Immunogenicity of COVID-19 vaccines in patients with hematologic malignancies: a systematic review and meta-analysis

Joanne S. K. Teh,^{1,2} Julien Coussement,^{1,2} Zoe C. F. Neoh,^{1,2} Tim Spelman,³ Smaro Lazarakis,⁴ Monica A. Slavin,^{1,2,5,6} and Benjamin W. Teh^{1,2,5}

¹Department of Infectious Diseases, ²National Centre for Infections in Cancer, and ³Department of Health Services Research, Peter MacCallum Cancer Centre, Melbourne, VIC, Australia; ⁴Health Sciences Library, Royal Melbourne Hospital, Parkville, VIC, Australia; ⁵Sir Peter MacCallum Department of Oncology, University of Melbourne, Parkville, VIC, Australia; and ⁶Victorian Infectious Diseases Service, Royal Melbourne Hospital, Melbourne, VIC, Australia

The objectives of this study were to assess the immunogenicity and safety of COVID-19 vaccines in patients with hematologic malignancies. A systematic review and meta-analysis of clinical studies of immune responses to COVID-19 vaccination stratified by underlying malignancy and published from January 1, 2021, to August 31, 2021, was conducted using MEDLINE, EMBASE, and Cochrane CENTRAL. Primary outcome was the rate of seropositivity after 2 doses of COVID-19 vaccine with rates of seropositivity after 1 dose, rates of positive neutralizing antibodies, cellular responses, and adverse events as secondary outcomes. Rates were pooled from single-arm studies while rates of seropositivity were compared against the rate in healthy controls for comparator studies using a random effects model and expressed as a pooled odds ratios with 95% confidence intervals. Forty-four studies (16 mixed group, 28 disease specific) with 7064 patients were included in the analysis (2331 after first dose, 4733 after second dose). Overall seropositivity rates were 62% to 66% after 2 doses of COVID-19 vaccine and 37% to 51% after 1 dose. The lowest seropositivity rate was 51% in patients with chronic lymphocytic leukemia and was highest in patients with acute leukemia (93%). After 2 doses, neutralizing antibody response rates were 57% to 60%, and cellular response rates were 40% to 75%. Active treatment, ongoing or recent treatment with targeted and CD-20 monoclonal antibody therapies within 12 months were associated with poor immune responses to COVID-19 vaccine. New approaches to prevention are urgently required to reduce COVID-19 infection morbidity and mortality in high-risk patient groups that respond poorly to COVID-19 vaccination.

Introduction

The COVID-19 pandemic caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) virus has resulted in global mortality of more than 3 million deaths. SARS-CoV-2 is likely to remain an endemic viral pathogen, with new variants continuously emerging. There is a significant burden of morbidity and mortality from COVID-19 infection in hematology patients. More than 80% of hematology patients require hospitalization, and up to 50% present with severe disease. Approximately 15% require admission to the intensive care unit, and mortality rates are high at 30% to 40%, depending on

Submitted 18 October 2021; accepted 17 November 2021; prepublished online on *Blood Advances* First Edition 1 December 2021; final version published online 28 March 2022. DOI 10.1182/bloodadvances.2021006333.

For publication-related data, please contact Benjamin W. Teh via email at ben.teh@ petermac.org.

The full-text version of this article contains a data supplement.

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underlying disease.²⁻⁷ Poor control of infection because of immune compromise lead to emergence of new variants which further complicate management.8

Vaccination is an effective public health measure for reducing the risk of infection and severe complications from COVID-19.9 However, patients with hematologic malignancies were excluded from the pivotal trials that preceded regulatory approvals of the currently recommended COVID-19 vaccines. 10-13 Understanding the impact of COVID-19 vaccines on patients with cancer is critical as social distancing restrictions are being eased in countries with relatively high rates of vaccination coverage. 14 Assessing immune response after vaccination is the basis of many vaccination studies in patients with hematologic malignancies and is often used as a surrogate marker of vaccine efficacy. ¹⁵⁻¹⁷ These studies are the basis for vaccination recommendations contained within international quidelines. 18,19 Therefore, this study was conducted to systematically review available data on the humoral and cellular immune responses to COVID-19 vaccination in patients with hematologic malignancies to build the evidence base for its utility.

The main objective of this systematic review was to assess the immunogenicity (ie, vaccine-induced immune response) of COVID-19 vaccines in patients with hematologic malignancies, stratified by underlying disease type. The secondary objective was to assess the safety of COVID-19 vaccines in the same patient groups.

Methods

This systematic review was preregistered with PROSPERO (CRD42021276851), and conducted in line with Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) auidelines.20

Types of studies and participants

Studies eligible for inclusion into the systematic review were those that investigated the vaccine-induced immunity in patients with hematologic malignancies who received at least 1 dose of COVID-19 vaccine. Randomized controlled trials (RCTs), guasi-RCTs, and observational studies were eligible for inclusion. All types of observational studies were included (eg, prospective and retrospective studies and studies with or without a control group). This systematic review evaluated all studies that reported on at least 1 of the following hematology patient groups: all patients with hematologic malignancies; patients with multiple myeloma (MM); patients with chronic lymphocytic leukemia (CLL); patients with lymphoma, Hodgkin lymphoma (HL), or non-Hodgkin lymphoma (NHL); patients with acute myeloid leukemia (AML) or acute lymphocytic leukemia (ALL); patients who are undergoing or have completed hematopoietic stem cell transplantation (HCT) or chimeric antigen receptor (CAR) T-cell therapy; and patients with myeloproliferative neoplasms (MPNs) and chronic myeloid leukemia (CML).

Types of intervention and evaluation

The type of intervention was the use of 1 or 2 doses of COVID-19 vaccine (of any type). Immune response to COVID-19 vaccination included humoral and cellular immune responses. For the purpose of this systematic review, humoral response consists of seropositivity which was defined by the SARS-CoV-2 spike/receptor binding domain-specific immunoglobulin G (IgG) level above the threshold

of detection for the assay used in each study. Rate of positive neutralizing antibody (nAb) response was defined by SARS-CoV-2-specific nAb level above the threshold established by the dedicated neutralization assay used in each study. A positive cellular response was defined by the appropriate increase in frequency of SARS-CoV-2-specific CD4⁺/CD8⁺ T cells after vaccination according to assays used. Where available, the immune response for each hematologic malignancy patient group was compared with a control group. If there was no comparator group, the immune response for each hematology patient group was described and summarized.

Outcome measures

The primary outcome was rates of seropositivity after 2 doses of COVID-19 vaccine stratified by disease groups. Secondary outcome measures were the rates of seropositivity after 1 dose of vaccine, rates of positive nAb response after 1 or 2 doses of COVID-19 vaccine, rates of positive cellular response after 1 or 2 doses of COVID-19 vaccine, and rates of systemic and/or local adverse events (AEs; whichever rate was higher) after 2 doses of COVID-19 vaccine.

Search strategy

Literature searches were conducted by an experienced research librarian (S.L.) using the Ovid MEDLINE, EMBASE, and Cochrane CENTRAL databases to identify relevant articles published in English from January 1, 2020, to August 31, 2021. A combination of subjects and keyword terms encompassing hematologic cancers, COVID-19, vaccines, and all their associated terms were used for the search. The terms used in our search included: hematologic neoplasms, leukemia, lymphoma, multiple myeloma, myeloproliferative disorders, myelodysplastic-myeloproliferative diseases, stem cell transplantation, bone marrow transplantation, chimeric antigen receptor therapy, COVID-19, SARS-CoV-2, vaccines, vaccination, immunize, immunization, BNT162b2, ChAdOx1, AZD 1222, mRNA-1273, Ad26, Ad5, and NVX-CoV2373. All word variations were searched, and medical subject headings were exploded. The detailed search strategy is summarized in the supplemental data.

Selection of studies

Studies were excluded from the systematic review if they did not measure or report immunogenicity after COVID-19 vaccination, if insufficient details were reported for the hematology patient groups in mixed-group studies, if there were <10 patients including case reports, if data were exclusive to pediatric patients younger than age 18 years, or if they were animal studies. Studies that were not peer reviewed and/or had not been published (eg, preprints, abstracts, and government reports) were excluded. Review articles and other publications without original data such as expert opinions, editorials, and consensus statements were also excluded.

Study eligibility was assessed by 2 independent reviewers (B.W.T., J.S.K.T.), and Covidence systematic review software (Veritas Health Innovation, Melbourne, Australia) was used to screen titles, abstracts, and full texts. Irrelevant reports were discarded, and the full texts of the other reports were accessed. Disagreements between the 2 main reviewers with respect to

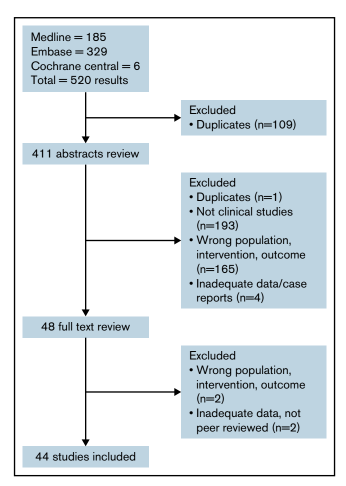


Figure 1. Flow diagram of studies identified, screened, and included.

study eligibility were resolved by discussion or via consultation with a third author.

