2)	16 362 (43.1)	58 973 (39.1)	2637 (70.2)	12230 (47.4)
Antigen	158 222 (56.8)	79111 (56.8)	79111 (56.8)		29987 (49.8)	21 579 (56.9)	91 993 (60.9)	1118 (29.8)	13545 (52.6)
Mean (SD) week of diagnosis	17.4 (4.9)	17.4 (4.9)	17.4 (4.9)	1	14.3 (6.9)	17.5 (3.9)	18.7 (3.2)	22.6 (4.4)	16.5 (5.2)
Age group (years):				0.82					
3-11	145 705 (52.3)	72822(52.3)	72883 (52.3)		39 390 (65.4)	23460 (61.8)	66831 (44.3)	277 (7.4)	15747 (61.1)
12-17	132937 (47.7)	66 499 (47.7)	66 438 (47.7)		20815 (34.6)	14481 (38.2)	84 135 (55.7)	3478 (92.6)	10028 (38.9)
Time since second dose of vaccinet:	(n=150966)	(n=77 557)	(n=73409)	<0.001					
<60 days	72515 (48.0)	39951 (51.5)	32 564 (44.4)		-	-	72515 (48.0)	-	-
≥60 days	78 451 (52.0)	37 606 (48.5)	40845 (55.6)		-	-	78451 (52.0)	-	-

Table 1 | Population characteristics after matching, by test result and vaccination status. Values are numbers (percentages) unless stated otherwise

RT-PCR=reverse transcription polymerase chain reaction; SD=standard deviation.

*Includes participants who were vaccinated in the two weeks before their test, those who received another vaccine, and those with inconsistent or incomplete vaccination data. tin fully vaccinated participants.

We did several subanalyses. Firstly, we split data according to the study period (on 25 December 2021) considering the change in the dominant variant (from delta to omicron). In the last week of the delta period, 68.2% of cases were identified as due to delta and 31.7% as due to omicron. During the first week of the omicron period, 79.2% of the studied cases were assigned to this variant.²² For a second subanalysis, we used another split in age subgroups (3-11 and 12-17 years).

Seeking evidence of waning immunity, in a third subanalysis we divided the fully vaccinated group into four levels according to the time between their second dose and the test (15-30, 31-45, 46-60, \geq 61 days). For a fourth subanalysis, we split the same level to assess vaccine effectiveness by product in the adolescent subpopulation, with all possible combinations of BNT162b2 and mRNA-1273 vaccines against both variants.

Fifthly, we did three sensitivity analyses. We assessed the effect of the type of test in the estimation by doing a subanalysis estimating RT-PCR and antigen based vaccine effectiveness separately. We fitted a linear regression model and used the non-parametric Mann-Whitney-Wilcoxon test to assess whether the difference between estimations was different from zero.

For the linear model, PCR based estimates were the predictor variable and antigen based estimates were the dependent variable, as most test negative, casecontrol studies use PCR tests to assign participants a case status. We then repeated the main analysis excluding matching pairs in epidemiological weeks with zero fully vaccinated, to assess how not holding the positivity assumption might bias our estimates. Lastly, we excluded participants who had a previous infection to assess whether these participants might be biasing our estimates.

Finally, for a fatality analysis, we matched participants with covid-19 associated death one to four to covid-19 negative participants by using the same matching criteria as the one used in the primary analysis estimating vaccine effectiveness against SARS-CoV-2 infection. On this second matched casecontrol group, we estimated vaccine effectiveness against mortality through an adjusted conditional logistic regression model.

Data preprocessing was carried out with PostgreSQL (PostgreSQL Global Development Group). We used R software (R Development Core Team, version 3.6.1) for all statistical analyses.

Table 2 | Vaccination schedule and vaccine effectiveness against infection, by combination and period. Values are numbers (percentages) unless stated otherwise

