The persistence of anti-Spike antibodies following two SARS-CoV-2 vaccine doses in patients on immunosuppressive therapy compared to healthy controls—a prospective cohort study

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Abstract
Background: The durability of vaccine-induced humoral immunity against SARS-CoV-2 in patients with immune-mediated inflammatory diseases (IMIDs) on immunosuppressive therapy is not known. The aim of this study was to compare the persistence of anti-Spike antibodies following two-dose SARS-CoV-2 vaccination between IMID patients and healthy controls and to identify factors associated with antibody decline.

Methods: IMID patients on immunosuppressive medication enrolled in the prospective observational Nor-væC study were included. Participants received two-dose SARS-CoV-2 vaccination. Serum collected at two time points following vaccination (first assessment within 6–48 days, second within 49–123 days) were analyzed for antibodies binding the receptor-binding domain (RBD) of the SARS-CoV-2 Spike protein. Multivariable regression models estimated percent reduction in anti-RBD over 30 days and factors associated with reduction.

Results: A total of 1108 patients (403 rheumatoid arthritis, 195 psoriatic arthritis, 195 spondyloarthritis, 124 ulcerative colitis, 191 Crohn’s disease) and 134 controls provided blood samples within the defined intervals (median 19 days [IQR 15–24] and 97 days [87–105] after second vaccine dose). Antibody levels were lower in patients compared...
Key messages

1. Anti-Spike antibody decline 4 months post-vaccination was significantly larger in patients (−83%) compared to controls (−66%).
2. Four months post-vaccination, antibody levels decreased to a low level (<200 BAU/ml) in 41% of IMID patients and 5% of controls.
3. Among therapies, the use of tumor necrosis factor inhibitors in mono- or combination therapy was associated with the most pronounced decline in anti-Spike antibodies.

Background

We rely on efficient vaccines for long-lasting protection against COVID-19 disease in order to counter the SARS-CoV-2 pandemic. SARS-CoV-2 vaccines have a proven efficacy in the general population, and antibody levels correlate to the degree of clinical protection against COVID-19 disease [1–3]. However, recent data suggest that patients with immune-mediated inflammatory disease (IMID) treated with certain immunosuppressants have an impaired serological response following vaccination, with lower anti-Spike antibody levels than the general population [4–15]. The impairment of the immune response varies between the different immunosuppressive therapies, where some seemingly do not affect the humoral response, while other drugs, such as rituximab and abatacept, have been found to profoundly reduce vaccine response [12, 15, 17–26].

IMIDs encompass a number of prevalent chronic diseases, including inflammatory joint diseases such as rheumatoid arthritis (RA), psoriatic arthritis (PsA), spondylarthritis (SpA), and inflammatory bowel disease (IBD); ulcerative colitis (UC); and Crohn’s disease (CD). Immunosuppressive medication, including tumor necrosis factor inhibitors (TNFi), non-TNFi biologic agents, metabolite inhibitors, and targeted small molecule drugs, are pivotal in the treatment of IMIDs [18–21]. Use of immunosuppressive therapies combined with a dysregulated immune system and an increased frequency of several co-morbidities increases the susceptibility to serious infections and vulnerability to adverse outcomes of infectious diseases in these patients [22–24]. There are, however, disparities between therapies regarding the degree of impact on the immune responses and thus the susceptibility to serious infections [12, 15, 17, 25, 26].

Evidence of a waning immune response over time is emerging in the healthy population, both in terms of antibody levels and protection against symptomatic COVID-19 disease [27–35]. Given the risk of severe COVID-19 disease faced by the IMID population [36–38], a possible weakened long-term immunological protection elicited by SARS-CoV-2 vaccines has been raised as a concern for this patient population. A recent study on patients with rheumatic diseases has reported a three-fold decrease in antibodies 6 months after two SARS-CoV mRNA vaccines [39], but the impact of various medications is unknown. Further, it is not known whether there is a difference in the perseverance of vaccine-induced immunity against SARS-CoV-2 between IMID patients and the general population. Data on the long-term effectiveness of the SARS-CoV-2 vaccines in immunosuppressed patients with IMIDs are needed to assess the protection against severe COVID-19 disease over time in this large at-risk population and to make decisions on the appropriate timing of booster doses.