Data extraction

For studies that fulfilled the inclusion criteria, data were extracted manually and independently by authors (J.S.K.T., J.C., B.W.T.) by using a predefined data extraction form. Extracted data elements included study design (enrollment, follow-up period, randomization/ allocation, laboratory analysis, predefined outcomes, adjusted analysis, funding source), participant information (inclusion criteria, number of participants, characteristics, disease, treatment), intervention (vaccine type, dose, schedule, comparator group), and outcomes (definitions, timing, AEs).

Assessment of risk of bias

Two review authors (Z.C.F.N., M.A.S.) independently assessed the risk of bias of each cohort study by using the Newcastle-Ottawa Scale.21 Each included study was assessed on the basis of 3 domains: (1) the selection of the study groups, (2) the comparability of the study groups, and (3) the ascertainment of outcome of interest. The rating system proposed by Sharmin et al²² was adopted; a good-quality study scored 3 or 4 stars in the selection domain, 1 or 2 stars in comparability domain, and 2 or 3 stars in the outcome domain. For single-arm studies, a good-quality study scored 3 stars in the selection domain and 2 or 3 stars in the outcome domain.

Measures of treatment effect

For each study, the number of patients who achieved seropositivity and the total number of patients who received vaccination was extracted and expressed as a proportion. For the meta-analysis, rate of response after Ad26 vaccine was analyzed at the same time point and therefore summarized as part of a 2-vaccine dose response. For each hematology disease group, the proportions from each study were pooled using the inverse variance method via the metaprop function in R version 4.1.1 (R Foundation for Statistical Computing, Vienna, Austria) and expressed as an overall proportion (response rate) with 95% confidence intervals (Cls).

For each study with a control group, the number of patients with seropositivity in each group and the total number of patients who received vaccination were extracted, expressed as a proportion with 95% Cls, and compared against each other. A meta-analysis was performed if there were 2 or more studies identified with a similar population and sufficient data for the outcomes of interest. The rate of seropositivity as a dichotomous outcome of interest across studies was summarized using a random effects model and expressed as a pooled odds ratios (ORs) with 95% Cls. Comparative analysis was performed using R 4.1.1 software.

Assessment of heterogeneity and missing data

Clinical and methodologic heterogeneity between included studies was assessed by comparing the key patient factors and study factors (vaccine type, type of assay used, threshold for positive response, duration of follow-up). Statistical heterogeneity was assessed by using the l^2 statistic and χ^2 with a P value \leq .05 considered significant for the presence of heterogeneity. Reason(s) for missing data were outlined if they were available. Because of time limitations, further information was not sought from the original or the corresponding author(s).

Subgroup analysis

Preplanned subgroup analysis was performed to evaluate the impact of active treatment (vs no active treatment), anti-CD-20 therapy in the last 12 months, including current CD-20 therapy (vs anti-CD-20 therapy 12 or more months ago), targeted therapy (defined as Bruton tyrosine kinase inhibitor [BTKi], BCL-2 antagonist venetoclax) in the last 12 months (vs no targeted therapy), timing of vaccination in relation to HCT (within 12 months vs longer than 12 months), and response by type of vaccine (BNT162b2 also known as Comirnaty or tozinameran [Pfizer] vs others, including messenger RNA-1273 or mRNA-1273 also known as Spikevax or elasomeran [Moderna], ChAdOx1 also known as Vaxzevria [AstraZeneca], or Ad26 [Janssen]). For subgroup analysis, only studies containing sufficient level of detail for the outcome of interest (eg, therapy <12 months) were included. For analysis by vaccine type, only studies reporting use and immune response of multiple vaccines types were included.

Results

Search results

A search of electronic databases yielded 520 results, and after duplicate results were excluded, 411 abstracts were reviewed. Forty-eight studies were identified for full-text review; subsequently, 44 studies fulfilled the inclusion criteria and underwent data

Table 1. Summary of study characteristics and outcomes for patients with hematologic malignancies by overall cohort and specific underlying disease

AEs	χ.	R	NR for hem	<u>κ</u> Ž	Ä	ŭ	χ.	N N	R	At least 1 AE; dose 1, 56 (9%) of 575; dose 2, 72 (13%) of 575	At least 1 AE; dose 1,88 (19%) of 154; dose 2, 55 (34%) of 163
Rate of positive nAb/cellular response	N.	χ χ	Ϋ́	Z.	22 days after first dose, ≥50%, 3 (5%) of 58 patients vs 50 (24%) of 213 controls	χ. Σ	ŭ Z	Y Z	Z Z	œ Z	Cetular response 14 days after second dose, 36 (53%) of 68
Seropositivity	Hem: 3-4 weeks after (72.%) of 2.5; (72.%) of 2.5; 3-4 weeks after second does, 17 (77%) of 2.2 20m. 17 (77%) of 2.9 (83%) of 66; (83%) of 66;	21 days after second dose, 36 (54%) of 67	21 days after first dose, 53 (43%) of 123; 28 days after second dose, 85 (71%) of 119	2-3 weeks after second dose, 34 (63%) of 54	22 days after first dose, ≥30%, 8 (14%) of 58 patients vs 114 (54%) of 213 controls	> 14days after second dose, 1088 (75%) of 1445	30-60 days after second dose, 235 (75%) of 315 vs 107 (99%) of 108 controls	28 days after second dose, 4 (40%) of 10	21 days after second dose, 41 (61%) of 67 vs 35 (100%) of 35 controls	7-21days after first dose, 481 (56%) of 857; NR for control group	≥3100 AU/nL threshold*: 28 days after first days after first 196; 14 days 196; 14 days doss, 91 (47%) doss, 91 (47%) oontrols
Analysis	SARS-CoV-2 spike, lgG Roche (=0.8 tul/mL = positive)	SARS-CoV-2 spike, IgG Beckman Couter (≥1.00 = positive)	SARS-CoV-2 spike, IgG Roche (≥0.8 IU/mL = positive)	SARS-CoV-2 spike, IgG Roche (≥0.8 IU/mL = positive)	nAb GenScript (=39% = positive; ≥50% = clinically relevant inhibition)	SARS-CoV-2 spike, IgG Roche (2.0.8 IU/mL = positive)	SARS-CoV-2 spike, IgG DaSorin (=12 AUmL = positive)	SARS-CoV-2 spike, IgG Abbott (≥50 AU/mL = positive)	SARS-CoV-2 spike, IgG in-house (OD 450 ≥3 = positive)	SARS-CoV-2 spike, IgG Abbort (≥50 AU/mL = positive)	SARS-CoV2 spike, lighthat though the Munt. Jigh Abbat (*50 Abbat
Vaccine type	BNT162b2, mRNA-1273, NR for hem	BNT162b2, mRNA-1273, 51%; 42%; unknown, 7%	BNT162b2	BNT162b2	BNT162b2, 76%; ChAdOx1, 24%	BNT162b2, 55%; mRNA-1273, 45%	BNT162b2	BNT162b2	BNT162b2, 46%; mRNA-1273, 54%	BNT162b2	BNT 162b2
M/F	NR for hem	35/32	NR for hem	32/22	28/30	574/871	2239200	NR for hem	36/31	404/453	117778
Median age, y (range or IQR)	NR for hem	71 (IQR, 65-77)	NR for hem	69 (IQR, 61-77)	75 (40-88)	68 (16-110)	70 (IQR, 61-77)	NR for hem	71 (24-90)	65 (IQR, 54-72)	69 (22-92)
No. of participants analyzed/ controls	25 106	67	123 34% myeloma, 38% L/ 9mphoma/WM, 28% AML/ MDS/MPNs 136	4	58 48% WM, 38% CLL, 14% NHL	1445.45% CLL, 55% NH-L, 55% NH, 15% MM, 45% acute leukena, 2% CML, 2% ONL, 2% ONL, 2% other	315 22% MPNs, 22% MPNs, 17% myeloma, 16% aggressive NNL, 11% cCLL, 7% CML, 5% HL, 5% acute leukema, 5% MDS	10	67 31% CLL, 63% NHL, 6% HL 35	857 88	195 27% mycloma, 22% NHL, 3% 15% ALL, 19% ALL, 9% ARPA, 7% 30 other, 7%
Study population/ comparator	Hem Solid tumor	Hem	Hem Solid tumor	Hem	Hem Control group	Hem	Hem Matched heality controls	Hem ≥80 y; health care worker controls ≥66 y (results NR)	Hem Health care worker controls	Hem Health care worker controls	Healthy controls
Type/location	Multicenter prospective cohort study Europe, North America	Single-center retrospective cohort study/ United States	Single-center prospective cohort study/ Austria	Single-center retrospective study/Israel	Single-center prospective cohort study/ Greece	Multicenter prospective cohort study/ United States	Single-center prospective cohort study/ Isnel	Single-center prospective cohort study/	Single-center prospective cohort study/ United States	Multicenter prospective cohort study/ Europe	Single-center retrospective cobort study/ France
Study	Mixed-group Addeo et al ⁶³	Agha et al ²⁴	Benda et al ²⁵	Cohen et al ²⁶	Gavriatopoulou et al ²⁷	Greenberger et al ²⁸	Herzog Tzarfati et a 20	lacano et al ³⁰	Jurgens et al ³¹	Maneikis et al ³²	Maland et al ³³

AEs, adverse events; ALL, acute lymphocytic leukemia; AML, acute myeloid leukemia; BAU, binding antibody unit; CLL, chronic lymphotic leukemia; COV, cut-off value; EC₅₀, median effective concentration; ET, essential thrombocythemia; F, female; GVHD, graft-versus-host disease; hem, hematology; HL, Hodgkin lymphoma; ICA, intracellular cytokine assay; ID₅₀, median infective dose; IEN-y, interferon-y; IL-2, interleukin-2; IOR, interquartile range; M, maltiple myeloma; ND, not determined; NE, not extractable; NHL, non-Hodgkin lymphoma; NR, not reported; OD, optical density; PBMC, peripheral blood mononuclear cell; PV, polycythemia vera; RBD, receptor-binding domain; SMM, smoldering multiple myeloma; WM, Waldenström macroglobulinemia.