		3-11 vears old	12-17 vears old	Vaccine effectiveness (95% CI)		
	Overall (n=150 966)	(n=59858)	(n=91108)	Delta period	Omicron period	
Cases	73409 (48.6)	29 200 (48.8)	44 209 (48.5)	-	-	
Vaccination schedule						
mRNA-1273 and mRNA-1273	22141 (14.7)	0 (0.0)	22141 (24.3)	70.2 (66.8 to 73.1)	17.9 (14.0 to 21.5)	
mRNA-1273 and BNT162b2	4197 (2.8)	0 (0.0)	4197 (4.6)	66.3 (54.0 to 75.4)	31.5 (26.3 to 36.4)	
BNT162b2 and mRNA-1273	573 (0.4)	0 (0.0)	573 (0.6)	88.9 (66.1 to 96.4)	40.6 (29.4 to 50.0)	
BNT162b2 and BNT162b2	64197 (42.5)	0 (0.0)	64 197 (70.5)	64.1 (60.5 to 67.3)	28.1 (25.2 to 30.8)	
BBIBP-CorV and BBIBP-CorV	59858 (39.6)	59858 (100.0)	0 (0.0)	61.2 (56.4 to 65.6)	16.0 (13.2 to 18.6)	
Cl. and fide and interval						

CI=confidence interval.

Study	Vaccine effectiveness, (95% Cl)		Vaccine e (eness, %)	:	
Delta period						
All	64.2 (61.6 to 66.5)					
3-11 years	61.2 (56.4 to 65.5)					
12-17 years	66.8 (63.9 to 69.5)					
Omicron per	riod					
All	19.9 (18.0 to 21.8)		•			
3-11 years	15.9 (13.2 to 18.6)					
12-17 years	26.0 (23.2 to 28.8)					
		0	20	40	60	80

Fig 3 | Vaccine effectiveness of two dose schedules against SARS-CoV-2 infection, by age group and study period. CI=confidence interval

Public and patient involvement

This was an unfunded study using routine surveillance data sources. No patients were involved in setting the research question or the outcome measures, nor were they involved in developing plans for design or implementation of the study. No patients were asked to advise on interpretation or writing up of results. There are no plans to directly disseminate the results of the research to study participants.

Results

The initial case-control cohort included 844460 participants, of whom 421359 sought RT-PCR testing and 423101 were tested with immunoassays for antigen detection, between 12 September 2021 and 23 April 2022. Those who tested positive (231181) had a higher proportion of reported comorbidities (5.2% v 4.9%; P<0.001), were less likely to be male (47.4% v 50.7%; P<0.001), had sought to be tested in more advanced epidemiological weeks and were more frequently tested with antigen detection (65.4% v 44.3%; P<0.001) (supplementary table A). Furthermore, they were slightly older (mean age 11.9 v 10.8 years; P<0.001) and had significant differences in the regional distribution (supplementary table B).

After matching, 139321 (60.3%) of 231181 cases with their corresponding controls remained for analysis. As a result of exact matching on presence of comorbidities, sex, type of test, province (supplementary table C), and week of testing, the distribution of those variables in cases and controls are equal. Age was matched with a tolerance of 1 year, yielding non-significant differences between the two groups (table 1).

The fully vaccinated participants represented 54% (150966/278642) of the matched case study group. Of these fully vaccinated participants, 73 409 (48.6%) were included as cases and the remainder as controls. In the 12-17 year old fully vaccinated subgroup, 86338/91108 (94.8%) received a homologous two dose schedule (64197 (70.5%) BNT162b2; 22141 (24.3%) mRNA-1273) and 4770 (5.2%) received a heterologous one (4197 (4.6%) mRNA-1273 and then BNT162b2; 573 (0.6%) BNT162b2 and then mRNA-1273). All children (3-11 years old) included as fully vaccinated received a two dose schedule of BBIBP-CorV (table 2). The median time from second dose to test was 66 (interguartile range 40) days for the 12-17 year old subgroup and 54 (32) days for the 3-11 year old subgroup.

Primary analysis on vaccine effectiveness in preventing infection

Figure 3 shows vaccine effectiveness in preventing infection according to the predominant variant and age group. Vaccine effectiveness was considerably higher in all age groups when we analysed only the period of delta variant predominance. Vaccine effectiveness was 64.2% (95% confidence interval 61.6% to 66.5%) for all ages, 61.2% (56.4% to 65.5%) for children (3-11 years), and 66.8% (63.9% to 69.5%) for adolescents (12-17 years). Estimates of vaccine effectiveness decreased markedly during the period of omicron predominance. Vaccine effectiveness was 19.9% (18.0% to 21.8%) for all ages, with a slightly higher value in adolescents who received mRNA based



Days since second dose

Fig 4 | Vaccine effectiveness (with 95% confidence interval) of two dose schemes against SARS-CoV-2 infection, by age group, study period, and days since second dose