The main aim of this study was to evaluate the persistence of the serologic response to two-dose SARS-CoV-2 vaccination in IMID patients and healthy controls within 4 months post-vaccination. Secondary aims were to compare the rate of decay of anti-Spike antibodies in patients and controls and to identify predictors of antibody
Methods

Study design and participants

The present study uses data from the ongoing Nor-vaC study (Norwegian study of vaccine response to COVID-19 vaccines in patients using immunosuppressive medication within rheumatology and gastroenterology). Nor-vaC is a prospective, observational study which included patients diagnosed with RA, PsA, SpA, UC, or CD who intended to receive SARS-CoV-2 vaccination. The study is conducted at two Norwegian hospitals; the Division of Rheumatology and Research at Diakonhjemmet Hospital (DH) and the Department of Gastroenterology at Akershus University Hospital (AHUS), both with large specialist clinics. Adult patients (aged ≥18 years) were recruited into Nor-vaC prior to the initiation of the Norwegian national vaccination program in February 2021. Eligible patients treated with immunosuppressive drugs were identified from hospital records at DH and AHUS and invited to participate in the study. Eligibility criteria are described in the Additional file 1: Section 1. Health care workers from DH and AHUS constituted the healthy control group. In the present study, we included patients and healthy controls who provided blood samples at both first (6–48 days) and second assessment (49–123 days) after the second vaccine dose, for details of inclusion, see Additional file 1: Fig. S1. Immunosuppressive medications were arranged into the following categories: Metabolite inhibitors (methotrexate, sulfasalazine, leflunomide, azathioprine and mercaptopurine), interleukin inhibitors (tocilizumab, ixekizumab, ustekinumab, secukinumab, risankizumab), vedolizumab (IBD patients only), abatacept (RA patients only), janus kinase inhibitors (JAKi), tumor necrosis factor inhibitors (TNFi) monotherapy, TNFi combination therapy (combined with a metabolite inhibitor or vedolizumab), and rituximab (RA patients only).

The study was approved by an independent ethics committee (Regional Committees for Medical and Health Research Ethics South East, reference numbers 235424, 135924) and institutional review boards. All patients and healthy controls signed informed consent. The study is registered at clinicaltrials.gov (NCT04798625).

Study procedures

According to the national vaccination program, as instructed by the Norwegian Institute of Public Health (NIPH) and administered by the public health system, all participants were vaccinated with two doses of SARS-CoV-2 vaccine, with the exception of those with prior COVID-19 disease who received one vaccine dose only. Initially, there were three SARS-CoV-2 vaccines available: BNT162b2, mRNA-1273, and ChAdOx1, until ChAdOx1 was withdrawn from the Norwegian vaccination program in March 2021 due to reports of serious side effects [40]. The interval between two doses of the mRNA vaccines was 3–6 weeks. All participants who had received one dose of ChAdOx1 vaccine were given one of the mRNA vaccines as the second dose after an interval of 9–12 weeks.

Data collection

Data was collected using electronic questionnaires handled by Services for Sensitive Data (TSD) at DH and Viedoc version 4 at AHUS. For patients; demographic data, medication use, disease activity, and information regarding previous COVID-19 were collected before vaccination. Disease activity was assessed by the Harvey-Bradshaw index and the Mayo Score in CD and UC patients, respectively and by patient-reported global assessment of disease activity for patients with RA, PsA and SpA. Age, gender, vaccine type, and COVID-19-related information were collected for healthy controls. Participants were asked to self-report COVID-19 disease at follow-up questionnaires. COVID-19 disease was defined by positive PCR and/or rapid antigen test. In addition, participants were linked by a unique personal identification number to the Norwegian Immunisation Registry (SYSVAK) and Norwegian Surveillance System for Communicable Diseases (MSIS) providing information on the date and type of vaccination received and the date of COVID-19 disease when applicable [41, 42].

Serological analyses

Participants were requested to donate serum samples 2–4 weeks and 3 months after the second vaccine dose. The Department of Immunology at OUH performed the antibody assessments. An in-house bead-based method was used to measure antibodies to the receptor-binding domain (RBD) at the full-length SARS-CoV-2 Spike protein and binding of the Spike-protein to the ACE-2 receptor. This method was validated against a micro-neutralization assay [43]. Results were given in Binding Antibody Units (BAU) per ml.

Antibody levels were categorized according to the official specifications at Oslo University Hospital as follows: less than 5 BAU/ml was defined as negative, 5–19 BAU/ml very weak positive, 20–199 BAU/ml weak positive, 200–1999 BAU/ml positive, 2000–8999 BAU/ml strong positive and 9000 BAU/ml or more very strong positive. 200 BAU/ml was defined as the threshold value for a positive result as 200 BAU/ml was determined to be the lower threshold for detection of neutralizing antibodies by the usage of a micro-neutralization assay [43].
Outcomes
The main outcome of this study was the persistence of antibodies after two-dose SARS-CoV-2 vaccination in IMID patients and controls, measured as the magnitude of the immunoglobulin G antibody levels to the receptor binding domain on the SARS-CoV-2 spike protein (anti-RBD) at 4 months. Additional outcomes were the percentage reduction in antibodies from first to second assessment in patients compared to controls.

Statistical analyses
Anti-RBD levels, estimated percentage change in anti-RBD levels, and time interval between first and second assessment of antibody levels were compared between groups using Student’s t-test and Mann-Whitney U test as appropriate. Participants with increasing anti-RBD levels between first and second serologic assessment post-vaccination were excluded from further analyses (Additional file 1: Fig. S1).

