

Incidence determinants and serological correlates of reactive symptoms following SARS-CoV-2 vaccination

Holt, H., Jolliffe, D. A., Talaei, M., Faustini, S., Vivaldi, G., Greenig, M., Richter, A. G., Lyons, R. A., Griffiths, C. J., Kee, F., Sheikh, A., Davies, G. A., Shaheen, S. O., & Martineau, A. R. (2023). Incidence determinants and serological correlates of reactive symptoms following SARS-CoV-2 vaccination. *NPJ vaccines, 8*, Article 26. Advance online publication. https://doi.org/10.1038/s41541-023-00614-0

Published in:

NPJ vaccines

Document Version: Publisher's PDF, also known as Version of record

Queen's University Belfast - Research Portal:

Link to publication record in Queen's University Belfast Research Portal

Publisher rights

Copyright 2023 the authors.

This is an open access article published under a Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution and reproduction in any medium, provided the author and source are cited.

General rights

Copyright for the publications made accessible via the Queen's University Belfast Research Portal is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy

The Research Portal is Queen's institutional repository that provides access to Queen's research output. Every effort has been made to ensure that content in the Research Portal does not infringe any person's rights, or applicable UK laws. If you discover content in the Research Portal that you believe breaches copyright or violates any law, please contact openaccess@qub.ac.uk.

Open Access

This research has been made openly available by Queen's academics and its Open Research team. We would love to hear how access to this research benefits you. – Share your feedback with us: http://go.qub.ac.uk/oa-feedback

ARTICLE OPEN

Check for updates

Incidence determinants and serological correlates of reactive symptoms following SARS-CoV-2 vaccination

Hayley Holt $1^{2,3}$, David A. Jolliffe^{1,3}, Mohammad Talaei 1^{1} , Sian Faustini⁴, Giulia Vivaldi $1^{2,3}$, Matthew Greenig¹, Alex G. Richter⁴, Ronan A. Lyons 5^{5} , Christopher J. Griffiths^{1,3}, Frank Kee⁶, Aziz Sheikh⁷, Gwyneth A. Davies⁵, Seif O. Shaheen¹ and Adrian R. Martineau^{1,2,3}

Prospective population-based studies investigating associations between reactive symptoms following SARS-CoV-2 vaccination and serologic responses to vaccination are lacking. We therefore conducted a study in 9003 adults from the UK general population receiving SARS-CoV-2 vaccines as part of the national vaccination programme. Titres of combined IgG/IgA/IgM responses to SARS-CoV-2 spike (S) glycoprotein were determined in eluates of dried blood spots collected from all participants before and after vaccination, 4262 (47.3%) participants experienced systemic reactive symptoms after a first vaccine dose. Factors associating with lower risk of such symptoms included older age (aOR per additional 10 years of age 0.85, 95% CI: 0.81–0.90), male vs. female sex (0.59, 0.53-0.65) and receipt of an mRNA vaccine vs. ChAdOx1 nCoV-19 (0.29, 0.26-0.32 for BNT162b2; 0.06, 0.01-0.26 for mRNA-1273). Higher risk of such symptoms was associated with SARS-CoV-2 seropositivity and COVID-19 symptoms prior to vaccination (2.23, 1.78–2.81), but not with SARS-CoV-2 seropositivity in the absence of COVID-19 symptoms (0.94, 0.81–1.09). Presence vs. absence of self-reported anxiety or depression at enrolment associated with higher risk of such symptoms (1.24, 1.12–1.39). Postvaccination anti-S titres were higher among participants who experienced reactive symptoms after vaccination vs. those who did not (P < 0.001). We conclude that factors influencing risk of systemic symptoms after SARS-CoV-2 vaccination include demographic characteristics, pre-vaccination SARS-CoV-2 serostatus and vaccine type. Participants experiencing reactive symptoms following SARS-CoV-2 vaccination had higher post-vaccination titres of IgG/A/M anti-S antibodies. Improved public understanding of the frequency of reactogenic symptoms and their positive association with vaccine immunogenicity could potentially increase vaccine uptake.

npj Vaccines (2023)8:26; https://doi.org/10.1038/s41541-023-00614-0

INTRODUCTION

The COVID-19 pandemic is the greatest threat in a generation to global health, having caused more than 6.1 million deaths to date¹. SARS-CoV-2 vaccination represents the mainstay of disease control, but uptake is suboptimal, largely owing to vaccine hesitancy². Fear of experiencing post-vaccination reactogenic symptoms - the physical manifestation of the inflammatory response to vaccination³—is often a contributing factor to such hesitancy⁴. Investigating the frequency with which post-vaccination reactogenic symptoms are experienced, risk factors for experiencing them and associations with vaccine immunogenicity has potential to yield information that will allow potential vaccinees to assess their risk of experiencing such symptoms, and to understand the significance of their presence or absence.

Existing studies investigating reactogenicity to first and second doses of SARS-CoV-2 vaccination have reported that post-vaccination symptoms are more commonly reported by women vs. men, in younger vs. older people, and in those who have had previous SARS-CoV-2 infection vs. those who have not⁵⁻⁹. However, the range of potential determinants explored by these studies has been limited: none have explored the potential impact of pre-vaccination anxiety or depression, specific comorbidities, medications or socio-economic factors which may either directly

influence risk of reactogenic symptoms, or confound associations with age and sex. Additionally, there is controversy regarding the relationship between post-vaccination symptoms from a full course of vaccination and SARS-CoV-2 vaccine immunogenicity, with some studies showing no relationship between post-vaccination symptoms and titres of antibodies to the SARS-CoV-2 spike (S) glycoprotein^{7,10}, one reporting a positive association overall⁸, and another reporting a positive association in men, but not in women¹¹. These studies were relatively small and were not population based, limiting both power and generalisability of their findings.

To address these limitations, we conducted a large, populationbased study in United Kingdom (UK) adults receiving SARS-CoV-2 vaccines, capturing detailed information on multiple potential determinants of reactogenic symptoms, and assaying combined IgG, IgA and IgM responses to SARS-CoV-2 spike protein before and after vaccination.

RESULTS

Participant flow and characteristics

Of the 17,796 participants who completed the COVIDENCE UK baseline questionnaire by September 2021, we excluded those

¹Wolfson Institute of Population Health, Barts and The London School of Medicine and Dentistry, Queen Mary University of London, UK. ²Bizard Institute, Barts and The London School of Medicine and Dentistry, Queen Mary University of London, London, UK. ³Asthma UK Centre for Applied Research, Queen Mary University of London, London, UK. ⁴Institute of Immunology and Immunotherapy, College of Medical and Dental Sciences, University of Birmingham, Birmingham, UK. ⁵Population Data Science, Swansea University Medical School, Singleton Park, Swansea, UK. ⁶Centre for Public Health Research (NI), Queen's University Belfast, Belfast, UK. ⁷Usher Institute, University of Edinburgh, Edinburgh, UK. ^{Sema}





who had no pre-vaccination serology (n = 6343) or who had not completed a full vaccination regimen (n = 2273). We also excluded those with missing data for independent variables required for minimal adjustment in the regression models