		Children (3-11 years old)		Adolescents (12-17 years old)		
Characteristic	Overall (n=220)	Controls (n=96)	Covid-19 death (n=24)	Controls (n=80)	Covid-19 death (n=20)	
Sex:						
Female	135 (61)	52 (54)	13 (54)	56 (70)	14 (70)	
Male	85 (39)	44 (46)	11 (46)	24 (30)	6 (30)	
Type of test:						
RT-PCR	155 (70)	84 (88)	21 (88)	40 (50)	10 (50)	
Antigen	65 (30)	12 (12)	3 (12)	40 (50)	10 (50)	
Vaccination status:						
Unvaccinated	66 (30)	34 (35)	15 (63)	6 (8)	11 (55)	
Partially vaccinated	26 (12)	10 (10)	2 (8)	9 (11)	5 (25)	
Fully vaccinated	109 (50)	41 (43)	6 (25)	60 (75)	2 (10)	
Boosted	2 (1)	0 (0)	0 (0)	2 (3)	0 (0)	
Other*	17 (8)	11 (11)	1 (4)	3 (4)	2 (10)	
Period:						
Delta	25 (11)	20 (21)	5 (21)	0 (0)	0 (0)	
Omicron	195 (89)	76 (79)	19 (79)	80 (100)	20 (100)	
Vaccine effectiveness against omicron related death (95% CI)	88.1 (70.7 to 95.2)	66.9 (6.4 to 89.8)		97.6 (81.0 to 99.7)	

Table 3 | Characteristics of covid-19 related deaths and matched controls. Values are numbers (percentages) unless stated otherwise

CI=confidence interval; RT-PCR=reverse transcription polymerase chain reaction.

*Includes participants who were vaccinated in the two weeks before their test, those who received another vaccine, and those with inconsistent or incomplete vaccination data.

vaccines (26.0%, 23.2% to 28.8%) than in children who received BBIBP-CorV (15.9%, 13.2% to 18.6%). Assessment of robustness by means of bootstrapping yielded almost identical results with narrower confidence intervals (supplementary table D).

Waning immunity

As shown in figure 4, during the delta period, vaccine effectiveness declined according to time since vaccination from 68.4% (64.1% to 72.2%) at 15-30 days to 65.2% (44.0% to 78.4%) at ≥ 61 days in children and from 74.8% (71.3% to 77.9%) to 56.3% (50.2% to 61.7%) in adolescents (supplementary table E). During the omicron period, the initial effectiveness was lower and the decline over time was steeper. Vaccine effectiveness dropped from 37.6% (34.2% to 40.8%) to 2.0% (1.8% to 5.6%) in children and from 55.8% (52.4% to 59.0%) to 12.4% (8.6% to 16.1%) in the adolescent subpopulation.

Subanalysis by vaccine type (adolescents)

As shown in table 2, all combinations of mRNA vaccine products showed similar results in adolescents. During the delta period, vaccine effectiveness was 70.2% (66.8% to 73.1%) for the homologous schedule of mRNA-1273 and 64.1% (60.5% to 67.3%) for BNT162b2. The heterologous schedule of mRNA-1273 then BNT162b2 yielded a vaccine effectiveness of 66.3% (54.0% to 75.4%), and that for BNT162b2 then mRNA-1273 was 88.9% (66.1% to 96.4%). In the omicron period, homologous schedules of mRNA-1273 and BNT162b2 showed a vaccine effectiveness of 17.9% (14.0% to 21.5%) and 28.1% (25.2% to 30.8%), respectively. Heterologous schedules showed 40.6% (29.4% to 50.0%) for participants receiving the BNT162b2 vaccine first and 31.5% (26.3% to 36.4%) for those who received the mRNA-1273 vaccine first.

Sensitivity analyses

On the hypothesis that estimations might be affected by the type of test used to select cases and controls, we analysed the results for RT-PCR and antigen test based vaccine effectiveness estimates (supplementary table F). The slope for the linear regression between RT-PCR and antigen based estimates of vaccine effectiveness was 0.933 (95% confidence interval 0.892 to 0.974), showing that antigen based vaccine effectiveness estimates tend to be slightly lower than RT-PCR based estimates, rejecting the null hypothesis that the slope equals one (P=0.004) (supplementary figure A). Additionally, we did a Mann-Whitney-Wilcoxon test to assess the relative width of the confidence interval between the two techniques. It yielded a nonsignificant P value of 0.61.