Multivariable linear regression
Factors associated with the 30 days estimated percent reduction in anti-RBD levels were identified by entering the following variables in three separate multivariable regression models including all participants; patients vs. controls, types of vaccination, and type of immunosuppressive medication. Diagnoses were entered into a fourth model that only included patients with a type of immunosuppressive medication. Diagnoses were entered into a fourth model that only included patients with a type of immunosuppressive medication entered as a possible confounder. Groups of immunosuppressive medications with less than 30 patients were excluded from the regression analyses. All models were corrected for age, gender, anti-RBD at first assessment, and time between blood samples. For details, see Additional file 1: Fig. S2 and Table S1.

The relationship between body mass index (BMI) and measures of disease activity and percent change in anti-RBD was explored in disease-specific models. To explore the impact of age on antibody decline, an age-stratified multivariable regression was performed.

Results
Population characteristics
Between 6th of April 2021, and 9th of November 2021, a total of 1108 patients (median age 54 years [IQR 43–64]; 617 women [56%]) and 134 controls (median age 46 years [IQR 35–56]; 111 women [83%]) provided two blood samples within the defined intervals for serologic assessment (median 19 days [IQR 15–24] and 97 days [87–105] after second vaccine dose), and were included in the present study (Additional file 1: Fig. S3). Characteristics of the participants are presented in Table 1.

Persistence of serological response
Anti-RBD levels were significantly lower in patients as compared to controls at first (median anti-RBD BAU/ml 2806 [IQR 1018–6068] vs. 6187 [IQR 4105–7496], p <0.001), and second assessment post-vaccination (median anti-RBD BAU/ml 608 [IQR 58–1053] vs. 1520 [979–3766], p <0.001) (Table 2). Changes in anti-RBD levels between the first and second assessment by medication groups and by diagnoses are presented in Fig. 1 and in Additional file 1: Fig. S4, respectively.

The median reduction of anti-RBD was significantly higher in patients (− 83% [IQR − 94 to − 66]) as compared to controls (− 66% [IQR − 79 to − 49]), p<0.001 (Table 2, Fig. 2a). As presented in Fig. 2b, the median change was greatest in the TNFi mono (− 86% [IQR − 96 to − 75], p<0.001) and TNFi combination (− 87% [IQR − 96 to − 74], p <0.01) groups compared to controls. The percent reduction in anti-RBD was significantly larger for all diagnoses, compared to controls, Additional file 1: Table S2.

At second assessment, 449 (41%) patients vs. 6 (5%) controls had low anti-RBD levels (<200 BAU/ml); p<0.0001, whereas only 148 (13.5%) patients vs. 58 (43%) controls had anti-RBD levels > 2000 BAU/ml, p<0.001 (Table 2). The distribution of anti-RBD levels by medication groups at first and second assessment are shown in Fig. 3a, b and in Additional file 1: Table S3. In patients treated with TNFi in mono- or combination therapy, 192 (41%) and 147 (56%) had anti-RBD < 200 BAU/ml at the second assessment, respectively (Fig. 3a, b, Additional file 1: Table S3). Only 46 (10%) and 19 (7%) of patients using TNFi monotherapy or TNFi combination therapy, respectively, had persisting anti-RBD levels > 2000 BAU/ml at the second assessment. The medication groups with the highest proportion of patients with anti-RBD>2000 BAU/ml at the second assessment were JAKi (9 patients [41%]) and Vedolizumab (14 patients [44%]); however, the number of patients in each of these medication groups was small (n=22 and n=32) (Fig. 3b). Additional file 1: Fig. S4 shows anti-RBD levels across diagnoses at both assessments. Of all diagnoses, the highest percentage of patients with anti-RBD <200 BAU/ml at second assessment, was found in CD (92 patients [48%]).

Factors influencing percent reduction in anti-RBD levels
In multivariable regression models (Table 3) a significantly greater estimated 30 days percent reduction in anti-RBD levels was found in patients (β = 6.4 [95% CI 8.4 to − 4.3]) compared to controls (p<0.001). The difference remained significant after adjusting for vaccine type. Age-stratified (< 50 and ≥ 50 years of age) multivariable regression showed similar results with a significantly greater estimated 30 days percent reduction in
### Table 1 Characteristics of the participants