	Characteristic	N (%)
Age, years	16–29.99	120 (1.3)
	30–39.99	341 (3.8)
	40–49.99	800 (8.9)
	50–59.99	2192 (24.4)
	60–69.99	3617 (40.2)
	≥70.00	1933 (21.5)
Sex	Female	6367 (70.7)
	Male	2363 (29.3)
Ethnicity	White	8676 (96.4)
	Mixed/multiple/other ethnic groups	197 (2.2)
	Asian/Asian British	94 (1.0)
	Black/African/Caribbean/Black British	36 (0.4)
Country of residence	England	7961 (88.4)
	Northern Ireland	139 (1.5)
	Scotland	563 (6.3)
	Wales	340 (3.8)
Highest educational level	Primary/Secondary	978 (10.9)
attained	Higher/further (A levels)	1265 (14.1)
	College	3991 (44.4)
	Post-graduate	2762 (30.7)
Body mass index ^a	<25, kg/m ²	4369 (48.6)
	25–30, kg/m ²	2907 (32.4)
	>30, kg/m ²	1709 (19.0)
Self-reported general health	Excellent	1878 (20.9)
	Very good	3590 (39.88
	Good	2326 (25.8)
	Fair	949 (10.5)
	Poor	260 (2.9)
Tobacco smoking status	Not a current smoker	8633 (95.9)
	Current smoker	370 (4.1)
E-cigarette status ^b	Not a current vaper	8778 (97.7)
	Current smoker	207 (2.3))
Alcohol consumption	None	2349 (26.1)
	1–7 units	3189 (35.4)
	8–14 units	1853 (20.6)
	15–21 units	901 (10.0)
	22–28 units	403 (4.5)
	>28 units	308 (3.4)
/accine type	2 doses ChAdOx1	5988 (66.5)
	2 doses BNT162b2	2864 (31.8)
	2 doses MRNA-1273	59 (0.66)
	2 doses Other ^c	92 (1.02)
Nonth of dose for first dose	O1 (Jan–Mar)	8262 (91.8)
	Q2 (Apr-Jun)	625 (6.9)
	O3 (Jul-Sep)	1 (0.01)
	04 (Oct-Dec)	115 (1.3)
Month of dose for	O1 (Jan-Mar)	751 (8 3)
second dose	O2 (Apr-lup)	7975 (88 6)
		274 (2.0)
	$Q_{2} (D_{\alpha} = -2ep)$	2/7 (3.0)
	24 (UCI-Dec)	5 (0.03)

Vape status missing for 18 participants.

^cSingle dose Janssen (n = 12), 2 doses Novavax (n = 6), 2 doses Valneva (n = 3), Comcov2 trial (n = 1), Not sure/do not know.

(n = 177), resulting in 9003 participants being eligible for this analysis (Supplementary Fig. 1, Supplementary Material). Selected baseline characteristics of participants included in the analysis are presented in Table 1. The mean age of those contributing data to analyses was 61.3 years (range: 16.6-88.0), 70.7% were female, 96.4% identified their ethnic origin as White, and 88.4% were living in England. A total of 6322 (70.2%) participants experienced at least one post-vaccination symptom. The mean number of days from the date of second vaccine dose to date of dried blood spot was 58.2 (s.d. 22.4). This differed slightly for participants who did vs. did not report systemic symptoms after vaccination (56.2 vs. 61.2 days, P < 0.001), but was no different for those who did vs. did not report local symptoms after vaccination (57.9 vs. 58.6 days, P = 0.23).

Incidence of reactive symptoms by dose order and vaccine type

Figure 1 presents the proportions of participants reporting systemic and local symptoms by dose order and vaccine type. Systemic symptoms following first dose were most commonly reported by participants who received ChAdOx1 (57.1%), followed by BNT162b2 (28.3%) and mRNA-1273 (3.4%; P < 0.001). Following the second dose, the proportion of participants experiencing systemic symptoms was similar for ChAdOx1 (27.9%) vs BNT162b2 (28.9%), but lower for mRNA-1273 (8.5%; P = 0.002). Local symptoms following first dose were most common for BNT162b2 (51.3%), followed by ChAdOx1 (40.3%) and mRNA-1273 (10.2%; P < 0.001). Local symptoms after the second dose were reported less frequently than after the first dose, with a similar distribution according to vaccine type as seen with the first dose (BNT162b2 43.7% vs. ChAdOx1 27.4% vs. mRNA-1273 8.5%; P < 0.001).

Determinants of systemic reactive symptoms following first vaccine dose

After adjustment for age and sex only, 23 variables were found to associate with risk of experiencing systemic symptoms following a first vaccine dose with P < 0.10 (Table 2), and 14 did not (Supplementary Table 3, Supplementary Material). After inclusion of the former group of factors in a multivariable model, the following variables were independently associated with higher risk



Fig. 1 Proportions of participants experiencing systemic or local reactive symptoms after receiving first and second doses of ChAdOx1, BNT162b2, and mRNA-1273 vaccines. Panel a shows systemic symptoms are more common with ChAdOx1 vaccine following first dose. Panel b shows local symptoms are more common with BNT162b2 vaccine following second dose.

r	J	ľ)	j
			1	3

			Minimally adjusted		Fully adjusted	
		N (%) symptomatic	aOR (95% CI)	Р	aOR (95% CI)	Р
Vaccine type	ChAdOx1	3420/5988 (57.0)	1.00		1.00	
	BNT162b2	810/2864 (28.3)	0.31 (0.28–0.34)	<0.001	0.29 (0.26–0.32)	<0.00
	MRNA-1273	2/59 (3.4)	0.03 (0.01-0.11)	< 0.001	0.06 (0.01–0.26)	< 0.00
	Other	30/92 (43.6)	0.34 (0.22–0.54)	< 0.001	0.39 (0.4–0.62)	< 0.00
Month of dose	O1 (Jan–Mar)	4080/8262 (49.4)	4.14 (3.33–5.14)	< 0.001	3.79 (3.01–4.76)	< 0.00
	O2 (Apr-Jun)	144/625 (23.0)	1.00		1.00	
	$O_3 (Jul-Sep)$	0/1 (0.00)	N/A		N/A ^a	
	Q4 (Oct-Dec)	38/1125 (31.3)	1.94 (1.24–3.02)	0.003	4,28 (2,69–6,80)	<0.00
Age vears	16-29.99	45/120 (37.5)	1.00	0.005	1.00	0.00
	30-39.99	121/341 (35.5)	0.92 (0.59–1.42)	0.69	0.82 (0.49–1.33)	0 44
	40-49.99	373/800 (46.6)	1 42 (0.96-2.12)	0.09	0.90 (0.57_1.42)	0.44
	50-59 99	1286/2192 (58.7)	7.42 (0.90 - 2.12) 2 36 (1 61 - 3 45)	<0.00	1.03(0.66-1.61)	0.05
	60, 69,99	1716/3617 (47.4)	2.50 (1.01 - 3.75) 1 56 (1 07 - 2 28)	0.001	0.72 (0.46 - 1.12)	0.50
	>70.00	721/1022 (27.2)	1.30(1.07-2.28) 1.10(0.75-1.61)	0.021	0.72(0.40-1.12) 0.58(0.37-0.92)	0.14
	≥70.00 P for trend	721/1955 (57.5)	1.10 (0.75-1.01)	0.04	0.38 (0.37 - 0.92)	-0.020
Sov	Fioritienu	2207/6267 (51.9)	1.00		0.85 (0.81-0.90)	<0.00
Sex	remaie	5297/0507 (51.8) 065/2626 (26.6)		<0.001		-
		905/2030 (30.0)	0.57 (0.52-0.63)	< 0.001	0.59 (0.53-0.65)	< 0.00
IMD rank, quartile	QT (most deprived)	906/2030 (44.7)	0.84 (0.75-0.95)	0.006	0.90 (0.79–1.02)	0.10
	02	1020/2160 (47.2)	0.95 (0.84–1.07)	0.39	0.95 (0.84–1.08)	0.44
	Q3	1129/2343 (48.2)	1.00 (0.89–1.12)	0.96	1.00 (0.89–1.13)	0.97
	Q4 (least deprived)	1182/2430 (48.7)	1.00		-	-
- 1	P for trend		1.00		0.96 (0.92–1.00)	0.070
lobacco smoking status	Not current smoker	409//8633 (4/.5)	1.00		1.00	0.04
AL 1 1 1 1 1 1	Current smoker	165/3/0 (44./)	0.83 (0.67–1.03)	0.09	0.80 (0.64–1.01)	0.06
Alcohol, units/wk	0	1170/2349 (49.8)	1.00	0.01	1.00	0.01
	1-7	1530/3189 (48.0)	0.95 (0.85-1.05)	0.31	1.01 (0.90–1.13)	0.91
	8–14	861/1853 (46.5)	0.93 (0.82–1.05)	0.25	1.02 (0.90–1.18)	0.68
	15–21	398/901 (44.2)	0.86 (0.73–1.01)	0.06	0.91 (0.77–1.08)	0.30
	22-28	1/8/403 (44.3)	0.91 (0.73–1.14)	0.42	0.98 (0.78–1.24)	0.89
	>28	125/308 (40.6)	0.80 (0.62–1.03)	0.08	0.84 (0.65–1.10)	0.21
	P for trend				0.98 (0.94–1.02)	0.26
Self-rated general health	Excellent	801/18,758 (42.7)	1.00		1.00	
	Very good	1620/3590 (45.1)	1.15 (1.02–1.29)	0.019	1.13 (1.00–1.28)	0.05
	Good	1162/2326 (50.0)	1.37 (1.21–1.56)	<0.001	1.38 (1.20–1.58)	<0.001
	Fair	530/949 (55.9)	1.67 (1.42–1.96)	<0.001	1.53 (1.28–1.83)	<0.001
	Poor	149/260 (57.3)	1.74 (1.33–2.27)	<0.001	1.54 (1.14–2.07)	0.005
	P for trend				1.15 (1.10–1.21)	<0.001
Pre-vaccination SARS-CoV-2 status	Seronegative	3560/7640 (46.6)	1.00		1.00	
	Seropositive asymptomatic	439/958 (45.8)	0.99 (0.86–1.13)	0.83	0.94 (0.81–1.09)	0.42
	Seropositive symptomatic	263/405 (64.9)	2.01 (1.62–2.48)	<0.001	2.23 (1.78–2.81)	<0.001
	P for trend				1.26 (1.15–1.38)	<0.001
Self-rated anxiety or depression	No	3040/6744 (45.1)	1.00		1.00	
	Yes	1220/2253 (54.2)	1.35 (1.23–1.49)	<0.001	1.24 (1.12–1.39)	<0.001
Asthma	No	3489/7552 (46.2)	1.00		1.00	
	Yes	773/1451 (53.3)	1.29 (1.15–1.45)	<0.001	1.12 (0.95–1.32)	0.18
Atopic disease ^b	No	3041/6684 (45.5)	1.00		1.00	
	Yes	1221/2319 (52.7)	1.28 (1.16–1.41)	<0.001	1.20 (1.08–1.34)	0.00
Arterial disease	No	4040/8514 (47.5)	1.00		1.00	
	Yes	222/489 (45.4)	1.21 (1.00–1.47)	0.049	1.31 (1.06–1.63)	0.014
Kidney disease	No	4164/8821 (47.2)	1.00		1.00	
	Voc	98/182 (53.9)	1 47 (1 00_1 00)	0.012	1 /0 /1 01 1 02)	0.043