To evaluate how not holding the positivity assumption could affect our estimations, we did a sensitivity analysis excluding matching pairs in epidemiological weeks with zero fully vaccinated participants for each age subgroup. This occurred only during the delta period, and for both age groups the confidence intervals had a wide overlap with the results reported in the main analysis (supplementary table G).

Next, we did a sensitivity analysis excluding participants with previous infection to evaluate whether our estimates might be biased due to previously acquired immunity. After exclusion of participants who fulfilled this criterion, vaccine effectiveness estimates had a wide overlap with the results reported in the main analysis (supplementary table H).

Vaccine effectiveness in preventing mortality

Fifty one deaths related to covid-19 were notified among cases, of which seven occurred in the delta period (all in unvaccinated participants). Of the 51, 30 were unvaccinated participants (16 children; 14 adolescents), eight were partially vaccinated (three children; five adolescents), nine were fully vaccinated (six children; three adolescents), and four had other status (two children; two adolescents). Two further deaths were excluded after validation that the cause of death was not related to covid-19. Following one-tofour case-control matching, 44 (90%) of 49 deceased participants were successfully matched (table 3). After adjusting a conditional logistic regression model, we found that vaccine effectiveness against mortality in the 3-17 year old group for the two dose schedule was 89.3% (73.9% to 95.6%). When we considered only the omicron period, vaccine effectiveness remained high at 88.1% (70.7% to 95.2%). Considering age subgroups, vaccine effectiveness against death related to SARS-CoV-2 infection during omicron predominance was 97.6% (81.0% to 99.7%) for adolescents and 66.9% (6.4% to 89.8%) for children.

Discussion

This study provides real world evidence for the effectiveness of mRNA-1273, BNT162b2, and BBIBP-CorV vaccines in preventing infection and mortality in children and adolescents in Argentina. To our knowledge, this is one of the first studies that reports vaccine effectiveness in children under 5 years old, in addition to evaluation of mortality and analysis of waning for BBIP-CoV and mRNA-1273 vaccines. Our results suggest that the primary vaccination schedule is effective in preventing mortality in children and adolescents with covid-19 regardless of the circulating SARS-CoV-2 variant. Vaccine effectiveness in preventing mortality in children vaccinated with BBIP-CoV was lower than in adolescents vaccinated with mRNA vaccines; however, owing to the scarcity of events, analysis of mortality by age group yielded results with wide confidence intervals. Our estimates also suggest that two doses of vaccine are effective in preventing SARS-CoV-2 infection in children and adolescents in the short term. Vaccine effectiveness was considerably higher when delta was the predominant circulating variant, but a significant decrease was observed over time, especially during omicron predominance.

Comparison with other studies

By the time of our study, only BBIBP-CorV was indicated in children, so we could not compare vaccine effectiveness in children by vaccine type. Evidence is variable taking into consideration that published studies differ in age groups and time to vaccination periods. Nevertheless, BNT162b2 vaccine effectiveness in preventing infection during omicron predominance has been evaluated, with vaccine effectiveness between 29.4% and 51% being reported.^{12 23 24} These results are similar to the vaccine effectiveness reported in our study and in other studies that evaluated inactivated vaccines (CoranaVac) and reported vaccine effectiveness of 39.8% and 38.2%.^{16 18}

In adolescents, we evaluated different vaccine schedules. Compared with the delta period, vaccine

effectiveness during the omicron period was lower for both evaluated schedules (homologous and heterologous), in consistency with the literature. A previous study of immunogenicity in adults in Argentina evaluated heterologous schedules of adenoviral vectored, inactivated, and mRNA vaccines and endorsed heterologous vaccination strategies.²⁵ In our study, which evaluated mRNA-1273 and BNT162b2 vaccines, heterologous schedules showed a comparable to superior vaccine effectiveness in comparison with homologous schedules. Nevertheless, further investigation may be needed to evaluate different vaccine schedules.