|                      | Controls (n=134) | Patients (n=1108) | Rheumatoid arthritis (n=403) | Psoriatic arthritis (n=195) | Spondyloarthritis (n=195) | Ulcerative colitis (n=124) | Crohn's disease (n=191) |
|----------------------|-----------------|-------------------|-----------------------------|-----------------------------|---------------------------|---------------------------|-------------------------|
| **Demographics**     |                 |                   |                             |                             |                           |                           |                         |
| Age (years), median (IQR) | 45.6 (34.9–56.1) | 54.3 (43.2–64.1)  | 61.2 (51.3–69.1)            | 57.9 (48.3–64.7)            | 50.5 (42.0–59.2)           | 45.7 (33.4–54.6)          | 43.0 (29.6–53.9)         |
| Female, n (%)        | 111 (83)        | 617 (56)          | 308 (76)                    | 104 (53)                    | 75 (38)                   | 53 (43)                   | 77 (40)                 |
| Male, n (%)          | 23 (17)         | 491 (44)          | 95 (24)                     | 91 (47)                     | 120 (62)                  | 71 (57)                   | 114 (60)                |
| **Patient reported disease activity, median (IQR)** |                 |                   |                             |                             |                           |                           |                         |
|                      |                 |                   |                             |                             |                           |                           |                         |
| **Vaccines, n(%)**   |                 |                   |                             |                             |                           |                           |                         |
| BNT162b2 & mRNA-1273 |                 |                   |                             |                             |                           |                           |                         |
| BNT162b2 & mRNA-1273 | 11 (8)          | 780 (70)          | 282 (70)                    | 135 (69)                    | 125 (64)                  | 94 (76)                   | 144 (75)                |
| mRNA-1273 & mRNA-1273| 43 (32)         | 285 (26)          | 108 (26.8)                  | 52 (27)                     | 60 (31)                   | 25 (20)                   | 40 (21)                 |
| Mixed*               | 80 (60)         | 37 (3)            | 12 (3)                      | 7 (3.5)                     | 10 (5)                    | 3 (2.4)                   | 5 (3)                   |
| COVID-19 infection and one vaccine dose |                 |                   |                             |                             |                           |                           |                         |
|                      |                 |                   |                             |                             |                           |                           |                         |
| **Medication, n(%) of the total number of patients** |                 |                   |                             |                             |                           |                           |                         |
| Rituximab            |                 | 31 (3)            | 31 (8)                      |                             |                           |                           |                         |
| Tumor necrosis factor inhibitors |                 |                   |                             |                             |                           |                           |                         |
| - Monotherapy        | 464 (42)        | 51 (12.5)         | 60 (31)                     | 163 (83.5)                  | 67 (54)                   | 123 (64)                  |                         |
| - Combination        | 261 (23.5)      | 106 (26)          | 64 (32)                     | 25 (13)                     | 24 (19)                   | 43 (23)                   |                         |
| Interleukin inhibitors |                 |                   |                             |                             |                           |                           |                         |
| - Tocilizumab        | 19 (2)          | 17 (4)            | 2 (1)                       |                             |                           |                           |                         |
| - Other interleukin inhibitors |                 |                   |                             |                             |                           |                           |                         |
| Janus kinase inhibitors | 22 (2)        | 13 (3)            | 2 (1)                       |                             |                           |                           |                         |
| Abatacept            | 6 (0.5)         | 6 (1.5)           |                             |                             |                           |                           |                         |
| Vedolizumab          | 32 (3)          | 22 (18)           |                             |                             |                           |                           |                         |
| Metabolite inhibitors |                 |                   |                             |                             |                           |                           |                         |
| - Methotrexate mono-therapy | 218 (19.5) | 161 (40)          | 53 (27)                     | 32 (27)                     |                           |                           |                         |
| - Other metabolite inhibitors |                 |                   |                             |                             |                           |                           |                         |
| Prednisolone mono-therapy | 13 (1.5)     | 7 (2)             | 2 (3)                       | 1 (0.5)                     |                           |                           |                         |

* Patients using prednisolone in doses <10mg/day in combination with other medication groups are included in all groups

* Patient reported disease activity at baseline, indicated on a visual analog scale 0–100

*Missing information in 23 patients

* ChAdOx1 + BNT162b2/mRNA-1273 or BNT162b2 + mRNA-1273

* In combination with metabolite inhibitors or vedolizumab

* Ustekinumab, risankizumab, secukinumab, iflexikumab

* Azathioprine, mercaptopurin, sulfasalazine, leflunomide in monotherapy or in combination with each other or in combination with methotrexate
Patients had lower anti-RBD levels at the first assessment, with a more pronounced decline than RA patients (β varied significantly across diagnoses. Patients with UC compared to controls. The reduction in anti-RBD levels 1273 as reference, mRNA-vaccine type using BNT162b2 ×<0.05. When comparing 2.9 [95% CI −0.2], −5.7 to p<0.05, respectively). Age and gender were not associated with the percentage change in anti-RBD levels.

In the disease-specific models (Additional file 1: Tables S4 and S5) markers of disease activity and BMI were not significantly associated with estimated 30 day reduction in anti-RBD levels.

**Discussion**

In this large observational study examining the persistence of anti-Spike antibodies after two-dose SARS-CoV-2 vaccination in IMID patients on immuno-suppressive therapies, we demonstrated that antibody levels declined considerably in both IMID patients and controls within 4 months after the second vaccine dose. Patients had lower anti-RBD levels at the first assessment, and a more rapid reduction in antibody levels, resulting in a higher proportion of patients with low antibody levels after 4 months compared to controls. The overall reduction of anti-RBD levels within 4 months after the second vaccine dose was significantly higher in patients (−83%) compared to controls (−66%). A considerably larger proportion of IMID patients (41%) compared to controls (5%) declined to low (<200 BAU/ml) antibody levels 4 months post vaccination. By medication groups, those treated with rituximab or TNFi mono- or combination therapy were more likely to have low antibody levels at 4 months after the second vaccine dose. Furthermore, a diagnosis of UC was associated with the highest antibody decline over time.