		N (%) symptomatic	Minimally adjusted		Fully adjusted	
			aOR (95% CI)	Р	aOR (95% CI)	Р
Cancer	Never	3891/8117 (47.9)	1.00		1.00	
	Previous	346/809 (42.8)	0.84 (0.72–0.97)	0.021	0.83 (0.71–0.97)	0.019
	Active	25/77 (32.5)	0.64 (0.39–1.05)	0.078	0.62 (0.37-1.05)	0.076
Statins	No	3620/7,377 (49.1)	1.00		1.00	
	Yes	642/1626 (39.5)	0.87 (0.77–0.98)	0.022	0.84 (0.74–0.97)	0.014
ACE inhibitors	No	3903/8101 (48.2)	1.00		1.00	
	Yes	359/902 (39.8)	0.82 (0.7–0.95)	0.008	0.80 (0.69–0.94)	0.007
Inhaled corticosteroids	No	3951/8415 (47.0)	1.00		1.00	
	Yes	311/588 (52.9)	1.24 (1.04–1.47)	0.015	0.98 (0.78–1.25)	0.89
Systemic immunosuppressants	No	4051/8607 (47.1)	1.00		1.00	
	Yes	211/396 (53.3)	1.29 (1.05–1.59)	0.014	1.10 (0.87–1.38)	0.423
Inhaled bronchodilators	No	3807/8160 (46.6)	1.00		1.00	
	Yes	453/837 (54.1)	1.31 (1.14–1.52)	<0.001	1.49 (0.63–3.50)	0.37
Beta-2 adrenergic agonists	No	3,822/8,192 (46.7)	1.00		1.00	
	Yes	440/811 (54.3)	1.31 (1.13–1.51)	<0.001	0.66 (0.27–1.60)	0.36
Multivitamin supplement	No	3258/7168 (45.5)	1.00		1.00	
	Yes	1004/1835 (54.7)	1.42 (1.28–1.58)	<0.001	1.41 (1.26–1.58)	<0.001
Vitamin D supplement	No	3213/6690 (48.1)	1.00		1.00	
	Yes	1049/2133 (45.3)	1.81 (0.97–3.38)	0.06	0.92 (0.83–1.02)	0.12

of reporting systemic symptoms after first vaccine dose: time of year (aOR 3.79, 95% CI: 3.01-4.76, for Q1 vs. Q2; aOR 4.28, 95% CI: 2.69–6.80, for Q4 vs. Q2), poorer self-rated general health (aOR per category of worsening health 1.15, 95% CI: 1.10-1.21), prevaccination SARS-CoV-2 seropositivity with symptoms vs. prevaccination SARS-CoV-2 seronegativity (aOR 2.23, 95% Cl: 1.78-2.81), presence vs. absence of self-rated anxiety or depression (aOR 1.24, 95% CI: 1.12-1.39), presence vs. absence of atopic disease (aOR 1.20, 95% CI: 1.08-1.34), presence vs. absence of arterial disease (aOR 1.31, 95% CI: 1.06–1.63), presence vs. absence of kidney disease (aOR 1.40, 95% CI: 1.01-1.93), and use vs. no use of multivitamin supplements (aOR 1.41, 95% CI: 1.26-1.58). Lower risk of developing systemic post-vaccination symptoms following first dose was independently associated with administration of mRNA vaccines vs. ChAdOx1 (aOR 0.29, 95% CI: 0.26-0.32 for BNT162b2 vs. ChAdOx1; aOR 0.06, 95% CI: 0.01-0.26 for mRNA-1273 vs. ChAdOx1), greater age (aOR per additional 10 years of age 0.85, 95% CI: 0.81-0.90), male vs. female sex at birth (aOR 0.59, 95% CI: 0.53-0.65), presence vs. absence of previous cancer (aOR 0.83, 95% CI: 0.71-0.97), use vs. no use of statins (aOR 0.84, 95% CI: 0.74-0.97), and use vs. no use of ACE inhibitors (aOR 0.80, 95% CI: 0.69 - 0.94).

Given that we showed no association between asymptomatic pre-vaccination SARS-CoV-2 infection and risk of systemic symptoms after a first vaccine dose (aOR 0.94, 95% Cl: 0.81–1.09, Table 2), we conducted an exploratory analysis to compare pre-vaccination anti-S titres between participants who experienced symptomatic vs. asymptomatic SARS-CoV-2 infection prior to vaccination: the former group had higher anti-S titres than the latter (P < 0.001, Supplementary Fig. 2).

In another exploratory analysis, we investigated whether these factors also associated independently with the mean total number of different systemic symptoms reported by the subset of participants reporting at least one such symptom. Results are displayed in Supplementary Table 4, Supplementary Material. Higher mean number of systemic symptoms reported associated independently with younger age (*P* for trend 0.01), female vs. male sex (1.62 vs. 1.54, P = 0.028), symptomatic SARS-CoV-2 infection vs. no SARS-CoV-2 infection pre-vaccination (1.76 vs. 1.58, P = 0.013), administration of ChAdOx1 vs. BNT162b2 (1.69 vs. 1.20, P < 0.001) and presence vs. absence of atopic disease (1.66 vs. 1.58, P = 0.019).