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Table 1. (continued)

AEs	NE for hem patients	Σ Z	Reported with Referent patient patient numbers	œ Z	NR for hem	At least 1 AE; 90 (53%) of 161 vs 29 (55%) of 55 controls	N R
Rate of positive nAb/cellular response	Cellular response 72 days after 72 days after 14 (82%) of 18 vs 14 (82%) of 18 vs 14 (82%) of 18 vs 16 (95%) of 18 vs (83%) of 18 vs (83%) of 18 vs (75%) of 48 vs 3 (75%) of 48 vs 3 (75%) of 48 vs 3 (75%) of 48 vs	Æ	Æ	œ Z	χ.	ĽZ	NR
Seropositivity	21 days after first does if 18 (18%) of 34 or 44 vs 32	56 days after frst dose, 88 (39%) of 160	21 days after frist does, 9 (121%) of 422; mysloma. 26 (122%) of 50 (1	21-28 days after second dose, 64 (62%) of 102	>7 days after second dose, 56 (85%) of 66 vs 134 (98%) of 134 sold tumors; rate NR for controls	14-21 days after second dose, 133 (78%) of 171 vs 63 (98%) of 64 controls	≥21 days after first dose, 52 (56%) of 93
Analysis	SARS-CoV-2 spike, 196 [=20 Ecc, 196 [=20 Ecc, 196 [=20 Ecc, 196]= positive). SARS-CoV-2 SARS-CoV-1 Trolls secreting 18N-y and/or IL-2 Poyotkine. Secreting cells per 10 ⁸ Pells per 10 ⁸ Pe	SARS-COV-2 spike, Ig5 Abbort (signal/cutof ratio ≥:14 = postive)	SARS-CoV2 spike, IgC DisSonn (=15 AU/mL = positve)	SARS-CoV2 spike, IgG; range of commercial kits using their own threshold	SARS-CoV-2 spike, IgA About (≥50 AU/mL = positive)	SARS-CoV-2 spike, IgG Roche (≥0.8 IU/mL = positive)	SARS-Co-V-2 spike, IgG Ortho clinical (≥1 signal/cutoff = positive)
Vaccine type	BNTie2b2	BNT162b2, 60%; mRN-1273, 31%; Ad26, 7%; ND, 1%	BN7182b2	BNT162b2, 93%; mRNA-1273, 7%	BNT162b2, 58%, mRNA-1273, 31%, Az26, 10%, mRNA 10%, uNknown, 2%	BNT162b2	BNT162b2, 52%; ChAdOx1, 48%
M/F	NE for hem	86/74	Myelona: 23/19; MPN: 26/24	67/35	NR for hem	96/75	55/38
Median age, y (range or IQR)	73 (IGR, 6E-80)	72 (66-79)	73 (47-79)/ 70 (28-90)	76 (33-93)	NR for hem	70 (38-94)	67 (47-87)
No. of participants analyzed/ controls	56: 68% B-cell marganores, marganores, marganores, marganores, resultanguaroes, or acute leukenin, 5% other 5% other marganores, see the seekenin, 5% other marganores, see the seekening	160: 36% and aggressive and 21% independing physician in 15% plasma cell leukermia, 12% CML, 9% other lymphona 6% mychoria elekermia leukermia mychoria elekermia leukermia mychoria elekermia leukermia	96 Sef-% MPN 36 Gef-% MPN 36 Gef-% MPN	102: 45% Iymphoma, 22% myeloma, 13% MDS/ AML, 10% CLL, 10% MPNs	6689% lymphoid malignancies, 27% myeloid malignancies, 33% plasma 134 ell leukemia 26	171 64	83
Study population/ comparator	Hem Heath care worker controls	Hen	Hem Oldertage (SB 9) control group)	Hem	Hem Solid tumors Healthy controls	Myeloma Healthy volunteers	Myeloma
Type/location	Multicenter prospective control autoly/ United Kingdom	Single-center errospective cohort audy/ United States	Single-center prospective prospective study/ that)	Multicenter retrospective cohort study/ France	Single-center prospective and retraspective cohort study/ United States	Single-center prospective cohort study/ Israel	Single-center retrospective cohort study/ United Kingdom
Study	Monin et al ³⁴	Olika et al ³⁵	Pmpinelli et a ⁹⁶	Re et al ³⁷	Thakkar et al ³⁸	Myeloma-specific Avivi et al ⁹⁹	Bird et al ⁴⁰

AEs, adverse events; ALL, acute lymphocytic leukemia; AML, acute myeloid leukemia; BAU, binding antibody unit; CLL, chronic lymphotic leukemia; COV, cut-off value; EC₅₀, median effective concentration; ET, essential thrombocythemia; F, female; GVHD, graft-versus-host disease; hem, hematology; HL, Hodgkin lymphoma; ICA, intracellular cytokine assay; ID₅₀, median infective dose; IFN-y, interferon-y; IL-2, interleukin-2; IOR, interquartile range; M, maltiple myeloma; ND, not determined; NE, not extractable; NHL, non-Hodgkin lymphoma; NR, not reported; OD, optical density; PBMC, peripheral blood mononuclear cell; PV, polycythemia vera; RBD, receptor-binding domain; SMM, smoldering multiple myeloma; WM, Waldenström macroglobulinemia.

Table 1. (continued)

	AEs	N.	MR.	ŭ.	Ϋ́ Y	First dose of Prist dose of Prist dose of 18 (18 24), of 18 (18 24) of 21 (21 %) of 21 %) of 21 (21 %) of 21 %	N.	At least 1 AE; 151 (47%) of 331	¥	First dose, 5.2 (13%) of 167 local restation; 21 (13%) of 167 (167 systemic reaction. Second dose, 68 (34%) of 167 local reaction; 21 (22%) of 167 systemic reaction
Rate of positive nAb/cellular	response	Υ Z	N.	^Σ	≥50%: 22 days after first dose, 4 (8%) of 48 vs 21 (20%) of 102 controls	≥ 50%, day 22 after risk close 65 (FW) of 276 (FW) of 226 controls	œ Z	nAb 14-21 days after second dass, 27 (60%) of 45	<u>۳</u> 2	Σ.
	Seropositivity	21 days after first dose, 17 (23%) of 74	>21 days after first dose, 17 (61%) of 28	14.21 days after first case, 20 (21%) of 96, 14.21 days after first second days. 30 (10%) of 96, 14.21 days second days. 46 (10%) of 100% of 1	≥30%: 22 days after first dose, of 12 (25%) of 48 vs 57 (55%) of 102 controls	=30%; day 22 after foot of 17 1.65 (8.4%) vs 1.65 (8.4%) vs 1.65 (8.4%) vs 1.65 (8.4%) vs day 20 after (184 day 20 after (184 day 20 after (184) of 276 vs 204 (90%) of 226 controls	51 days after second dose, 219 (84%) of 260 vs 67 (100%) of 67 controls	14-21 days after second dass, 160 (43%) of 373	14-21 days after second dose, 25 (54%) of 46	14-21 days after second dose, 66 (40%), of 167 patients vs 167 patients vs controls
	Analysis	SARS-CoV-2 spike, IgG DiaSorin (≥34 AU/mL = positive)	SARS-CoV-2 spike, IgG Abbort (COV ≥50 = positive)	SARS-CoV 2 spike, fig. of a rhouse (190 ar house (190 ar house) (200 U/mt and response) circuit response)	SARS-CoV-2 nAb GenScript (≥30% = positive; ≥50% = clinically relevant)	SARS CoV 2 nAb Cor Society (Core Section of Core Section of Co	SARS-CoV-2 spike, IgG in-house (≥5 AU/mL = positive)	SARS-COV.2 spike, (g. DiaSorn (als AUmt.) positive), About (>50 Umt.) positive), inchouse (>1,1	SARS-CoV-2 spike, IgG MaGLUMI (≥1.1 = positive)	SARS-CoV 2 spike, gaples, gapl
	Vaccine type	BNT162b2, 77%; ChAdOx1, 23%	BNT162b2, 42%; ChAdOx1, 58%	BNT162b2, 50%; mRNA-1273, 50%.	BNT162b2	BMT16262, 78%; ChAcOxt, 22%	BNT162b2, 69%, mRNA-1273, 27%; unknown, 4%	BNT162b2	BNT162b2	Herishanu et al ¹⁴ Single-center CLL patients 167 71 (63-76) 112/56 BMT162b2 SARS-CoV-2 spike, 14-21 days after NR prospective Age and sex 52 (50%) of Foote second dose, 66 (40%) of Foote school dose, 67 (40%) of Foote school dose, 68 (40%) of Foote school dose, 68 (40%) of Foote school dose, 68 (40%) of Foote school dose, 69 (40%) of Foote school dose, 60 (40%) of Foo
	M/F	49/33	67/42	61/42	29/19	151/126	185/135	222/151	29/17	112/65
Median age, y	(range or IQR)	68 (40-85)	89	(68-58) 89	83 (59-92)	74 (62-80)	68 (38-93)	70 (40-89)	Ľ	71 (63-76)
No. of participants analyzed/	controls	82	105-28 patients sampled	2 - 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2 -	48 102	7.7% 7.7% myelona, 1.5% eMA 9% 2.26 Md 5.9%	320 260 sampled 67	373	46	52
Study population/	comparator	Myeloma	Myeloma	Myeloma Helithy Comnois Pre-COVID-19 controls	Myeloma Matched controls	Matched controls	Myeloma Matched health care worker controls	CLL patients	CLL patients	OLL patients Age and sec- matched controls
	Type/location	Single-center prospective cohort study/ Germany	Multicenter web- based prospec- tive cohort study/ United Kingdom	Single-center prospective prospective cohort stuly/ United States	Single-center prospective cohort study/ Greece	Single-center propercive prince of study/ Greece	Single-center prospective and retrospective cohort study/ United States	Multicenter prospective cohort study/ Israel	Single-center prospective cohort study/ Italy	Single-center prospective prospective cohort study/ Isanel
	Study	Ghandili et al ⁴¹	Ramasamy et al ⁴²	Stampfer et a ⁴⁴³	Terpos et al ⁴⁴	Terpos et al ⁴⁵	Van Oekelen et al⁴e	CLL-specific Benjamini et al ⁴⁷	Del Poeta et al⁴ ⁸	Herishanu et al ¹⁹