We have previously reported that patients with IMIDs have an attenuated SARS-CoV-2 vaccine response compared to healthy controls at the first assessment after a standard two-dose regimen [12, 15]. We now report continued low antibody levels in this 4-month follow-up study. A threshold antibody level giving actual, clinical protection against breakthrough COVID-19 infection has not yet been established. However, evidence for a correlation between antibody levels and the protective immunity against COVID-19 infection is emerging [3, 44–48]. Gilbert et al. demonstrated a decreasing risk of serious COVID-19 disease with increasing antibody level and that the protection against COVID-19 induced by the mRNA-1273 vaccine was improved with increasing levels of antibodies [3]. Given the considerable decrease in antibody levels demonstrated in this present study and recently reported by others [31, 39], the duration of

| Table 2 Serological response at first and second assessment following two-dose vaccination in patients and controls |
|---------------------------------------------------------------|
| **Controls** | **Patients** |
| **First sera assessment** | **Second sera assessment** | **First sera assessment** | **Second sera assessment** |
| Median anti-RBD level (IQR) | 6187 (4105–7496) | 2806 (1018–6068) | 608 (58–1053) |
| Median change in anti-RBD level (IQR) | −3332 (−5096 to −2206) | −2039 (−4304 to −806) | 1273 (−8.4 to −5.9) |
| Median percent change in anti-RBD level (IQR) | −66 (−79 to −49) | −83 (−94 to −66) | 10.7 | 8.6 [95% CI 5.9, −10.4 to 8.6] |
| Mean (SD) number of days between first and second assessment | 75 (16) | 75 (17) | 75 (16) | 75 (17) |
| Anti-RBD <5, n (%) | 0 | 0 | 17 (1.5) | 56 (5) |
| Anti-RBD 5–19, n (%) | 0 | 0 | 13 (1) | 74 (6.5) |
| Anti-RBD 20–199, n (%) | 0 | 6 (5) | 62 (6) | 319 (29) |
| Anti-RBD 200–1999, n (%) | 10 (7.5) | 70 (52) | 366 (33) | 511 (46) |
| Anti-RBD 2000–8999, n (%) | 107 (80) | 58 (43) | 599 (54) | 145 (13) |
| Anti-RBD ≥ 9000, n (%) | 17 (12.5) | 0 | 51 (4.5) | 3 (0.5) |

First sera assessment 6–48 days after second vaccine dose. Second sera assessment 49–123 days after second vaccine dose. Serological response is anti-SARS-CoV-2 IgG antibodies to the receptor binding domain (RBD) measured as BAU/ml binding antibody units/ml, IQR inter quartile range

* Change in median anti-RBD level compared across groups by Mann-Whitney U test: p<0.001

<sup>a</sup> Percent change in anti-RBD level compared between groups by Mann-Whitney U test: p<0.001

<sup>**</sup> Mean number of days between first and second assessment compared between groups by Student’s t-test p=0.77

anti-RBD levels in patients compared to controls for both age categories (data not shown). Use of TNFi in mono- or combination therapy was associated with a larger reduction in anti-RBD levels (β = 8.6 [95% CI −10.7 to 6.5]), β = 8.1 [95% CI −10.4 to −5.9], p<0.001, respectively), compared to controls. The reduction in anti-RBD levels varied significantly across diagnoses. Patients with UC had a more pronounced decline than RA patients (β = 2.9 [95% CI −5.7 to −0.2], p<0.05. When comparing vaccine type using BNT162b2 × 2 as reference, mRNA-1273 × 2 or mixed vaccine type was significantly associated with a lower reduction in antibodies (β 4.4 [95% CI 3.0–5.9], p<0.001, and β 2.8 [95% CI 0.2–5.4], p<0.05, respectively).
protection against COVID-19 afforded by the vaccines remains uncertain. In this current study, we show that anti-RBD levels decrease more rapidly in IMID patients than in healthy controls. Recent studies in the general population have demonstrated waning antibody levels over time and reduced long-term protection against COVID-19 disease induced by SARS-CoV-2 vaccines [27–34]. Antibodies have also been shown to decline following SARS-CoV-2 infections after a peak between 20 and 30 days after onset of symptoms, although most individuals had high levels of IgG up to 94 days after infections [47]. Short-term serological responses have been investigated in patients with autoimmune diseases treated with immunosuppressive medications, demonstrating lower serological response rates and antibody levels after two-dose SARS-CoV-2 vaccination [5, 6, 9, 11, 17, 49]. However, there are limited data available regarding the persistence of humoral immunity in immunosuppressed patients. A recent study by Levin et al. assessing humoral response over 6 months after the second vaccine dose, reported that antibody levels were severely depleted in immunosuppressed patients compared to those without immunosuppressive therapies (only 13 patients were followed until 6 months post-vaccination). In contrast to our study, they found similar rates of antibody reduction between those with and without immunosuppression [31]. The present study demonstrates a greater reduction in antibody levels in patients (83%) than recently found by Frey et al., who reported a 64% reduction in antibodies. 