Determinants of systemic symptoms following a second dose of vaccine were broadly similar, in that administration of BNT162b2 vs. ChAdOx1, vaccination in Q1 vs. Q2, poorer vs. better self-rated general health, pre-vaccination SARS-CoV-2 seropositivity with symptoms vs. pre-vaccination SARS-CoV-2 seronegativity, presence vs. absence of self-rated anxiety or depression, presence vs. absence of atopic disease, and use vs. no use of multivitamin supplements associated with higher risk of reporting symptoms and greater age, and male vs. female sex at birth all associated with lower risk. Additionally, a shorter inter-dose interval associated with reduced risk of systemic symptoms after the second vaccine dose (aOR 0.61, 95% CI: 0.46-0.83 for <6 weeks vs. >10 weeks; aOR 0.86, 95% CI: 0.76-0.96 for 6-10 weeks vs. >10 weeks), as did higher alcohol intake (aOR per increasing category 0.93, 95% Cl: 0.89-0.96) and use vs. no use of anticholinergic medication (aOR 0.73, 95% CI: 0.53-0.99; Supplementary Table 5, Supplementary Material).

Determinants of local reactive symptoms following first vaccine dose

After adjustment for age and sex only, 33 variables were found to associate with risk of experiencing local symptoms following a first vaccine dose with P < 0.10 (Table 3) and 8 were not (Supplementary Table 6, Supplementary Material). After inclusion of all of these factors in a multivariable model, the following variables were independently associated with higher risk of reporting local symptoms after first dose: administration of BNT162b2 vs. ChAdOx1 (aOR 1.95 95% CI: 1.74–2.18), administration of vaccine

		Minimally adjusted Fully adjusted				
		N (%) symptomatic	aOR (95% CI)	P	aOR (95% CI)	Р
Vaccine type	ChAdOx1	2414/5988 (40.3)	1.00		1.00	
	BNT162b2	1470/2864 (51.3)	1.78 (1.62–1.96)	< 0.001	1.95 (1.74–2.18)	< 0.001
	MRNA-1273	6/59 (10.2)	0.16 (0.07–0.38)	<0.001	N/A	N/A
	Other	24/92 (26.1)	0.50 (0.31-0.81)	0.005	0.47 (0.27-0.82)	0.00
Timing	Before 12:00	1520/3193 (47.6)	1 00	0.005	0.17 (0.27 0.02)	1 00
	12:00-14:00	745/1425 (52.3)	1 23 (1 08–1 41)	0.002	1 23 (1 07–1 41)	0.004
	14:00-17:00	1097/2200 (49.9)	1.09 (0.98–1.22)	0.002	1.05 (0.93–1.18)	0.00
	After 17:00	509/1030 (49.4)	1.03 (0.89–1.19)	0.720	1.00 (0.85–1.17)	0.99
Month of dose	O1 (lan-Mar)	3722/8262 (45.1)	5.07 (4.04–6.35)	<0.001	1.11 (0.81–1.52)	0.55
	Ω^2 (Apr-lup)	129/625 (20.6)	1 00		1.00	0.51
	$O_3 (u -Sep)$	0/1 (0.00)	N/A		N/A	
	O4 (Oct-Dec)	63/115 (54.8)	6 90 (4 46-10 7)	<0.001	0.89 (0.52–1.53)	0.68
Age vears	16-29.99	50/120 (41 7)	1.00	<0.001	1.00	0.00
	30_39.99	128/341 (37.5)	0.84 (0.55_1.29)	0.43	0.51 (0.21_1.28)	0.15
	40-49.99	389/800 (48.6)	1 29 (0.87_1.91)	0.45	0.46 (0.20-1.09)	0.15
	50 50 00	1164/2102 (53.1)	1.25(0.07 - 1.91) 1 57 (1 08 2 20)	0.20	0.40 (0.20 - 1.00)	0.00
	60, 69,99	1546/2617 (42.7)	1.09 (0.75 1.58)	0.66	0.17 (0.07-0.38)	~0.002
	>70.00	637/1033 (33.0)	0.77 (0.53 - 1.30)	0.00	0.17 (0.07 - 0.38)	<0.00
	\geq 70.00	037/1933 (33.0)	0.77 (0.55-1.15)	0.10	0.10(0.04-0.23)	<0.00
Sov	Fibritiend	2101/6267 (10 0)	1.00		1.00	<0.00
Jex .	Malo	810/2636 (30 7)	0.50 (0.46, 0.55)	<0.001	0.48 (0.43_0.54)	~0.00
Education	Primany/Sacandany	291/079 (20.0)	0.30(0.40-0.33)	<0.001	0.48 (0.43-0.34)	0.00
Education	Higher/Eurther	552/1265 (427)	0.05 (0.92 1.00)	0.012	0.78(0.00-0.94)	0.003
		555/1205 (45.7) 1738/2001 (43.2)	0.95 (0.85 - 1.09)	0.40	0.97 (0.82 - 1.34)	0.00
	College Dest graduate	1720/3991 (43.3)	1.00	0.24	0.90 (0.85 - 1.08)	0.50
	Post-graduate	1240/2702 (45.2)	1.00		1.00 (rer)	0.02
		1520/2250 (45 5)	1 10 (1 06 1 22)	0.000	0.94 (0.89-0.99)	0.024
vigorous physical exercise, hr/wk	0	1529/3359 (45.5)	1.19 (1.06–1.32)	0.002	1.02 (0.89–1.17)	0.74
	1-3	1450/3323 (43.6)	1.09 (0.97-1.21)	0.15	1.05 (0.92-1.19)	0.49
	≥ 4	926/2301 (40.3)	1.00		1.00	0.70
	P for trend	1205/2022 (46.1)	1 16 (1 04 1 20)	0.000	0.99 (0.92-1.06)	0.78
Light physical exercise, hr/wk	0-4	1305/2832 (46.1)	1.16 (1.04–1.29)	0.006	1.05 (0.92-1.19)	0.49
	5-9	1311/2983 (44.0)	1.07 (0.96–1.18)	0.24	1.07 (0.95–1.21)	0.27
	≥10 D fan twas d	1293/3172 (40.8)	1.00		1.00	0.46
	P for trend	2672/0500 (42.2)	1.00		0.98 (0.91–1.04)	0.46
Dietary restrictions	None	36/9/8500 (43.3)	1.00	0.15		1.00
	Vegetarian	196/397 (49.4)	1.16 (0.95–1.43)	0.15	1.36 (1.05–1.76)	0.019
	Vegan	36/9/8500 (43.3)	0.70 (0.47–1.04)	0.080	0.64 (0.40–1.03)	0.067
Self-rated general health	Excellent	/11/18/8 (37.8)	1.00		1.00	
	Very good	1491/3590 (41.5)	1.21 (1.08–1.36)	0.001	1.19 (1.04–1.36)	0.013
	Good	1099/2326 (47.3)	1.50 (1.32–1.70)	< 0.001	1.43 (1.23–1.68)	< 0.00
	Fair	481/949 (50.7)	1.64 (1.40–1.93)	<0.001	1.49 (1.21–1.83)	< 0.001
	Poor	132/260 (50.8)	1.60 (1.23–2.09)	0.001	1.41(1.00–1.99)	0.048
	P for trend				1.14 (1.08–1.21)	<0.001
Body mass index, kg/m²	<25	1882/4369 (43.1)	1.00		1.00	
	25–30	1226/2907 (42.2)	1.04 (0.94–1.14)	0.38	0.95 (0.85–1.07)	0.38
	>30	799/1709 (46.8)	1.12 (1.00–1.27)	0.05	0.88 (0.76–1.02)	0.08
Pre-vaccination SARS-CoV-2 status	Seronegative	3270/7640 (42.8)	1.00			1.00
	Seropositive asymptomatic	407/958 (42.5)	1.00 (0.87–1.15)	0.96	0.96 (0.81–1.12)	0.58
	Seropositive symptomatic	237/405 (58.5)	1.74 (1.41–2.14)	<0.001	1.84 (1.43–2.38)	<0.001
	P for trend				1.19 (1.08–1.32)	0.001