Abs, adverse events; ALL, acute lymphocytic leukemia; AML, acute myeloid leukemia; BAU, binding antibody unit; CLL, chronic lymphotic leukemia; COV, cut-off value; EC₅₀, median effective concentration; ET, essential monocolons, and the match of the m

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Table 1. (continued)

AEs	NR	Z.	A.		Ν.	X.	Ä	At least 1 AE; 60 (15%) of (15%) of (18%) of (18%) of (18 local AE; 23 (20%) of 118 systemic		č.	뜻
Rate of positive nAb/cellular response	N.	œ Z	Z.		œ	æ Z	χ Z	χ Σ		<u>압</u>	Σ Z
Seropositivity	43 days after first close, servin 22 (44%) arm 22 (44%) arm 22 (44%) are placed blood for serving the days controls; 18 days after second close, a	21 days after second dose, 23 (52%) of 44	22 days after second dose, 49 (58%) of 84		14-56 days after complete of watchiston of watchiston, vaccination, va	28 days after second dose, 83 (51%) of 162	14 days after first close, 32 (£4%) of 56 patients vs 66 (100%) of 66 controls; 14-28 days after second close, 61 (711%) of 86 patients vs 85 (100%) of 86 controls;	14-21 days after second dose, 73 (49%) of 149 vs 64 (99%) of 148 vs 64 (99%) of 16 controls; 38 (49%) of 80 indohen NHL; 34 (49%) of 89 aggressive NHL		42 days after first dose, 21 (38%) of 55	28 days after second dose, 69 (78%) of 88; >41 60 AU/mL 28 days after second dose, 82 (59%) of 88
Analysis	SARS-CoV-2 spike, lgG Rochie (≈0.8 IU/mL positive), dried blood sampling Roche (ratio ≥1.0 = positive)	SARS-CoV-2 spike, lgG DiaSorin (≥15 AU/mL = positive)	SARS-CoV-2 spike, lgG Abbott (≥50 UmL = positive); SARS- CoV-2 RBD IgG (>1.1 = positive)		SARS-CoV-2 spike, lgG Berkad (27.1.0 = positive)	SARS-CoV-2 spike, lgG Abbott (≥50 IU/mL = positive)	SARS-CoV-2 spike, lgG Meso Scale Discovery (C-CoE BAUML = positive); RBD lgG (C-C-73 BAUML = positive)	SARS-CoV-2 spike, gG Roche (GC0 NU/mL = positive)		SARS-CoV-2 spike, IgG Ortho clinical (≥1 signal/cutoff = positive)	SARS-CoV-2 spike, IgG Abbott (>21 AU/m. = positive; >4160 AU/m. = neutralization)
Vaccine type	BNT16282, 52%; ChArod1, 48%	BNT162b2, 57%; mRNA-1273, 43%	BNT162b2		BNT16262, 47%; mRAN-1273, 52%; Ad26, 1%;	BNT162b2	BNT162b2, ChAdOx1	BNT162b2		BNT162b2, 38%; ChAdOx1, 62%	BNT162b2
M/F	159/140	23/21	53/29		45/41	89/73	81/48	88/61		34/21	47/41
Median age, y (range or IQR)	69 (IOR, 63.74)	71 (37-89)	69 (44-87)		70 (35-91)	65 (52-73)	69 (IOR, 57.74)	64 (20-92)		50 (18-73)	ŭ
No. of participants analyzed/ controls	299 93	44	48		96 147 154	162: 88% NHL, 12% HL	129 recruited, 119 analyzed / 150:68% incloent B-cell NHL, 29% aggressive B-cell NHL, 10% HL, 3% other	149 53% indolent NHL, 47% aggressive NHL 65		OD OD	98
Study population/ comparator	CLL patents Healthy agen-matched controls	CLL patients	CLL patients		B-cel lymphona Nursing hone controls Health care worker controls	Lymphoma	Lymphoma Healthy controls	Lymphoma, B-cell NHL Healthy controls		Allo-HCT >3 months	Allo-HCT
Type/location	Single-center prospective cohort study/ United Kingdom	Single-center retrospective cohort study United States	Multicenter prospective observation stuch/		Single-center prospective conbort study/ United States	Multicenter prospective cohort study/ Israel	Multicenter prospective prospective cohort study interim analysis/ United Kingdom	Single-center prospective cohort study/ Isnel		Single-center retrospective cohort study/ United Kingdom	Single-center retrospective cohort study/ France
Study	Pany et al ⁴⁰	Roeker et al ^{5 1}	Tadmor et al ⁶²	Lymphoma- specific	Ghione et al ⁵³	Gurion et al ⁵⁴	Lim et al ⁶⁶	Perry et al ⁵⁶	HCT-specific	Easdale et al ⁵⁷	Redjoul et al ⁶⁶

AEs, adverse events; ALL, acute lymphocytic leukemia; AML, acute myeloid leukemia; BAU, binding antibody unit; CLL, chronic lymphotic leukemia; COV, cut-off value; EC₅₀, median effective concentration; ET, essential thrombocythemia; F, female; GVHD, graft-versus-host disease; hem, hematology; HL, Hodgkin lymphoma; ICA, intracellular cytokine assay; ID₅₀, median infective dose; IFN-y, interferon-y; IL-2, interleukin-2; IQR, interquartile range; M, male; MF, myelofibrosis; MGUS, monoclonal gammopathy of unknown significance; MM, multiple myeloma; ND, not determined; NE, not extractable; NHL, non-Hodgkin lymphoma; NR, not reported; OD, optical density; PBMC, peripheral blood mononuclear cell; PV, polycythemia vera; RBD, receptor-binding domain; SMM, smoldering multiple myeloma; WM, Waldenström macroglobulinemia.