**Fig. 1** Levels of anti-RBD antibodies at the first and second assessment according to medication group. The orange bars show anti-RBD levels at the first assessment and the purple bars show anti-RBD at the second assessment, 6–48 and 49–123 days after the second vaccine dose, respectively. Bars indicate the lower and upper quartiles. Horizontal lines inside the bars indicate the median. Vertical lines through the bars show the minimum (Q1–1.5×IQR) and maximum value (Q3+1.5×IQR). Dots indicate outliers. A cut-off at 200 BAU/ml is indicated by a red line. MTX mono, methotrexate monotherapy; ILi, interleukin inhibitors including tocilizumab, ustekinumab, ixekizumab, risankizumab, secukinumab; VED, vedolizumab; JAKi, janus kinase inhibitor; TNFi mono, tumor necrosis factor inhibitor in monotherapy; TNFi comb, tumor necrosis factor inhibitor in combination with metabolite inhibitor(s) or vedolizumab; RTX, rituximab. All groups include patients using prednisolone in doses <10mg/day in combination with other medication.
6 months after two-dose SARS-CoV-2 vaccination in 326 patients with rheumatic diseases on immunosuppressive therapy [39]; however, no controls were available. A more rapid decline in antibody levels in IMID populations could be due to the continued low level of immunoglobulin production in this population, rather than an increased clearance of antibodies.

Immunity to SARS-CoV-2 also involves cellular immune responses, and evidence of a good T-cell response despite poor humoral response is emerging [12, 50–52]. The association between a persistent humoral immune response and a cellular response has not been fully elucidated. A recent study by Chen et al. followed 27 patients who had recovered from
Fig. 3  Percent distribution of anti-RBD levels at the first (a) and second (b) assessment in patients and healthy controls. 

(a) Percent distribution of anti-RBD levels at the first assessment 6–48 days after the second vaccine dose in controls and in patients according to medication groups.

(b) Percent distribution of anti-RBD levels at the second assessment 49–123 days after the second vaccine dose in controls and in patients according to medication group. MTX mono, methotrexate monotherapy; ILi, interleukin inhibitors including tocilizumab, ustekinumab, ilsezikumab, risankizumab, secukinumab; VED, vedolizumab; JAKi, janus kinase inhibitor; TNFi mono, tumor necrosis factor inhibitor in monotherapy; TNFi comb, tumor necrosis factor inhibitor in combination with metabolite inhibitor(s) or vedolizumab; RTX, rituximab. All groups include patients using prednisolone in doses <10mg/day in combination with other medication.
SARS-CoV-2 infection. Although the anti-RBD IgG level had decreased significantly by approximately 7 months post infection, the study reports that the SARS-CoV-2 specific CD4+ T-cell response persisted, with no significant change during the follow-up period [50].

There are some limitations to this present study. Patients were older than controls and the gender distribution for both cohorts was not equal. However, age was not an effect modifier of the association between patients and controls and the rate of antibody decline. The majority of patients received vaccine type BNT162b2, while healthy controls received a combination of one ChAdOx1 and one mRNA vaccine or two mRNA-1273 vaccines. There were differences in the type of immunosuppressive medication prescribed to patient groups. Only RA patients are treated with rituximab, and given the known negative effect of rituximab on the ability to mount a serologic response, the comparison of serologic response between RA and other diagnoses is difficult. We could not fully adjust for channeling bias in the regression models.

The estimation of the percent reduction in anti-RBD levels during 30 days was based on two samples per individual, consequently, the model may not fully capture the changing rate of decay at different time points. However, the time between the first and second assessments was not significantly different between patients and controls.