Table 3 continued

			Minimally adjust	ted	Fully adjusted		
		N (%) symptomatic	aOR (95% CI)	Р	aOR (95% CI)	Р	
Self-rated anxiety or depression	No	2806/6744 (41.6)	1.00		1.00		
	Yes	1106/2253 (49.1)	1.23 (1.11–1.35)	<0.001	1.17 (1.03–1.32)	0.018	
Asthma	No	3206/7552 (42.5)	1.00		1.00		
	Yes	708/1451 (48.8)	1.23 (1.10–1.38)	<0.001	1.17 (0.97–1.41)	0.10	
Atopic disease ^a	No	2781/6684 (41.6)	1.00		1.00		
	Yes	1133/2319 (48.8)	1.27 (1.15–1.40)	<0.001	1.15 (1.02–1.30)	0.019	
Diabetic status	No diabetes	3612/8278 (43.6)	1.00		1.00		
	Pre-diabetes	120/291 (41.0)	1.03 (0.81–1.32)	0.79	0.88 (0.67–1.17)	0.38	
	Type 1 diabetes	35/61 (57.4)	1.84 (1.09–3.09)	0.022	1.02 (0.57–1.84)	0.94	
	Type 2 diabetes	141/363 (38.6)	0.99 (0.79–1.24)	0.95	0.94 (0.71–1.25)	0.68	
Heart disease	No	3769/8642 (43.6)	1.00		1.00		
	Yes	145/361 (40.2)	1.23 (0.99–1.54)	0.07	0.78 (0.49–1.22)	0.27	
Arterial disease	No	3708/8514 (43.5)	1.00		1.00		
	Yes	206/489 (42.1)	1.33 (1.10–1.62)	0.004	1.43 (0.90–2.28)	0.13	
Major neurological condition	No	3802/8752 (43.4)	1.00		1.00		
, ,	Yes	112/251 (44.6)	1.29 (1.00–1.68)	0.05	1.00 (0.66–1.51)	1.00	
Immunodeficiency	No	3883/8950 (43.4)	1.00		1.00		
,	Yes	31/53 (58.5)	2.03 (1.16-3.55)	0.013	1.80 (0.92-3.47)	0.09	
Statins	No	3272/7377 (44.4)	1.00		1.00		
	Yes	642/1626 (39.5)	1.18 (1.04–1.33)	0.008	1.09 (0.94–1.28)	0.26	
Proton pump inhibitors	No	3354/7752 (43.3)	1.00	0.000	1.00	0.20	
	Yes	560/1251 (44.8)	1.00	0.005	0.98 (0.84–1.14)	0.77	
Inhaled corticosteroids	No	3622/8415 (43.0)	1.00	0.005	1.00	0.77	
	Yes	292/588 (49.7)	1.00	0.006	1.00 (0.79_1.32)	0.89	
SSRIs	No	3619/8430 (42.9)	1.00	0.000	1.02 (0.75 1.52)	0.05	
55115	Yes	295/573 (51 5)	1.00	0.022	1.06 (0.86–1.32)	0.57	
Thiazidas	No	3782/8708 (43.4)	1.00	0.022	1.00 (0.00 1.52)	0.57	
Thatices	Vec	132/205 (44.8)	1.00	0.07	1.00	0 14	
Inhaled bronchodilators	No	3807/8160 (46.6)	1.25 (0.50-1.55)	0.07	1.24 (0.24-1.03)	0.14	
	No	A52/927 (54.1)	1.00	<0.001	1.00	0.27	
Sodium-ducose co-transporter-2 (SCIT2)	No	3001/8055 (43.6)	1.00	<0.001	1.49 (0.05-5.50)	0.57	
inhibitors	No	13/48 (27.1)	0.57 (0.30 1.00)	0.00	0.48 (0.22 1.05)	0.07	
Anti platalat drugs	No	2666/9404 (42.6)	1.00	0.09	1.00	0.07	
Anti-platelet drugs	No	249/500 (41.4)	1.00	0.011	0.00 (0.54 1.44)	0.61	
Poto 2 adronargie agoniste	No	240/399 (41.4)	1.20 (1.00-1.50)	0.011	1.00	0.01	
beta-2 adrenergic agonists	NO	3622/6192 (40.7)	1.00	-0.001	1.00	0.26	
Acairia	Ne	440/011 (34.3)	1.51 (1.15-1.51)	<0.001	1.00	0.50	
Aspinn	NO	3/10/8531 (43.0)	1.00	0.010	1.00	0.44	
PCC see singly d	res	198/4/2 (42.0)	1.29 (1.06-1.56)	0.012	1.22 (0.74–2.02)	0.44	
BCG vaccinated	NO	414/1087 (38.1)	1.00	0.001	1.00	0.012	
	Yes	319///084 (45.1)	1.25 (1.09–1.43)	0.001	1.21 (1.04–1.41)	0.013	
	NO	3238//108 (45.5)	1.00	-0.001	1.00	10.001	
	Yes	1004/1835 (54.7)	1.42 (1.28–1.58)	<0.001	1.41 (1.26–1.58)	<0.001	
Vitamin D supplement	No	3213/6690 (48.1)	1.00		1.00		
	Yes	1049/2133 (45.3)	1.81 (0.97–3.38)	0.06	0.92 (0.83–1.02)	0.12	

at lunchtime (i.e. 12:00–14:00) vs. morning (i.e. before 12:00; aOR 1.23, 95% Cl: 1.07–1.41), vegetarian vs. unrestricted diet (aOR 1.36, 95% Cl: 1.05–1.76), poorer vs. better self-rated general health (aOR per worsening health category 1.14, 95% Cl: 1.08–1.21), prevaccination SARS-CoV-2 seropositivity with symptoms vs. pre-

vaccination SARS-CoV-2 seronegativity (aOR 1.84, 95% Cl: 1.43–2.38), presence vs. absence of self-rated anxiety or depression (aOR 1.17, 95% Cl: 1.03–1.32), presence vs. absence of atopic disease (aOR 1.15, 95% Cl: 1.02–1.30), BCG vaccinated vs. not (aOR 1.21, 95% Cl: 1.04–1.41), and use vs. no use of multivitamin

	Ν	Mean titre (s.d.)	Adjusted mean difference (95% Cl) ^a	Ρ	P for trend
Local reactive symptoms					
Never	2150	2.64 (2.23)	– (ref)	-	0.007 ^b
After one dose only	1212	2.78 (2.22)	0.05 (-0.10 to 0.19)	0.534	
After both doses	995	3.03 (2.76)	0.23 (0.07 to 0.38)	0.004	
Systemic reactive sympton	ms				
Never	1437	2.54 (2.06)	– (ref)	-	<0.001 ^c
After one dose only	1812	2.83 (2.42)	0.25 (0.11 to 0.39)	0.001	
After both doses	1108	2.95 (2.25)	0.33 (0.17 to 0.49)	<0.001	
Local reactive symptoms					
Never	720	4.54 (3.30)	– (ref)	-	0.029 ^d
After one dose only	501	4.78 (3.06)	0.30 (-0.05 to 0.64)	0.097	
After both doses	805	4.93 (3.21)	0.34 (0.04 to 0.65)	0.027	
Systemic reactive sympton	ms				
Never	1120	4.49 (3.00)	– (ref)	-	<0.001 ^e
After one dose only	548	4.81 (3.13)	0.30 (-0.01 to 0.61)	0.058	
After both doses	358	5.51 (3.80)	0.75 (0.38 to 1.11)	<0.001	

^aAdjusted for pre-vaccination anti-S titre and number of days from date of second vaccine dose to date of post-vaccination antibody measurement. ^bAssociated beta coefficient per increasing category 0.11 (95% CI: 0.03–0.18).

^cAssociated beta coefficient per increasing category 0.17 (95% CI: 0.09–0.25).

^dAssociated beta coefficient per increasing category 0.17 (95% Cl: 0.02–0.32).