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Table 1. (continued)

Study	Type/location	Study population/ comparator	No. of participants analyzed/ controls	Median age, y (range or IQR)	M/F	Vaccine type	Analysis	Seropositivity	Rate of positive nAb/cellular response	AEs
Ram et al ⁹⁹	Single-center prospective cohort study/ Israel	Allo-HCT and CAR T cells >3 months	80: 83% allo-HCT, 17% CAR T cells	65 (23-83)	44/37	BNT162b2	SARS-CoV2 spike, IgG Roahe (=0.08 U/mL = (=0.08 U/mL = CoV2-especific cells ELISPOT (RN, IL-2) (4 spots per well = spots per well =	7-14 days after second close, 47 (75%), or (75%), or (86%) of 14 CAR T cells	Cellular responses 2-14 days after scored dose, 7 (19%), of 19% of 8 (alc-HCT, 6 (50%), of 12 CAR T cells	At least 1 AE; first doce, 11 (14%) of 80, 3 (4%) of 80 (2VHD; second doce, 18 (24%) of 74 (3VHD) of 74 (3(W)
Dhakal et al ⁶⁰	Single-center retrospective cohort stud/ United States	Auto-HCT, allo-HCT, CAR T cells	130: 45 auto-HCT, 71 allo-HCT, 14 CAR T cells	Auto-HCT, 66 (45- 75); alo-HCT, 6 (25-77); CAR T cels, NR	뜻	BNT162b2, 59%; mRNA-1273, 38%; Ad26, 5%.		14 days after completion of vacconation, 27 (60%) of 45 auto-HCT; 49 (38%) of 71 allo-HCT; 3 (21%) of 14 CAR T cells	Σ	œ Z
MPN- and CML- specific										
Caocci et al ⁶ 1	Single-center prospective cohort study/ Italy	MPN	20: 65% MF, 30% ET, 5% PV	66 (48-82)	R	BNT162b2	SARS-CoV-2 spike, IgG DiaSorin (≥15 AU/mL = positive)	42 days after second dose, 13 (65%) of 20	ĽZ	N.
Chowdhury et al ⁶²	Single-center retrospective cohort/ United Kingdom	CML and MPN Health care workers age >60 years	59 232	62 (IQR, 52-73)	27/32	BNT162b2, 37%; ChAdOx1, 63%	SARS-CoV-2 spike, lgG.Abbott (≥50 AU/mL = positive)	≥2 weeks after first dose, 34 (57%) of 59 vs 224 (97%) of 232 controls	E.	œ Z
Gugielmelli et al ⁶³	Single-center prospective cohort study/ Italy	MPN Healthy controls	30 43% MF, 33% PV, 23% ET 14	NR overall	10/20	BNT162b2, 83%; mRNA-1273, 17%	SARS-CoV-2 spike/ RBD igG (NS)	21 to 28 days after first dose, 18 (60%) of 30 vs 14 (100%) of 14 controls	nAb, 21-28 days after first dose, 13 (43%) of 30 vs 14 (100%) of 14 controls	Z Z
Harrington et al ^{e4}	Single-center prospective orbot study/ United Kingdom/	CML	9	45 (23-74)	12/4	BNT1 82b.2	SARS-CoV2 spke, igd inhouse (igd inhouse (igs postwe); SARS-Downer); SARS-CoV2 reutralize (ii) inhouse (iii) inhouse (iii) inhouse (iii) inhouse (iii) inhouse (iii) inhouse inmesse inhouse (iii) inhouse inmesse inpostive)	21 days after first dose, 14 (89%) of 16	nAb 21 days after met doss, 6 (98%) of 16; cellular response, 14 (80%) of 15	Local AEs, 8 (50%) of 16 systemic AEs, 9 (56%) of 16
Harrington et al ¹⁰⁵	Single-center prospective cohort study/ United Kingdom	NdM	2	58 (36-72)	7721	BNT1 82b.2	SARS-CoV2 spike, igd inhouse (1.25 = positive); SARS-CoV2 reutralizing inhouse (1050 = COV2 T cells (COV2 T cells (COV2 T cells (COV2 T cells (COV2 T cells increase = positive); SARS-COV2 T cells (Mere (idd increase = positive)); SARS-COV2 T cells (COV2 T cells increase = positive); SARS-COV2 T cells (COV2 T cells in	21 days after first dose, 16 (76%) of 21	nAb 21 days after first close, 18 (86%) of 21; cludar response (CD4), 15 (75%) of 20	At least 1 AE; 12 (57%) of 21 (67%) of 21 (62%) of (48%) of 21 systemic
Pimpinelli et al ⁶⁶	Single-center prospective cohort study/ kaly	MPN	42: 40% ET, 36% PV, 24% MF	72 (52-82)	20/22	BNT162b2	SARS-CoV-2 spike, IgG DiaSorin (≥15 AU/mL = positive)	21 days after first dose, 23 (55%) of 42; 14 days after second dose, 36 (86%) of 42	A.N.	N N

AEs, adverse events; ALL, acute lymphocytic leukemia; AML, acute myeloid leukemia; BAU, binding antibody unit; CLL, chronic lymphotic leukemia; COV, cut-off value; EC₅₀, median effective concentration; ET, essential thrombocythemia; F, female; GVHD, graft-versus-host disease; hem, hematology; HL, Hodgkin lymphoma; ICA, intracellular cytokine assay; ID₅₀, median infective dose; IFN-γ, interferon-γ; IL-2, interleukin-2; IOR, interquartile range; M, male; MF, myelofibrosis; MGUS, monoclonal gammopathy of unknown significance; MM, multiple myeloma; ND, not determined; NE, nor extractable; NHL, non-Hodgkin lymphoma; NR, not reported; OD, optical density; PBMC, peripheral blood mononuclear cell; PV, polycythemia vera; RBD, receptor-binding domain; SMM, smoldering multiple myeloma; WM, Waldenström macroglobulinemia.
*Excluded from meta-analysis.

Table 2. Summary of pooled seropositivity rates for patients with hematologic malignancies by underlying disease type and number of vaccine doses

			•	•									Ì
	,	Single-arm studies	dies	Malignancy arm of cor	nparator st	udies (Malignancy arm of comparator studies Control arm of comparator studies	rator stu		Intervention vs control cohort	ontro	coho	Ŧ
	No. of patients	Pooled response rate (95% CI)	e f² (%) P	Pooled response rate (95% CI)	f (%)	ام	Pooled response rate (95% CI)	f² (%)	 a	OR (95% CI)	ρí	f² (%)	۵
Overall: Patients with hematologic malignancies	es												
After second dose	4733	0.62 (0.55-0.70)	92 <.01	1 0.66 (0.57-0.75)	93	<.01	0.99 (0.97-1.00)	53	0.01 0	0.04 (0.02-0.08)	<.01	> 02	<.01
After first dose	2331	0.51 (0.38-0.64)	92 <.01	1 0.37 (0.23-0.51)	06	<.01	0.78 (0.62-0.95)	86	<.01 0	0.10 (0.04-0.29)	<.01	> 98	<.01
Myeloma													
After second dose	1218	0.80 (0.64-0.95)	85 <.01	1 0.76 (0.70-0.82)	06	<.01	0.98 (0.95-1.00)	71	<.01	0.09 (0.03-0.29)	<.01	49	90:
After first dose	685	0.43 (0.18-0.68)	91 <.01	1 0.29 (0.09-0.48)	81	<.01	0.64 (0.42-0.87)	9/	<.01 0	0.23 (0.05-0.99)	.05	49	Ξ
CIL													
After second dose	1446	0.51 (0.37-0.65)	91 <.01	1 0.51 (0.34-0.68)	26	90.	1.00 (0.99-1.00)	0	.07 0	0.01 (0.01-0.03)	<.01	0	o.
After first dose	111	Single study		0.18 (0.00-1.00)	88	<.01	0.92 (0.52-1.00)	0	.32 0	0.03 (0.00-0.80)	.05	0	.60
Lymphoma													
After second dose	1227	0.52 (0.28-0.75)	97 <.01	1 0.55 (0.35-0.76)	84	<.01	0.99 (0.98-1.00)	0	.72 0	0.02 (0.01-0.02)	<.01	0	1.0
After first dose	69	No studies available		0.33 (0.00-1.00)	66	<.01	0.95 (0.09-1.00)	71	90.	0.01 (0.0-1.24)	.05	0	.7
HSCT and cellular therapies													
After second dose	401	0.61 (0.42-0.80)	88 .05	Single study			Single study			Single study			
After first dose	21	Single study		No studies available			No studies available		Š	No studies available			
Acute leukemia and MDS													
After second dose	113	0.93 (0.80-1.00)	29 .24	. Single study			Single study			Single study			
After first dose	13	Single study		Single study			Single study			Single study			
MPN and CML													
After second dose	281	0.88 (0.72-1.00)	70 .01	0.87 (0.81-0.92)	0	.83	0.99 (0.98-1.00)	0	06:	0.07 (0.0-1.55)	90.	0	.77
After first dose	222	0.71 (0.30-1.00)	76 .01	0.54 (0.37-0.71)	0	.46	0.85 (0.51-1.00)	06	<.01 0	0.13 (0.01-1.71)	60:	88	<.01
Second of MO commonly of the description of the common of	oin old Maria	TOOH . cienceling bigle.	cito; o a o to a o d	solder the contract of	OCW.	Jan. holo	an MON company of the	itonofilona olon	or in one	9			

AEs, adverse events; CLL, chronic lymphotic leukemia; CML, chronic myeloid leukemia; HSCT, hematopoietic stem cell transplantation; MDS, myelodysplastic syndrome; MPN, myeloproliferative neoplasms.

extraction (Figure 1). Overall, there were 44 studies of COVID-19 vaccination in patients with hematologic malignancies. Sixteen studies²³⁻³⁸ involved mixed group of patients with various underlying hematologic malignancies, and 28 studies focused on specific malignancies and treatments: 8 myeloma studies,³⁹⁻⁴⁶ 6 CLL studies,⁴⁷⁻⁵² 4 lymphoma studies,⁵³⁻⁵⁶ 4 studies of HCT^{57,58} and chimeric antigen receptor (CAR) T-cell therapy,^{59,60} and 6 MPN studies.⁶¹⁻⁶⁶ A total of 7064 patients were included in the analysis (2331 after first dose and 4733 after second dose). Characteristics of patients and primary and secondary outcomes in the included studies are summarized in Table 1.