| Table 3 | Linear regression models of estimated 30 days percent reduction in anti-RBD level |
|---------|---------------------------------------------------------------------------------|
|         | Model 1  | Model 2  | Model 3  | Model 4  |
|         | β (95% CI) | β (95% CI) | β (95% CI) | β (95% CI) |
| Demographics |         |         |         |         |
| Age in years | 0.0 (‑0.0–0.1) | 0.1 (‑0.0–0.1) | -0.0 (‑0.1–0.0) | -0.0 (‑0.1–0.0) |
| Female gender | 1.2 (‑0.1–2.4) | 1.2 (‑0.1–2.4) | 0.7 (‑0.6–1.9) | 0.7 (‑0.6–2.1) |
| Patients vs controls | -6.4 (‑8.4–‑4.3) ** | -4.9 (‑7.4–‑2.4) * | - | - |
| Vaccine type |         |         |         |         |
| BNT162b2 x 2 | - | - | - | - |
| COVID-19 infection and vaccine | 8.3 (‑0.2–16.8) | 2.8 (0.2–5.4) * | 4.4 (3.0–5.9) ** | - |
| mRNA-1273 x 2 | 2.8 (0.2–5.4) * | - | - | - |
| Medication |         |         |         |         |
| Methotrexate monotherapy | -1.9 (‑4.3–0.5) | - | - | - |
| Interleukin inhibitors | -0.9 (‑4.5–2.7) | 1.8 (‑1.7–5.3) | - | - |
| Vedolizumab | 0.7 (‑3.3–4.7) | 5.1 (0.7–9.5) ** | - | - |
| TNF mono | -8.6 (‑10.7–‑6.5) ** | -6.2 (‑8.4–‑4.1) ** | - | - |
| TNF comb | -8.1 (‑10.4–‑5.9) ** | -6.0 (‑8.0–‑4.0) ** | - | - |
| Rituximab | -0.3 (‑4.8–4.2) | 1.0 (‑3.3–5.3) | - | - |
| Diagnosis |         |         |         |         |
| Rheumatoid arthritis | - | - | - | - |
| Psoriatic arthritis | 0.5 (‑1.4–2.5) | - | - | - |
| Spondyloarthritides | 0.8 (‑1.4–3.1) | - | - | - |
| Ulcerative colitis | 2.9 (‑5.7–‑0.2) * | -2.3 (‑4.7–‑0.1) | - | - |
| Crohn’s disease | -2.3 (‑4.7–‑0.1) | -2.3 (‑4.7–‑0.1) | - | - |
| R²adjusted | 0.21 | 0.23 | 0.26 | 0.26 |

Linear regression models. Dependent variable of each model is estimated reduction in anti-RBD level in 30 days
All models are adjusted for anti-RBD levels at first assessment and time between first and second sera assessment. Please see Table S1 for details
Variables selected by forwards stepwise selection
Model 1 includes age, sex and patients vs. controls
Model 2 includes age, sex, patients vs. controls and type of vaccine
Model 3 includes age, sex, type immunosuppressive medication vs controls
Model 4 includes age, sex, type immunosuppressive medication and diagnosis. Controls were excluded from this model due to collinearity. (Methotrexate monotherapy and rheumatoid arthritis are comparators)
*p < 0.05, **p < 0.001
a ChAdOx1 + BNT162b2/mRNA-1273 or BNT162b2 + mRNA-1273
We cannot exclude the possibility of residual confounding due to the study design. Patients who did not provide serum samples at the second assessment were excluded from our analyses. This group may include those who had a high initial anti-RBD response at the first assessment and who therefore had low motivation to provide further serum samples. Participants with increasing anti-RBD antibodies between assessments were excluded from the present study with the assumption that many of these have been exposed to the SARS-CoV-2 virus. However, there is a possibility that some of these participants with increasing antibody levels were late peakers [31]. Additionally, we do not have methods available to identify those previously infected with COVID-19, and we cannot exclude that participants who were subjected to asymptomatic COVID-19 disease before vaccination were wrongly included in the study as COVID-19-naïve. However, the prevalence in Norway was low at that point of time, and the system in place for tracing infection makes this less likely.

To our knowledge, this is the largest study to date assessing the persistence of serologic response after SARS-CoV-2 vaccination in IMID patients treated with immunosuppressive medication compared to healthy controls. The prospective study design and a study population consisting of a large patient cohort and controls are two of the strengths of this study. Sera drawn at two time-points after the second vaccine dose enabled us to study how the vaccine response change over a longer period of time. Further, an important strength is the long time period from the second vaccine dose until the second blood sampling. This distinguishes our study from the majority of previous work on IMID patients that have assessed vaccine response at one time-point closer to the date of the second vaccination. Additionally, the generalizability of our results is increased by the composition of the patient population constituted by several autoimmune diseases from both rheumatology and gastroenterology, allowing assessment of vaccine response across a range of diagnoses and immunosuppressive treatment regimens.

Conclusions
In conclusion, our results are important when planning vaccine regimens for IMID patients and when prioritizing groups for additional vaccine doses. Our work supports a program of three initial vaccine doses and that further booster doses may be of particular importance in IMID patients who may also need this earlier than the general population.

Abbreviations
AHUS: Akerhus University Hospital; BAU: Binding Antibody Unit; CD: Crohn’s disease; DHI: Diakonhjemmet Hospital; IBD: Inflammatory bowel disease; IMID: Immune-mediated inflammatory disease; IQR: Inter-quartile range; JAKi: Janus kinases inhibitors; MSIS: Norwegian Surveillance System for Communicable Diseases; NIPH: Norwegian Institute of Public Health; OUH: Oslo University Hospital; PCR: Polymerase chain reaction; PsA: Psoriatic arthritis; RA: Rheumatoid arthritis; RBD: Receptor-binding domain; SD: Standard deviation; SpA: Spondyloarthritis; SYSVAK: Norwegian Immunisation Registry; TNFi: Tumor necrosis factor inhibitors; TSD: Services for sensitive data; UC: Ulcerative colitis.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12916-022-02587-8.