^eAssociated beta coefficient per increasing category 0.36 (95% Cl: 0.19–0.53).

supplements (aOR 1.41, 95% CI: 1.26-1.58). Lower risk of developing local post-vaccination symptoms following first dose was independently associated with greater age (aOR per additional 10 years of age 0.62, 95% CI: 0.59-0.66), male vs. female sex at birth (aOR 0.48, 95% Cl: 0.43-0.54), and lower educational level (aOR per decreasing education category 0.94, 95% CI: 0.89-0.99). Determinants of local symptoms following a second dose of vaccine were broadly similar, in that administration of BNT162b2 vs. ChAdOx1, poorer vs. better self-rated general health, pre-vaccination SARS-CoV-2 seropositivity with symptoms vs. pre-vaccination SARS-CoV-2 seronegativity, presence vs. absence of self-rated anxiety or depression, presence vs. absence of atopic disease, and use vs. no use of multivitamin supplements associated with higher risk of reporting local symptoms and greater age, male vs. female sex at birth and lower educational level associated with lower risk. Additionally, use vs. no use of H2receptor antagonists associated with increased risk of local symptoms after second dose (aOR 2.05, 95% CI: 1.21-3.47), while use vs. no use of anticholinergics (aOR 0.73, 95% CI: 0.53-0.99) and previous history of cancer vs. no such history (aOR 0.83, 95% Cl: 0.70-0.99) both associated with reduced risk; Supplementary Table 7, Supplementary Material).

Association between reactive symptoms and post-vaccination anti-S titres

Table 4 displays mean post-vaccination antibody titres by frequency of post-vaccination symptoms, stratified by vaccine type (ChAdOx1 vs BNT162b2) and adjusted for duration between date of second dose and date of post-vaccination antibody measurement and pre-vaccination anti-S titres. For both ChAdOx1 and BNT162b2, a dose–response relationship between increasing symptom frequency and higher post-vaccination antibody titres was observed for both local and systemic symptoms (*P* for trend \leq 0.02). A sensitivity analysis excluding participants who had SARS-CoV-2 infection prior to vaccination (as evidenced by presence of anti-S antibodies and/or a positive RT-PCR or antigen

test for SARS-CoV-2) yielded consistent findings (Supplementary Table 8, Supplementary Material).

Association between long COVID and risk of reactive symptoms following vaccination

Supplementary Table 9 presents results of an exploratory analysis restricted to participants with evidence of SARS-CoV-2 infection prior to vaccination, conducted to determine whether presence or absence of self-reported 'long COVID' associated with risk of reactive symptoms following SARS-CoV-2 vaccination. After adjustment for all covariates included in the final models described above, we found that self-reported long COVID associated independently with increased risk of both systemic and local reactive symptoms after a first vaccine dose (systemic, aOR 1.73, 95% Cl: 1.12–3.35; local, aOR 1.94, 95% Cl: 1.41–2.66). However, self-reported long COVID did not associate independently with risk of either systemic or local reactive symptoms after a second vaccine dose.

DISCUSSION

This large, prospective, population-based study represents the most comprehensive investigation of incidence, determinants and serological correlates of reactogenic symptoms following SARS-CoV-2 vaccination that has been conducted to date. We found that administration of ChAdOx1, younger age, female sex, poorer self-rated general health, pre-vaccination COVID-19, presence of self-rated anxiety or depression at enrolment, presence of atopic disease, and use of multivitamin supplements associated independently with increased odds of reporting both systemic and local symptoms following SARS-CoV-2 vaccination. A sub-set of these factors (younger age, female sex, pre-vaccination COVID-19, administration of ChAdOx1 and presence of atopic disease) also associated independently with a higher mean number of systemic symptoms reported. Those who received BNT162b2 vaccine, as compared to those who received ChAdOx1, were more likely to report local reactogenic symptoms and less likely to report

systemic symptoms. Importantly, for both ChAdOx1 and BNT162b2, we found a dose-response relationship between increasing frequency of both local and systemic reactogenic symptoms and higher post-vaccination anti-S titres, that persisted when we excluded participants who had evidence of SARS-CoV-2 infection prior to vaccination.

Our findings are in keeping with those of other studies that have reported increased risk of post-vaccination symptoms in women vs. men^{6,7,11}, in younger vs. older people^{5–8} and in those who had COVID-19 prior to vaccination⁶. However, we advance knowledge in this area by showing that participants who had asymptomatic SARS-CoV-2 infection prior to vaccination (as evidenced by presence of anti-S antibodies in the absence of any self-report of COVID-19 symptoms) were not at higher risk of post-vaccination symptoms than those who were anti-S seronegative prior to vaccination. This might reflect the presence of weaker pre-vaccination memory responses to SARS-CoV-2 in those who had asymptomatic vs. symptomatic SARS-CoV-2 infection, as evidenced by our demonstration of lower pre-vaccination anti-S titres in the former group. We also show for the first time that those reporting anxiety or depression at enrolment, those taking multivitamin supplements and vegetarians are more likely to report post-vaccination symptoms. The former association may reflect increased perception of physical symptoms that has previously been reported in people with anxiety and depression¹². The latter might arise as a result of immunomodulatory effects of micronutrient supplements, which could augment reactogenic symptoms by boosting vaccine immunogenicity¹³; alternatively, participants taking micronutrient supplements or who maintain a vegetarian diet may be more health-conscious^{14,15}, and thus have an increased awareness of post-vaccination symptoms. The incidence of systemic post-vaccination reactogenic symptoms reported in our study is broadly in keeping with that reported by other studies^{5–8}, although incidence of local symptoms reported in this study is lower than for others5-8; this may reflect the relatively high proportion of participants receiving ChAdOx1 vs. mRNA vaccines in our study vs. others. Additionally, we highlight that our serological outcome was a combined IgG/IgA/IgM response to spike protein measured using a semiguantitative ELISA. This contrasts with other studies investigating associations between vaccine-related symptoms and serologic responses to SARS-CoV-2vaccination, which primarily use quantitative levels of IgG anti-S antibodies.

Our study has several strengths. Its prospective nature reduces the potential for reverse causation to explain the associations we report; for example, we show that anxiety is a prequel to reporting symptoms, and not a consequence of experiencing them. We investigated a comprehensive range of potential determinants of reactogenic symptoms, and captured detailed symptom profiles within days to weeks of vaccine administration, thereby reducing imprecision due to poor symptom recall. Additionally, the availability of pre- and post-vaccination anti-S status and COVID-19 symptom data allowed us to dissect apart the influence of symptomatic vs. asymptomatic SARS-CoV-2 infection on risk of reactogenic symptoms. A further strength is the use of an ELISA with high sensitivity and specificity that targets three different types of antibody (IgG/IgA/IgM), increasing the probability of identifying a past infection over other studies which primarily use quantitative levels of IgG anti-spike antibodies¹⁶. Additionally, we used dried blood spots for our sampling: although these yield a low volume of sample and their results may be impacted by variations in haematocrit, their use has been found to significantly reduce processing failures compared with microtubes¹⁷, which are currently used by large seroprevalence surveys¹⁸

Our study also has some limitations. First, COVIDENCE UK is a self-selected cohort, and thus several groups—such as people younger than 30 years, people of lower socioeconomic status, and non-White ethnic groups—are under-represented. Second, in

common with other studies in this field, we lack objective data relating to characterisation of reactions to vaccination; accordingly, the presence or absence of reported symptoms represents a function of the presence or absence of a physiological reaction to vaccination and the perception of that reaction. Third, those who had vaccines other than BNT162b2 or ChAdOx1 are underrepresented, and this limited our power to investigate reactogenic symptoms following administration of other SARS-CoV-2 vaccines. Fourth, the duration of symptoms was not considered. Lastly, as with any observational study, we cannot exclude the possibility that the associations we report may be explained by residual and/ or unmeasured confounding. We sought to minimise this by capture of, and mutual adjustment for, a comprehensive panel of potential confounders.

In conclusion, we found that factors that associate with lower risk of systemic post-vaccination symptoms include greater age, male sex, and receipt of an mRNA vaccine. Factors that associate with a higher risk of systemic post-vaccination symptoms include history of symptomatic SARS-CoV-2 infection prior to vaccination, presence of self-rated anxiety or depression and use of a multivitamin supplement. Finally, we found vaccine reactogenicity was associated with higher levels of antibody titres for both BNT162b2 and ChAdOx1 vaccines. Public communication of this message may provide a degree of reassurance to those who have experienced symptoms after SARS-CoV-2 vaccination and allow them to better understand the relationship between experiencing such symptoms and vaccine efficacy; this may in turn provide added motivation for them to receive further vaccine doses.