Risk of bias and quality assessment

Overall, 23 studies (52%) were evaluated as being of good or fair quality (good, 11; fair, 12) with low risk of bias, whereas the remaining 21 studies were rated as poor quality (high risk of bias) primarily because of lack of details regarding patient selection, demonstration that outcome of interest was not present at the start of study, comparability of cohorts (design or analysis), and duration of follow-up. Risk of bias and quality assessment of studies are summarized in supplemental Table 1.

Seropositivity rates

There were 44 studies measuring humoral immune responses in patients with hematologic malignancies. The majority of studies evaluated the use of BNT162b2 and used SARS-CoV-2 spike-specific IgG levels above the threshold of detection (seropositivity). Twenty-three studies (52%) reported sufficient data on the spike/receptor binding domain IgG antibody levels (median value or geometric mean titer) after 1 dose (6 studies), 27,35,37,41,62,63 2 doses (13 studies), 24,26,29,30,38,39,46,48,49,55,59-61 and after both 1 and 2 doses (4 studies). Antibody levels achieved were lower compared with those in control cohorts after 1 and 2 doses of COVID-19 vaccine. Antibody levels between doses was 5-fold to 15-fold, whereas the fourth study by Parry et al 50 reported a 132-fold increase between the first and second doses in 12 of 299 patients with CLL.

Several studies attempted to establish potential thresholds for clinical protection with higher antibody levels than the seropositivity threshold used. Malard et al³³ correlated SARS-CoV-2 spike IgG levels with nAb levels, and ≥3100 arbitrary units (AU)/mL on the Abbott assav was predictive of an nAb level ≥30%. Using this threshold, the rate of humoral response was 47% in hematology patients compared with 97% in healthy controls after a second dose of the BNT162b2 vaccine.33 Pimpinelli et al36 also evaluated humoral responses using a higher threshold of ≥80 AU/mL (Dia-Sorin, Saluggia, Italy) and noted response rates of 55% for patients with myeloma and 84% for patients with MPNs compared with 97% for healthy controls. Other studies correlated SARS-CoV-2 IgG levels with plague reduction neutralization tests or with levels seen in clinical vaccination trials. Stampfer et al 43 defined >250 IU/mL (in-house assay) as a clinically relevant response that was attained by 45% of patients with myeloma after 2 doses of a COVID-19 vaccine. For HCT patients, Redjoul et al⁵⁸ used a threshold of 4160 AU/mL (Abbott Laboratories, Wiesbaden, Germany), and 59% of patients achieved this response after 2 doses.

Overall pooled seropositivity rates in patients with hematologic malignancies were 62% in single-arm studies and 66% in comparator studies after 2 doses of COVID-19 vaccination. After a single dose, the pooled seropositivity rates were 51% in single-arm studies and 37% in comparator studies. Compared with healthy or older matched controls, patients with hematologic malignancies were less likely to achieve seropositivity with ORs of 0.04 (95% Cl, 0.02-0.08; P < .01) after 2 doses (Figure 2A) and 0.10 (95% Cl, 0.04-0.29; P < .01) after a single dose (Figure 2B) of COVID-19 vaccine. Heterogeneity was 70% and 86%, respectively (P < .01). Pooled seropositivity rates by disease type and vaccine doses are summarized in Table 2 and Figure 3. Overall results were graded as having moderate quality due to statistical and clinical heterogeneity and the proportion of studies with a high risk of bias (48%).

Patients with myeloma. There were 8 dedicated studies and 9 other studies that reported immune response rates in patients with myeloma (supplemental Table 2). For the 2 studies by Terpos et al, 44,45 only nAbs at 2 levels were reported; a threshold of ≥30% was used to define seropositivity and a threshold of ≥50% was used to define rate of positive nAbs. There were 1218 patients with myeloma who received 2 doses of vaccine, and seropositivity rates were 76% to 80%. Seropositivity rates were 29% to 43% after 1 dose of COVID-19 vaccine in 685 patients. The OR for achieving seropositivity was 0.09 (95% CI, 0.03-0.29; P < .01) in patients with myeloma compared with those in the healthy control group after 2 vaccine doses and 0.23 (95% CI, 0.05-0.99; P = .05) after 1 dose (Table 2).

Patients with CLL. A total of 1557 patients with CLL in 12 studies (6 CLL specific, 6 hematology) were included in this review (supplemental Table 3). Pooled seropositivity rates were 51% in 1446 patients with CLL after 2 doses of COVID-19 vaccine. Rate of seropositivity was 18% after 1 dose of COVID-19 vaccine, and it was 37% in a single-arm study by Ollila et al. ORs for seropositivity were 0.01 (95% Cl, 0.01-0.03; P < .01) for patients with CLL after 2 doses and 0.03 (95% Cl, 0.00-0.80; P = .05) after 1 dose of vaccine compared with that for healthy controls.

Patients with lymphoma. For 1296 patients with lymphoma across 11 studies (supplemental Table 4), pooled seropositivity rates were 52% to 55% after 2 doses and 33% after 1 dose of COVID-19 vaccine. ORs for achieving seropositivity were 0.01 to 0.02 for patients with lymphoma compared with a healthy cohort (Table 2).

Patients after HCT and CAR T-cell therapy. A total of 6 studies included in this review reported on immune responses after HCT and CAR T-cell therapy (supplemental Table 5). Two studies involved patients who received an allogeneic HCT (allo-HCT)^{57,58}: One study included patients who received an allo-HCT and those who received CAR T-cell therapy,⁵⁹ and the other included all patients who received HCTs and those who received CAR T-cell therapy.⁶⁰ Two studies were larger studies of hematology patients with subsets of patients treated with HCT or CAR T cells.^{28,29} There were 422 patients of whom 401 received 2 doses of COVID-19 vaccine and the remainder received 1 dose. Pooled seropositivity rate was 61% after 2 doses of COVID-19 vaccine. The seropositivity rate was 74% after allo-HCT and 31% after treatment with CAR

Table 3. Summary of pooled rates of positive nAb, cellular responses, and AEs

	Positive nAb response		Positive cellular response		Atleas	Atleast 1 AE rate		
	Pooled response rate	95% CI	95% CI Pooled response rate	95% CI	Pooled response rate 95% CI I^2 (%)	95% CI	f (%)	Ь
Overall: Patients v	Overall: Patients with hematologic malignancies							
After second dose	After second dose Single arm: 0.60 (single study)		Single arm: 0.40	0.00-1.00	0.36	0.24-0.48	62	<.01
	Malignancy arm of comparator studies: 0.57 (single study)		Malignancy arm of comparator studies: 0.75 (single study)					
After first dose	Single arm: 0.63	0.00-1.00	0.00-1.00 Single arm: 0.86	0.00-1.00	0.39	0.18-0.60	26	<0.01
	Malignancy arm of comparator studies: 0.18	0.00-0.44	0.00-0.44 Malignancy arm of comparator studies: 0.33 (single study)					

T cells. In a single study with a healthy control cohort, seropositivity rate was 81% in patients who received an autologous HCT (auto-HCT) compared with 99% in an age-matched group without haematologic malignancy.²⁹ In the only study of immune response after 1 dose of COVID-19 vaccine, the rate of seropositivity was 38% in patients who received an allo-HCT.57

Patients with acute leukemia or myelodysplastic syndrome (MDS). There were no dedicated studies of COVID-19 vaccination in patients with acute leukemia. A subset of 126 patients with acute leukemia or MDS were reported in 6 studies (supplemental Table 6), and the seropositivity rate was 93% after 2 doses of COVID-19 vaccine. In a single study, the seropositivity rate was 80% in patients with acute leukemia and 94% in patients with MDS compared with 99% in the control group.²⁹ No patients with acute leukemia mounted an immune response after a single vaccine dose compared with 86% of controls.34

Patients with MPNs or CML. Of 12 studies (supplemental Table 7) encompassing 503 patients with MPNs or CML (281 after 2 doses, 222 after a single dose), rates of seropositivity were 87% to 88% after 2 doses and 54% to 71% after 1 dose of vaccine. Compared with a healthy patient cohort, the OR was 0.07 (95% Cl, 0.0-1.55; P = .06) for patients with MPNs or CML achieving seropositivity with 2 doses of COVID-19 vaccine and 0.13 (95% Cl, 0.01-1.71; P = .09) after a single dose.