In addition, our results are consistent with other published studies that found high short term protection induced by the primary vaccination regimen against SARS-CoV-2 infection during the delta predominance period.²⁶⁻²⁸ However, since the emergence and spread of the omicron variant of concern, a reduction in vaccine effectiveness in preventing infection has been observed in adults and children.^{13 29-31} A population based study in Norway reported that one and two doses of mRNA vaccine (BNT162b2) in adolescents protected against infection with the delta variant, reaching an effectiveness of 91% after the primary vaccination schedule and declining to 53% for the omicron variant.²⁸ Fowlkes and colleagues described a decrease in vaccine effectiveness against infection in adolescents for the primary mRNA vaccine series (BNT162b2) from 87% to 59%, comparing delta and omicron variants.¹² In Chile, Jara and colleagues reported vaccine effectiveness in preventing infection for a complete schedule of inactivated vaccine (Sinovac's CoronaVac) in 6-11 year old children of 75.8% during the delta circulating period. In another study, Jara and colleagues reported a lower vaccine effectiveness of 38.2% during omicron circulation for the same vaccine in 3-5 year old children.^{15 16} Vaccine effectiveness reported by Jara and colleagues is similar to that observed with BBIBP-CorV in children in our study.

Available evidence suggests that the vaccine induced immune response could decrease over time, with a reduction in the effectiveness of SARS-CoV-2 vaccines in preventing infection, particularly in the context of the emergence of new variants of the virus with potential ability to partially evade the host immune response.^{32 33} Our results highlight not only a decrease in effectiveness when comparing delta with omicron but also, and especially during omicron predominance, a significant decrease with time since vaccination. The decrease observed with BBIBP-CorV in children in our study is consistent with published evidence. Fleming-Dutra and colleagues did a test negative, casecontrol analysis to evaluate the vaccine effectiveness of BNT162b2 in preventing infection among children and adolescents during omicron variant predominance and reported that two months after the second dose the estimated vaccine effectiveness had declined in both age groups.³⁴ Similarly, Dorabawila and colleagues concluded that the risk of infection

and hospital admission was higher in unvaccinated children and adolescents, although the protection declined over time from vaccination.³⁰ Sacco and colleagues reported a decrease in vaccine effectiveness for BNT162b2 in preventing infection over time, and Klein and colleagues found no evidence of protection for adolescents aged 12-17 years after receipt of two doses of BNT162b2 ≥150 days earlier during omicron predominance.^{23 29} In a Norwegian study, protection against omicron after a complete primary regimen in adolescents was reduced to 23% from 63 days post-vaccination, whereas a smaller reduction was documented for delta.²⁸ Amir and colleagues observed a rapid reduction in protection against omicron of two doses of mRNA vaccine (BNT162b2) in adolescents aged 12-15, with an increase in the rate of confirmed infections from 60 days after the second dose.³⁵ Evidence of a decrease in vaccine effectiveness after inactivated vaccines is lacking.

Although the previously cited studies suggest a reduction in vaccine effectiveness for SARS-Cov-2 infection over time, observational studies have shown that vaccine effectiveness against severe disease is higher and more stable. Data from Brazil and Scotland in adolescents aged 12-17 years during the omicron wave showed a reduction in BNT162b2 vaccine effectiveness for symptomatic infection from 98 days after receipt of the primary schedule. However, protection against hospital admission and death with covid-19 remained above 80% in the same time interval.³⁶ In Canada, vaccine effectiveness for two doses of BNT162b2 against severe disease with omicron in the same age group was 85%, with a stable trend over time.³⁷ In children, Tan and colleagues reported a vaccine effectiveness of 82.7% in preventing hospital admission for BNT162b2.38 For inactivated vaccines, vaccine effectiveness of 59.2% and 64.6% have been reported for prevention of hospital admission with Sinovac's CoronaVac.^{16 18}

Our results are consistent with a study conducted in Buenos Aires Province (Argentina) that reported a two dose vaccine effectiveness of 78.0% for hospital admission in 3-17 year olds evaluated during delta/ omicron circulation.¹⁷ As expected, we found that vaccine effectiveness for infection was lower and vaccine effectiveness for mortality was higher than these estimations for protection against hospital admissions, especially in adolescents. In this study, estimated effectiveness against hospital admissions was higher when delta was predominant, although it remained higher than 65% during the omicron period.¹⁷ This study, which took place in an Argentinian province, was the first one to evaluate vaccine effectiveness of BBIBP-CorV in children between 3 and 11 years old. However, it did not evaluate vaccine effectiveness in preventing infection or mortality or decrease in vaccine effectiveness over time.17