METHODS

Study design, setting and participants

Our study was nested within COVIDENCE UK, which is a prospective longitudinal population-based observational study of coronavirus disease in the UK population (www.qmul.ac.uk/ covidence)^{16,17,19}. The study was launched on 1st May 2020, and closed to enrolment on 6th October 2021. Inclusion criteria for the cohort as a whole were age 16 years or more and residence in the UK at the time of enrolment. Participants were invited via a national media campaign to complete an online baseline questionnaire to capture information on potential symptoms of COVID-19 experienced since 1st February 2020, results of any COVID-19 tests, and details of a wide range factors that may influence risk of COVID-19 and response to vaccination, as described below. Follow-up questionnaires were administered at monthly intervals to capture incident test-confirmed COVID-19, potential symptoms of COVID-19, administration of SARS-CoV-2 vaccines and post-vaccination symptoms. Study participants were invited to provide dried blood spots for testing for combined IgG, IgA and IgM responses to the SARS-CoV-2 spike protein before and after SARS-CoV-2 vaccination as described elsewhere¹⁶⁻¹⁸. Analyses for the current study were restricted to the subset of participants whose pre-vaccination anti-S serostatus was determined; who completed a full primary course of SARS-CoV-2 vaccination; and who reported their post vaccination symptoms. This paper reports findings from analyses of data collected up to 1st September 2021.

Sponsorship, registration, ethics and reporting

The study was sponsored by Queen Mary University of London and approved by Leicester South Research Ethics Committee (ref 20/EM/0117). All participants provided informed consent by completing online consent forms. The study is registered with ClinicalTrials.gov (NCT04330599).

Independent variables

Fifty-six putative risk factors for symptoms following administration of a SARS-CoV-2 vaccine were selected a priori, covering sociodemographic, occupational and lifestyle factors; longstanding medical conditions and prescribed medication use; Bacille Calmette Guérin vaccine status; and diet and supplemental micronutrient intake (Supplementary Tables 1 and 2 Supplementary Material). These factors, which were obtained from the baseline and follow-up questionnaires, were included as independent variables in our models. To produce patient-level covariates for each class of medications investigated, participant responses were mapped to drug classes listed in the British National Formulary, or the DrugBank and Electronic Medicines Compendium databases if not explicitly listed in the British National Formulary, as previously described¹⁹. Index of Multiple Deprivation (IMD) 2019 scores were assigned according to participants' postcodes, and categorised into guartiles. Duration of follow-up was defined as the number of days between the date of enrolment and the date of dried blood spot collection.

Outcomes

The primary outcome for the current study was new symptoms following administration of the first dose of a SARS-CoV-2 vaccine. Symptoms were classified as being systemic (namely fatigue, headache, fever/high temperature [37.8 °C or greater] or myalgia) or local (namely tenderness, soreness, swelling, redness or a painful heavy feeling at the injection site and axillary or cervical lymphadenopathy). The questions used to capture these symptoms are presented in Supplementary Table 2, Supplementary Material. Secondary outcomes were incidence of symptoms after second vaccine doses, and titres of combined IgG, IgA and IgM anti-Spike antibody responses following vaccination, which were determined in eluates of postal dried blood spot samples as described below.

Procedures for antibody testing

Procedures for the pre- and post-vaccination antibody studies are reported elsewhere^{16,17}. Briefly, participants in these studies were sent kits containing instructions, lancets, and blood spot collection cards, to be posted back to the study team. Once returned, the samples were logged by the study team and sent in batches to the Clinical Immunology Service at the Institute of Immunology and Immunotherapy of the University of Birmingham (Birmingham, UK). Up to two more test kits were offered to participants whose initial samples were found to be insufficient for testing.

To elute antibody from DBS cards, we isolated individual preperforated DBS spots by using a sterile pipette tip and placed them into a universal tube at a ratio of 1 spot to 250 µL 0.05% phosphate-buffered saline (PBS)-Tween 20 (PBS-T) (PBS, xoid; Tween-20; Sigma-Aldrich, https://www.sigmaaldrich.com). We briefly vortexed and incubated tubes overnight at room temperature. We then harvested DBS eluate into a microtube and centrifuged it at $10,600 \times q$ for 10 min at room temperature. Eluates were stored at 4 °C for <14 days, prior to semi-quantitative determination of antibody titres. This was done using a commercially available ELISA that measures combined IgG, IgA, and IgM (IgGAM) responses to the SARS-CoV-2 trimeric spike glycoprotein (product code MK654; The Binding Site [TBS], Birmingham, UK). The SARS-CoV-2 spike used is a soluble, stabilised, trimeric glycoprotein truncated at the transmembrane region. 8,9 This assay has been CE-marked with 98.3% (95% CI: 96.4-99.4) specificity and 98.6% (92.6-100.0) sensitivity following RT-PCR-confirmed mild-to-moderate COVID-19 that did not result in hospitalisation.10 A cut-off ratio relative to the TBS cut-off calibrators was determined by plotting 624 pre-2019 negatives in a frequency histogram. A cut-off coefficient was then established for IgGAM (1.31), with ratio values classed as positive (\geq 1) or negative (<1). Dried blood spots were pre-diluted at a 1:40 dilution with 0.05% PBS-Tween using a Dynex Revelation automated absorbance microplate reader (Dynex Technologies; Chantilly, VA, USA). Plates were developed after 10 min using 3,3',5,5'-tetra-methylbenzidine core and orthophosphoric acid used as a stop solution (both TBS). Optical densities at 450 nm were measured using the Dynex Revelation. Results of ELISA for detection of anti-Spike antibodies in dried blood spot eluates have previously been shown to have almost perfect agreement with those performed on serum (Cohen's kappa = 0.83)¹⁸.

Sample size and statistical methods

Using the Stata powerlog programme, we estimated that a minimum sample size of 10,964 would be required to detect a difference of at least 2% in the proportion of exposed vs. unexposed participants experiencing a given binary outcome [equivalent to an odds ratio (OR) of 1.08], with 90% power, for a binary exposure with maximum variability (probability 0.50 changing to 0.52) and a moderate correlation (R2 = 0.4) with other variables in a logistic regression model, using a two-sided test and 5% significance. This post-vaccination symptom analysis was a pragmatic study including all participants meeting the inclusion criteria, with no sample size specified.

Logistic regression models were used to estimate odds ratios (ORs) and 95% confidence intervals (CIs) for associations between potential determinants of post-vaccination symptoms and their occurrence, first in a minimally adjusted, then fully adjusted models. Factors associating independently with each outcome at the P < 0.10 significance level were selected for inclusion in the multivariable model. Both the minimally adjusted and fully adjusted models were controlled for age (<30 years, 30 to <40 years, 40 to <50 years, 50 to <60 years, 60 to <70 years, and ≥70 years) and sex (male vs. female). Correction for multiple comparisons was not applied, on the grounds that we were testing a priori hypotheses for all risk factors investigated²⁰. Unpaired two-sided t tests were used to compare mean anti-S titres and mean number of days from the date of second vaccine dose to the date of blood sampling between participants who did vs. did not experience reactive symptoms. Pearson chi-square tests were used to compare proportions of participants reporting post-vaccination symfptoms by dose order and vaccine type. Linear regression was used in an exploratory analysis to evaluate determinants of the total number of systemic symptoms reported following a first dose of vaccine, and to test the association between frequency of reporting symptoms (never vs. after one vaccine dose only vs. after both vaccine doses) and mean postvaccination antibody titres.

COVID-19 severity was classified into two groups: 'asymptomatic' (non-hospitalised seropositive participants, who either did not report any symptoms of acute respiratory infection or whose symptoms were classified as having <50% probability of being due to COVID-19, using the symptom algorithm by Menni and colleagues²¹) and 'symptomatic' (hospitalised and nonhospitalised seropositive participants who reported symptoms of acute respiratory infection that were classified as having ≥50% probability of being due to COVID-19, using the symptom algorithm.)²¹.

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

DATA AVAILABILITY

The data that support the findings of this study are available from the corresponding, Professor Adrian Martineau, author upon request. Project name is COVIDENCE UK.

CODE AVAILABILITY

The code written to implement statistical analyses for this study is available from h.holt@qmul.ac.uk upon reasonable request.