Additional file 1: Section 1. Inclusion and Exclusion Criteria. Section 2. Supplementary Figures S1-S4. Fig. S1. Flow-chart of analyses population. Fig. S2. Margins plot of the estimated 30 days reduction at different intervals between first and second serum assessment. Fig. S3. Scatter plot of antibody levels and timing of blood sampling. Fig. S4. Levels of anti-RBD antibodies at first and second assessment across diagnoses. Section 3. Supplementary Tables S1-S5. Table S1. Predictors of anti-RBD level at second assessment. Table S2. Median percentage change in anti-RBD level between first and second assessment across diagnoses. Table S3. Serological response at first and second assessment by medication group. Table S4. Linear regression models of estimated percent reduction in anti-RBD level in 30 days in patients with inflammatory joint diseases. Table S5. Linear regression models of estimated percent reduction in anti-RBD level in 30 days in patients with inflammatory bowel diseases.

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Authors’ contributions
All authors critically revised the manuscript, approved the final submitted version, and took the responsibility for the completeness and accuracy of the data and analyses. All authors had full access to all the data in the study and have made the final decision to submit the manuscript for publication. IEC, UJ, SAP, GLG, SWS, ATT, FTJ, and JS have verified the underlying data. All authors read and approved the final manuscript. IEC, UJ, GLG, SWS, KKJ, FLJ, and GG conceived and designed the present study. GLG, SWS, KKJ, FLJ, ATT, JTV, SAP, IEC, and UJ oversaw the implementation of the study. GLG, SWS, SAP, KKJ, ATT, and UJ collected the data. IEC, UJ, GLG, SWS, LAM, FLJ, JS, ATT, KKJ and SAP interpreted data and drafted the manuscript. FLJ developed the assay used for serological assessment. FLJ and TTT performed the serological analysis. JS was the study statistician. ATT, DJW, TTK, EAH, SM, GG, GBK, and JJ contributed to the study conception and design.

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Availability of data and materials
Data are available upon reasonable request after we have published all data on our predefined research objectives. The data will only be made available after the submission of a project plan outlining the reason for the request and any proposed analyses and will have to be approved by the Nor‑vaC steering group. Project proposals can be submitted to the project leader (gurololvik.goll@diakonysk.no). Data sharing will have to follow appropriate regulations.

Declarations

Ethics approval and consent to participate
Ethical approval for the Nor‑vaC study was granted by the National Committee for Research Ethics (reference numbers: REK‑235424, REK‑135924). Patients and healthy controls signed informed consent prior to inclusion.

Consent for publication
Not applicable.

Competing interests
TKK reports grants from AbbVie, Amgen, BMS, MSD, Novartis, Pfizer, UCB, consulting fees from AbbVie, Amgen, Biogen, Celltrion, Eli Lilly, Genentech, Gilead, Mylan, Novartis, Pfizer, Sandzio, Sanofi, speakers bureaus Amgen, Celltrion, Eiras, Evapharma, Evopharma, Hilma, Okta, Sandzio, Sanofi. JH reports grants from AbbVie, Pharmacomos, Ferring, consulting fees from AbbVie, Boehringer Ingelheim, BMS, Celltrion, Ferring, Gilhead, Janssen Cilag, MSD, Napp Pharma, Novartis, Orion Pharma, Pfizer, Pharmacomos, Takeda, Sandzio, Unimedica Pharma, speakers bureaus Abbvie, Astro Pharma, Boehringer Ingelheim, BMS, Celltrion, Ferring, Gilhead, Hilma, Janssen Cilag, Meda, MSD, Napp Pharma, Novartis, Orion Pharma, Pfizer, Pharmacomos, Roche, Takeda, Sandzio. LAM reports funding from KG Jøbsen foundation, support for infrastructure and biobanking from the University of Oslo and Oslo University Hospital, grants from the Coalition of Epidemic Preparedness Innovations (CEPI), speakers bureaus Novartis, Cellgene. JTV reports grant from the Coalition of Epidemic Preparedness Innovations (CEPI). GG reports consulting fees from the Norwegian System of Compensation to Patients, AstraZeneca, speakers bureaus Bayer, Sanofi Pasteur, and Theramo Fisher. FLI reports grant from the Coalition of Epidemic Preparedness Innovations (CEPI), grant from South‑East region Health authority: KKJ reports speakers bureaus Roche and BMS, advisory board Celtrion and Norgine. GLG reports funding from The Karin Fossun foundation, Diakonhjemmet Hospital, Oslo University Hospital, Akershus University Hospital, Trygve Glydtedt og frues Foundation, South‑East region Health authority, consulting fees AbbVie and Pfizer, speaker’s fees AbbVie, Pfizer, Sandzio, Orion Pharma, Novartis and UCB, advisory board Pfizer, AbbVie. IEC, I, SWS, ATT, TTT, JS, SAP, SM, DJW, GBK, and EAH report nothing to disclose.

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