Rates of nAb and cellular response and AEs

Only 7 studies (16%) reported nAb responses and 5 studies (11%) reported cellular responses. After 2 doses of COVID-19 vaccine, 57% of patients with myeloma achieved a positive nAb response compared with 81% of the control group, whereas 60% of patients with CLL achieved a positive nAb response in a subpopulation of a single-arm study. 45,47 After 1 dose of COVID-19 vaccine, the overall pooled rate for a positive nAb response was 18% for all patients with hematologic malignancies (Table 3). The pooled rate of 2 single-arm studies of patients with MPNs was 63%. After COVID-19 vaccination, the rate of achieving a positive cellular response was 40% to 75% for all patients after 2 doses and 33% to 86% after a single dose. In a single study, the cellular response rates after 2 doses of COVID-19 vaccine were 19% for patients who received treatment with allo-HCT and 50% for those who received treatment with CAR T cells.⁵⁹ In patients with MPNs, this rate was 86% after 1 dose. 64,65

In 10 studies (22%), at least 1 systemic or local AE was reported. Overall, the pooled rate of at least 1 AE was 36% after 2 doses and 39% after a single dose (Table 3). Overall, local and systemic AEs were mild (grade 1 to 2) except for a single study in which systemic AEs rates were 2%. 32,33,39,45,47,49,56,64,65 In patients who received an HCT, a 5% rate of exacerbation of graft-versus-host disease (grade 1 to 2) and grade 3 to 4 neutropenia or thrombocytopenia (self-resolved) was noted.⁵⁹ Most commonly reported AEs were injection site pain, fatigue, myalgia, headache, and fever. 32,33,39,45,47,49,56,59,64,65

Subgroup analysis

Study characteristics and outcomes of studies included in subgroup analyses are summarized in supplemental Tables 8-12. Receiving 2 doses of vaccine during active therapy was associated with

Table 4. Summary of subgroup analysis by active treatment, timing of CD20 antibody therapy, HCT, receipt of targeted therapies, and type of COVID-19 vaccine

	Intervention arm			Control arm			Intervention vs cont	trol c	ohort	
	Pooled response rate (95% CI)	I ² (%)	P	Pooled response rate (95% CI)	I ² (%)	P	OR (95% CI) Heterogeneity	P	/ 2(%)	P
Active therapy vs no	active therapy									
After second dose	0.28 (0.08-0.48)	83	<.01	0.62 (0.39-0.86)	94	<.01	0.21 (0.06-0.67)	.02	75	<.01
After first dose	0.42 (0.09-0.75)	49	.08	0.81 (0.66-0.95)	0	.62	0.19 (0.03-1.19)	.06	24	.27
CD-20 antibody ther	apy within 12 months vs CD-20 th	erapy 1	2 or mo	ore months						
After second dose	0.19 (0.00-0.50)	88	<.01	0.61 (0.41-0.82)	88	<.01	0.08 (0.01-0.59)	.02	57	.04
After first dose	Single study			Single study			Single study			
Targeted therapy										
After second dose	0.35 (0.19-0.52)	94	<.01	No controls			No analysis			
After first dose	Single study			No controls			No analysis			
HSCT or cellular the	rapy within 12 months vs after 12	e mont	hs							
After second dose	0.66 (0.43-0.88)	0	.63	0.68 (0.40-0.95)	45	.16	0.96 (0.11-8.74)	.94	36	.21
After first dose	Single study			Single study			Single study			
BNT162b2 vs non-Bi	NT162b2 vaccine type*									
After second dose	0.77 (0.40-1.00)	98	<.01	0.81 (0.56-1.00)	87	<.01	1.08 (0.20-5.92)	.90	64	.04
After first dose	0.51 (0.13-0.90)	86	<.01	0.64 (0.36-0.91)	73	.01	0.59 (0.22-1.60)	.19	7	.36

^{*}Non-BNT162b2 includes mRNA-1273, ChAdOx1, and Ad26 vaccines.

seropositivity rates of 28% compared with a rate of 62% when patients were not receiving active therapy with an OR of 0.21 (95% Cl, 0.06-0.67; P = .02). Lower seropositivity rates (19% vs 61%) were reported with vaccination during or within 12 months of CD-20 antibody therapy compared with vaccination 12 or more months after completion of therapy. Use of targeted therapy was associated with a pooled seropositivity rate of 35% after 2 doses of COVID-19 vaccine. Seropositivity rates did not differ by timing of vaccination in relation to HCT (66% vs 68%) or vaccine type (BNT162b2 vs others, including mRNA-1273, ChAdOx1, Ad26; [51% vs 64%] [77% vs 81%]) after 1 or 2 doses. Table 4 summarizes the seropositivity rates by each subgroup analyzed.

Sensitivity analysis

Excluding single-arm studies that were assessed as poor quality or as having a high risk of bias did not alter the pooled seropositivity rates, but in comparator studies, OR was 0.17 (95% Cl, 0.04-0.75; P = .03) after 1 dose instead of 0.10 (supplemental Table 13).

Discussion

In this systematic review and meta-analysis of more than 7000 patients with hematologic malignancies, rate of seropositivity was 62% to 66% after 2 doses and 37% to 51% after 1 dose of COVID-19 vaccine. Compared with age-matched and non-agematched healthy controls (primarily health care workers), odds of achieving seropositivity were significantly lower by 96% after 2 doses of COVID-19 vaccine and 90% after 1 dose. Statistical heterogeneity was substantial at more than 70%, and studies were also clinically heterogenous because of the variety of underlying hematologic malignancies, lack of standardized laboratory platforms by which to measure immune response to vaccination, and variable follow-up periods. Reassuringly, reported rates of at least 1 AE after vaccination were lower at 36% compared with rates of AEs (51% to 88%) reported in clinical trials of the general population. 10,13

Among different hematologic malignancies, pooled seropositivity rates after 2 doses were highest at close to 90% in patients with acute leukemia, MDS, or MPNs and the lowest at 51% in patients with CLL. Patients with CLL respond poorly to vaccination, especially to new or novel antigens compared with recall antigens from previous infections or vaccination.⁶⁷ Poor response is compounded by use of B-cell-depleting and targeted therapies such as CD-20 monoclonal antibodies (mAb) and BTKi's.67 Immune response to other vaccines, including the seasonal influenza vaccine, is negatively impacted by these therapies, and the poor response persists for at least 6 to 12 months after cessation of therapy.⁶⁸

A similar negative impact of treatment on COVID-19 vaccine responses was noted in subgroup analysis of published studies. Rates of seropositivity were lower in the setting of active treatment (28%), and the lowest response rates were reported in the setting of current or CD-20 mAb therapy within 12 months (19%), targeted therapies (35%), and after therapy with CAR T cells (31%). Seropositivity rates were 2 to 3 times higher at 62% when patients were vaccinated when they were not receiving active therapy and 61% when vaccinated at 12 or more months after completion of mAb CD-20 therapy.

Unvaccinated patients with CLL have a high burden of morbidity from COVID-19 infection with close to 90% of patients requiring hospital admission, 35% requiring admission to intensive care units, and a mortality rate of 33%.4 Yet patients with CLL have poor humoral responses to vaccination: 18% after 1 dose and 51% after 2 doses. In patients with CLL, immune suppression from underlying disease and ongoing treatments such as anti-CD20 mAb's and BTKi's continue to pose a risk for SARS-CoV-2 infection and concurrently limit protective responses from vaccination. Although vaccination remains highly recommended, new strategies are required to

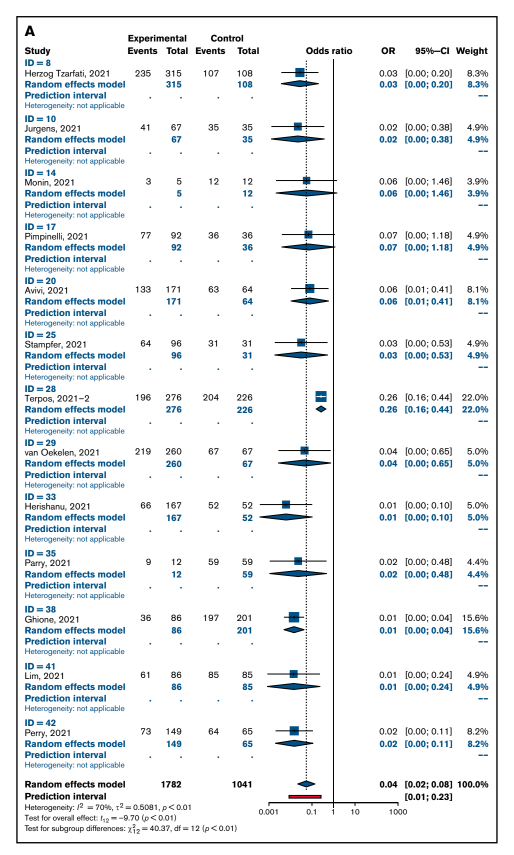


Figure 2. OR for achieving seropositivity in patients with hematologic malignancies vs healthy control group after 2 doses (A) and 1 dose of COVID-19 vaccine (B).