Our study evaluated vaccine effectiveness in children and adolescents, including different types of diagnostic tests. Although statistically significant, the real life implications of these differences might not be important for public health agents. According to our results, antigen based estimates tended to be 6.7% lower (P=0.004) than the PCR based one for the same population and provide results with a comparable variance. Therefore, the advantages of using antigen tests alone or in addition to RT-PCR tests in vaccine effectiveness studies would outweigh the slight differences in estimations. Tan and colleagues reported differences in vaccine effectiveness by type of diagnostic test; however, they highlighted that in disease management protocols, the country included PCR testing for people with more severe symptoms or coexisting conditions, so they used PCR results as proxy of increased severity of illness.³⁸

Strengths and limitations of study

The study has several strengths. Firstly, the sample size was large and included all children and adolescents tested and reported to the National Surveillance System in Argentina during the study period. Secondly, the high quality of the epidemiological surveillance (especially for laboratory and mortality data) and vaccination records used confers robustness to the results.

Limitations of our study include that some information, such as symptoms and hospital admissions, was incomplete and was consequently not included in any analysis. Also, as persistent potential confounders, misclassification, or selection bias might have been present, we decided to minimise them by matching with strict criteria on all the available confounders and doing univariate conditional logistic regression analyses. We also did several sensitivity analyses to assess the effect of some unmeasured phenomena. Nevertheless, unobserved confounding might exist and bias odds ratio estimations. The generalisability of our results is limited to the subpopulation who had access to testing. Additionally, as misclassification might have occurred as a result of non-100% specificity and sensitivity of covid-19 tests, we included a sensitivity analysis to assess how test related problems might affect the vaccine effectiveness estimations, which showed that antigen based estimates of vaccine effectiveness tended to be lower. Other limitations include that each age group used different vaccines, so any conclusion based on their differences should not be attributed exclusively to the vaccine technology or to age related factors. Finally, information about variants could not be established at an individual level, and we used the prevalent variant from surveillance data when analysing our results.

Conclusions

Vaccine effectiveness remained high for preventing mortality in children and adolescents regardless of the predominant circulating variant. Vaccines were effective in preventing SARS-CoV-2 infection in the short term after vaccination, being lower when the omicron variant was predominant and decreasing sharply over time. Heterologous mRNA schedules showed comparable to superior vaccine effectiveness in

comparison with homologous schedules. In summary, vaccinating children is an important public health measure that will prevent mortality in this population, especially in periods of high viral circulation.

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Contributors: AR, CV, and JMC conceived and supervised the study. AR was in charge of the project administration. AR and SO designed the methods. SO performed and interpreted the formal analysis. AR, SO, CV, MDVJ, MP, MEC, MP, and ANI conceptualised the study and wrote the original draft. CMG critically revised the manuscript. MPB, MBM, and ML developed the supporting algorithms and curated the data. All authors reviewed, edited and gave final approval for the version to be submitted. AR is the guarantor. The corresponding author attests that all listed authors meet authorship criteria and that no others meeting the criteria have been omitted.

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Competing interests: All authors have completed the ICMJE uniform disclosure form at https://www.icmje.org/disclosure-of-interest/ and declare: no support from any organisation for the submitted work; JMC and CV were involved in the decision making process of the vaccination campaign in Argentina; no other relationships or activities that could appear to have influenced the submitted work.

Ethical approval: Covid-19 surveillance was included as an obligatory notifiable disease by RM 680/2020 (https://www. boletinoficial.gob.ar/detalleAviso/primera/227324/20200331). As secondary data were used, the study protocol was subject to an internal review by the ethical area of the Research Department of the Ministry of Health and was found to be fully compliant with all regulatory requirements. As no regulatory issues were identified, a full ethical review was deemed not to be necessary.

Data sharing: The data that support the findings of this study will be available for researchers who provide a methodologically sound proposal after it is approved, on request from the corresponding author.

The lead author (the manuscript's guarantor) affirms that the manuscript is an honest, accurate, and transparent account of the study being reported; that no important aspects of the study have been omitted; and that any discrepancies from the study as planned (and, if relevant, registered) have been explained.

Dissemination to participants and related patient and public communities: Results from this research will be shared through conferences with expert immunisation advisory committees for Argentina and public health officials overseeing Argentina's covid-19 immunisation campaign. The findings will also be further disseminated through the National Health Ministry website, social media channels, and news media.

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Web appendix: Supplementary materials