Received: 25 May 2022; Accepted: 27 January 2023; Published online: 25 February 2023

REFERENCES

- 1. Johns Hopkins Coronavirus Resource Center. Coronavirus Resource Center. Available at https://coronavirus.jhu.edu2022.
- Sallam, M. COVID-19 vaccine hesitancy worldwide: a concise systematic review of vaccine acceptance rates. Vaccines 9, 160 (2021).
- Herve, C., Laupeze, B., Del Giudice, G., Didierlaurent, A. M. & Tavares Da Silva, F. The how's and what's of vaccine reactogenicity. *NPJ Vaccines.* 4, 39 (2019).
- Rief, W. Fear of adverse effects and COVID-19 vaccine hesitancy: recommendations of the treatment expectation expert group. JAMA Health Forum 2, e210804 (2021).
- Chapin-Bardales, J., Gee, J. & Myers, T. Reactogenicity following receipt of mRNA-Based COVID-19 Vaccines. J. Am. Med. Assoc. 325, 2201–2202 (2021). 06 01.
- Menni, C. et al. Vaccine side-effects and SARS-CoV-2 infection after vaccination in users of the COVID Symptom Study app in the UK: a prospective observational study. *Lancet Infect. Dis.* 21, 939–949 (2021). 07.
- Coggins, S. A. et al. Adverse effects and antibody titers in response to the BNT162b2 mRNA COVID-19 vaccine in a prospective study of healthcare workers. Open Forum Infect. Dis. 9, ofab575 (2022).
- Naaber, P. et al. Dynamics of antibody response to BNT162b2 vaccine after six months: a longitudinal prospective study. *Lancet Reg. Health Eur.* 10, 100208 (2021).
- Krammer, F. et al. Antibody responses in seropositive persons after a single dose of SARS-CoV-2 mRNA vaccine. N. Engl. J. Med. 384, 1372–1374 (2021).
- Hwang, Y. H. et al. Can reactogenicity predict immunogenicity after COVID-19 vaccination? Korean J. Intern. Med. 36, 1486–1491 (2021), 11.
- Bauernfeind, S. et al. Association between reactogenicity and immunogenicity after vaccination with BNT162b2. Vaccines (Basel). 9(Sep), 1089 (2021). 27.
- Howren, M. B. & Suls, J. The symptom perception hypothesis revised: depression and anxiety play different roles in concurrent and retrospective physical symptom reporting. J. Pers. Soc. Psychol. 100, 182–195 (2011).
- Calder, P. C. et al. Micronutrients to support vaccine immunogenicity and efficacy. Vaccines 10, 568 (2022).
- Dickinson A., MacKay D. Health habits and other characteristics of dietary supplement users: a review. (1475–2891 (Electronic)).
- Bedford, J. L. & Barr, S. I. Diets and selected lifestyle practices of self-defined adult vegetarians from a population-based sample suggest they are more 'health conscious'. Int. J. Behav. Nutr. Phys. Act. 2, 4–4 (2005).
- Talaei, M. et al. Determinants of pre-vaccination antibody responses to SARS-CoV-2: a population-based longitudinal study (COVIDENCE UK). BMC Med. 20, 87 (2022).
- Jolliffe, D. A. et al. Determinants of antibody responses to SARS-CoV-2 vaccines: population-based longitudinal study (COVIDENCE UK). *Vaccines (Basel)*. 10, 1601 (2022).
- Cook, A. M. et al. Validation of a combined ELISA to detect IgG, IgA and IgM antibody responses to SARS-CoV-2 in mild or moderate non-hospitalised patients. J. Immunol. Methods 494, 113046 (2021).
- Holt, H. et al. Risk factors for developing COVID-19: a population-based longitudinal study (COVIDENCE UK). *Thorax* 77, 900–912 (2021).
- Rothman, K. J. No adjustments are needed for multiple comparisons. *Epidemiology* 1(Jan), 43–46 (1990).
- Menni, C. et al. Real-time tracking of self-reported symptoms to predict potential COVID-19. Nat. Med. 26, 1037–1040 (2020).

ACKNOWLEDGEMENTS

We thank all participants of COVIDENCE UK, and the following organisations who supported study recruitment: Asthma UK, the British Heart Foundation, the British Lung Foundation, the British Obesity Society, Cancer Research UK, Diabetes UK, Future Publishing, Kidney Care UK, Kidney Wales, Mumsnet, the National Kidney Federation, the National Rheumatoid Arthritis Society, the North West London Health Research Register (DISCOVER), Primary Immunodeficiency UK, the Race Equality Foundation, SWM Health, the Terence Higgins Trust, and Vasculitis UK. MT was supported by a grant from Barts Charity (MGU0570). Funded by Barts Charity, Health Data Research UK. This study was supported by Barts Charity (refs MGU0459 and MGU0466), Pharma Nord, the Fischer Family Foundation, DSM Nutritional Products, the Exilarch's Foundation, the Karl R Pfleger Foundation, the AIM Foundation, Synergy Biologics, Cytoplan, the UK National Institute for Health and Care Research Clinical

Research Network (refs 52255 and 52257), the HDR UK BREATHE Hub, the UK Research and Innovation Industrial Strategy Challenge Fund (ref MC_PC_19004), Thornton & Ross, Warburtons, Hyphens Pharma, and a personal donation from Matthew Isaacs (a philanthropist without financial interests constituting a potential conflict). MT was supported by a grant from the Rosetrees Trust and The Bloom Foundation (ref M771) until May 2021 and has been supported by the Barts Charity since then (ref MGU0570). The funders had no role in the study design or in the collection, analysis, or interpretation of data, writing of the report, or the decision to submit the article for publication.

AUTHOR CONTRIBUTIONS

A.R.M. wrote the study protocol, with input from H.H., M.T. and S.O.S. H.H., M.T., J.S., S.F., G.A.D., R.A.L., C.J.G., F.K., A.S. and A.R.M. contributed to questionnaire development and design. HH co-ordinated and managed the study, with input from A.R.M., D.A.J., M.T., J.S., S.O.S. and S.M. H.H., J.S., A.R.M. and S.O.S. supported recruitment. S.F. and A.G.R. developed, validated and performed laboratory assays. M.T., H.H., M.G. and D.A.J. contributed to data management and coding medication data. H.H., M.T. and G.V. directly accessed and verified the data. Statistical analyses were done by H.H., with input from S.O.S., A.R.M. and G.V. H.H. and A.R.M. wrote the first draft of the report. All authors read and approved the final paper. H.H., M.T., D.A.J., S.O.S., A.R.M. and G.V. had full access to all data in the study, and H.H. had final responsibility for the decision to submit for publication.

COMPETING INTERESTS

J.S. declares receipt of payments from Reach plc for news stories written about recruitment to, and findings of, the COVIDENCE UK study. A.S. is a member of the Scottish Government Chief Medical Officer's COVID-19 Advisory Group and its Standing Committee on Pandemics. He is also a member of the UK Government's NERVTAG's Risk Stratification Subgroup. ARM declares receipt of funding in the last 36 months to support vitamin D research from the following companies who manufacture or sell vitamin D supplements: Pharma Nord Ltd, DSM Nutritional Products Ltd, Thornton & Ross Ltd and Hyphens Pharma Ltd, ARM also declares support for attending meetings from the following companies who manufacture or sell vitamin D supplements: Pharma Nord Ltd and Abiogen Pharma Ltd. ARM also declares participation on the Data and Safety Monitoring Board for the Chair, DSMB, VITALITY trial (Vitamin D for Adolescents with HIV to reduce musculoskeletal morbidity and immunopathology). ARM also declares unpaid work as a Programme Committee member for the Vitamin D Workshop. ARM also declares receipt of vitamin D capsules for clinical trial use from Pharma Nord Ltd, Synergy Biologics Ltd and Cytoplan Ltd. All other authors declare have no competing interests.

ADDITIONAL INFORMATION

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41541-023-00614-0.

Correspondence and requests for materials should be addressed to Hayley Holt.

Reprints and permission information is available at http://www.nature.com/ reprints

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http:// creativecommons.org/licenses/by/4.0/.

© The Author(s) 2023