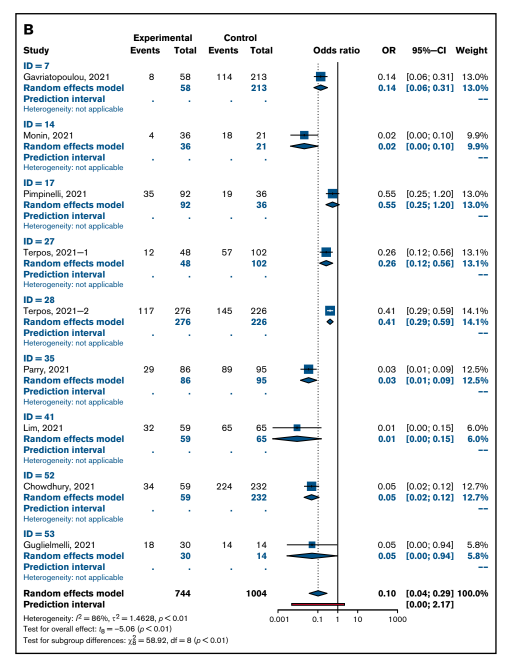


Figure 2. (continued)

further improve immune responses in this high-risk patient population. In other groups of immune compromised patients, the use of an additional dose of COVID-19 vaccine improved serologic response rates by 37%, and the use of heterologous vaccination schedules (mixing vaccine types) appears promising. 69-71 International health groups now recommend a third dose of COVID-19 vaccine for immune compromised patients, including those with hematologic malignancies. 72,73 Heterologous prime boost or use of high-dose vaccine formulations have been used in randomized trials of seasonal influenza vaccination in hematology and in patients who have received an HCT with mixed success. 17,74 In addition, other preventative approaches such as the use of anti-SARS-CoV-2 mAb

therapy as pre-exposure (NCT04625725) and post-exposure prophylaxis require further evaluation in vulnerable patient groups who respond poorly to vaccination.⁷⁵

Serologic responses are classically used as surrogate end points for clinical efficacy in clinical trials of vaccination in patients with hematologic malignancies. 15-17 Serologic thresholds for protection (seroprotection) after vaccination have been established for infections such as seasonal influenza.17 In the majority of studies included in this review, however, the outcome of interest was seropositivity as defined by antibody levels above the detection threshold. This is not equivalent to seroprotection because thresholds

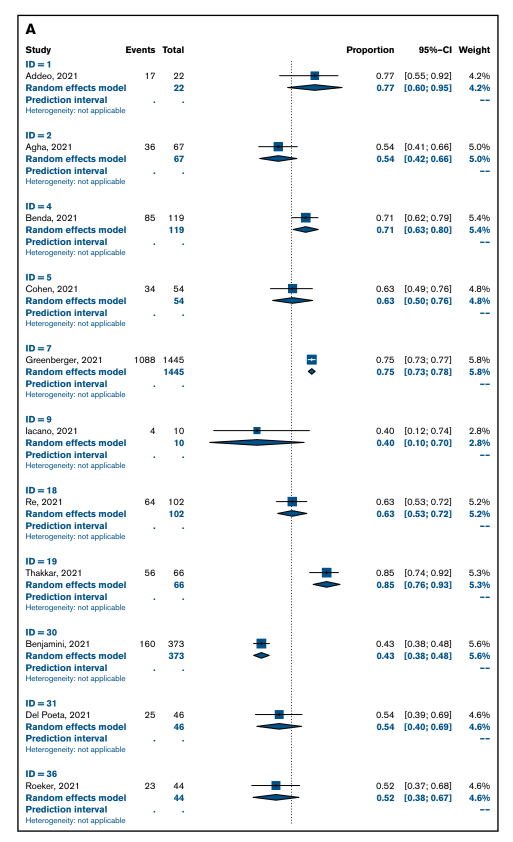


Figure 3. Pooled rates of seropositivity in single-arm studies involving patients with hematologic malignancies after 2 doses (A) and 1 dose of COVID-19 vaccine (B).

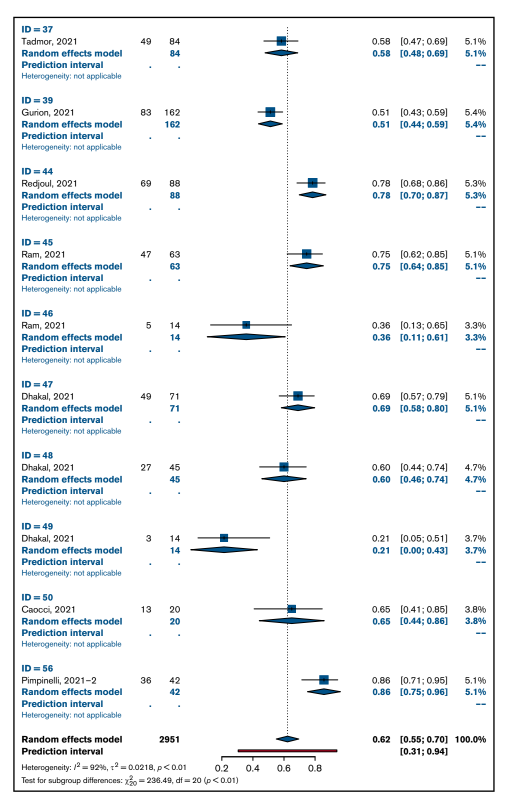


Figure 3. (continued)

have not been standardized nor have they been established across the variety of commercial and research platforms used by these studies. Some authors such as Malard et al, 33 Stampfer et al, 43 and Redjoul et al⁵⁸ have attempted to define serologic thresholds that correlate with nAb and clinical protection. Unsurprisingly, a lower proportion of patients (by 20% to 35% compared with pooled

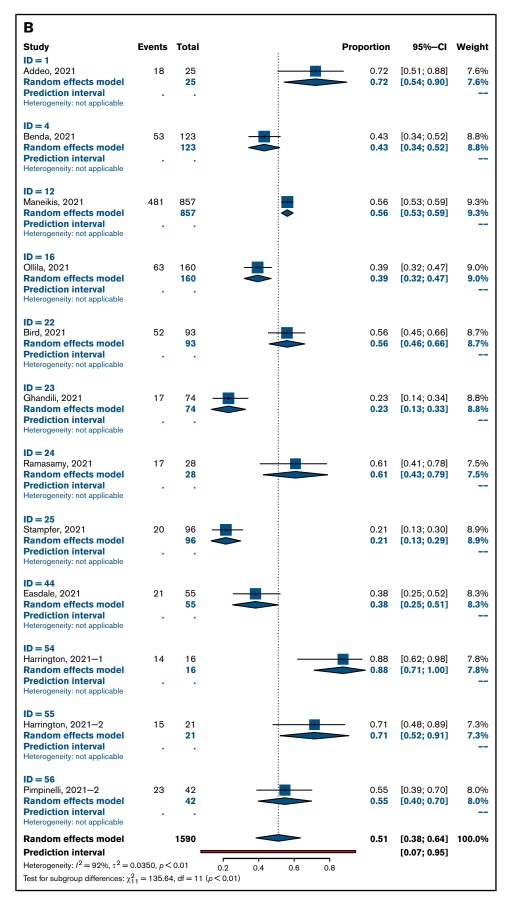


Figure 3. (continued)

rates) achieved these higher antibody thresholds. 33,43,58 For serologic response measurement to guide clinical management of COVID-19 vaccination, further work is required to achieve harmonization across testing platforms and to derive and validate thresholds that correlate with clinical protection.

Measurement of serologic responses offers only a glimpse of the potential breadth of immune responses to COVID-19 vaccination. Both nAb and cellular responses to vaccination play complementary and vital roles in protection against COVID-19 and remain under-reported. 76-78 In 16% of studies, at least 18% of hematology patients achieved a positive nAb response. Positive cellular response rates were at least 15% higher than nAb responses after 1 or 2 doses of vaccine. Although serologic responses have been used as surrogate end points in vaccination studies of immune compromised patients, further large studies are required to identify new immune markers for vaccine response and to determine the efficacy of vaccination.

This review has several limitations. In particular, the findings are of moderate quality because of significant clinical and statistical heterogeneity of included studies and the proportion of studies of poorer quality. In line with other established studies of vaccination in hematology patients, only immune response data were analyzed because clinical efficacy data were limited.

In conclusion, this systematic review and meta-analysis has comprehensively summarized the latest data on response to COVID-19 vaccination in patients with hematologic malignancies. Overall, seropositive rates were reasonable at 66% after 2 doses of vaccine. Higher-risk patient groups were identified, namely patients with CLL and patients receiving active therapy,

including targeted and CD-20 mAb therapies. New approaches to treating high-risk patients who are poor responders to vaccination are urgently required.

Acknowledgments

B.W.T. is supported by the Australian Government Medical Research Future Fund Investigator Fellowship. M.A.S is supported by a National Health and Medical Research Council Investigator Fellowship.

Authorship

Contribution: B.W.T coordinated the study, performed abstract and full-text screening and data extraction, and wrote the manuscript; J.S.K.T. performed abstract and full-text screening and data extraction; J.C. developed the study protocol and performed data extraction; S.L. developed the search strategy; T.S. conducted the statistical analysis; Z.C.F.N. and M.A.S. assessed the risk of bias in the studies; and all authors reviewed and revised the manuscript.

Conflict-of-interest disclosure: M.A.S. has received research funding and honoraria from Pfizer and Merck Sharp & Dohme and research funding from Gilead. B.W.T has received research funding and honoraria from CSL-Behring, Merck Sharp & Dohme, and Sanofi. The remaining authors declare no competing financial interests.

ORCID profiles: J.S.K.T., 0000-0001-7825-691X; J.C., 0000-0002-4302-6599; B.W.T., 0000-0003-0213-5470.

Correspondence: Benjamin W. Teh, Department of Infectious Diseases, Peter MacCallum Cancer Centre, 305 Grattan Street, Melbourne, Vic 3000, Australia; e-mail: ben.teh@petermac.org.

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