

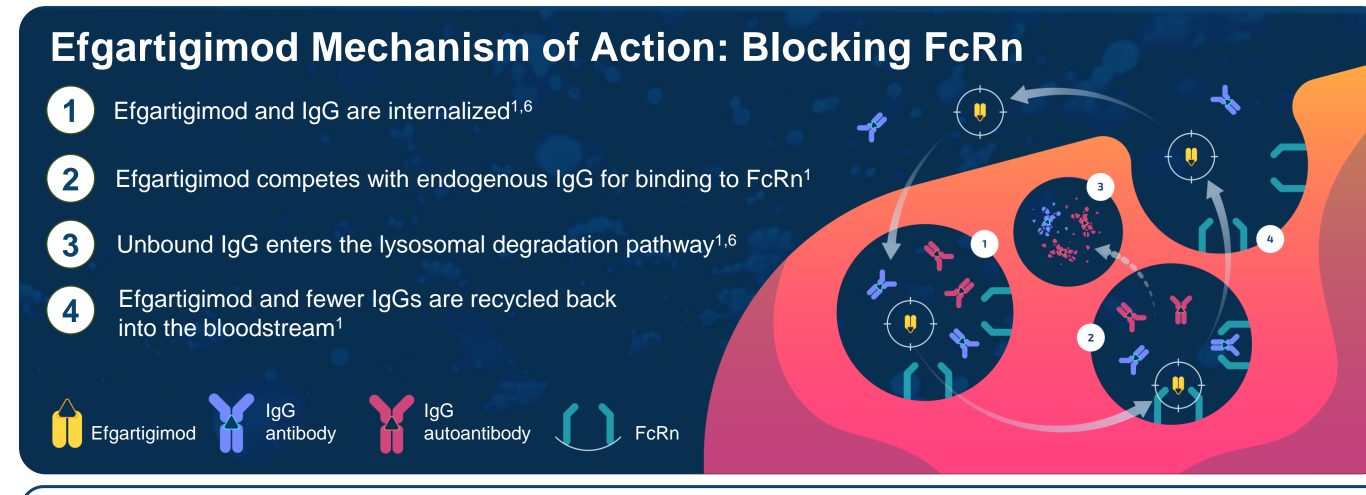
# COVID-19 Vaccination Response in Participants Across Clinical Trials Investigating Efgartigimod in Autoimmune Diseases

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#### INTRODUCTION

- Efgartigimod is an IgG1 antibody Fc fragment that has been engineered for increased affinity to FcRn compared to endogenous IgG, and is uniquely composed of the only part of the IgG antibody that normally binds FcRn¹
- Efgartigimod selectively reduces IgG by blocking FcRn-mediated IgG recycling without impacting antibody production or other parts of the immune system, and does not decrease albumin<sup>1-3</sup>
- Efgartigimod PH20 SC is a coformulation of efgartigimod and recombinant human hyaluronidase PH20 (rHuPH20), which allows for rapid SC administration of larger volumes<sup>4,5</sup>



Some immunosuppressive therapies used in the treatment of autoimmune diseases impair immune response to vaccines<sup>7</sup>

- Some immunosuppressive and B-cell-depleting therapies can reduce immunogenicity of vaccines, including vaccines to SARS-CoV-28,9
- In previous studies, efgartigimod did not impair generation of IgG responses to antigenic challenges, and levels of both naturally and vaccine-induced protective antibody titers closely followed total IgG reduction kinetics<sup>3,10,11</sup>

#### **METHODS**

#### Assess the effects of efgartigimod IV and efgartigimod PH20 SC treatment on humoral immune responses to **OBJECTIVE** COVID-19 vaccination in participants with gMG, CIDP, ITP, and PV across multiple clinical trials<sup>a</sup>

#### SARS-CoV-2-IgG-RBD, -S, -N, and -NEUT Titers Measured during gMG, CIDP, ITP, and PV studies atb,c: Max EFG PDd: PreVacc: ≥4 wk post Vacc: Cyclic dosing: Prevaccination ≥4 wk after vaccination Once-weekly/Q2W dosing: ~1 wk after 4th efgartigimod ≥4 wk after 1st efgartigimod (last available value (at maximum efgartigimod PD effect) (at maximum vaccine effect)

gwi G			CIDP		F\		
<u>ADAPT</u>	ADAPT+	ADAPT-SC	ADAPT-SC+	<u>ADHERE</u>	<u>ADVANCE</u>	<u>ADDRESS</u>	ADDRESS+
3	3, OLE	3	3, OLE	2	3	3	3, OLE
EFG IV	10 mg/kg	EFG IV 10 mg/kg or EFG PH20 SC	EFG PH20 SC 1000 mg	EFG PH20 SC 1000 mg	EFG IV 10 mg/kg		<b>SC 1000 mg</b> days 1 and 8)
≤3 cycles of 4 once-weekly infusions	≤19 cycles of 4 once-weekly infusions	1 cycle of 4 once-weekly administrations	≤11 cycles of 4 once-weekly injections	Once-weekly injections	4 once-weekly injections, then once-weekly or Q2W infusions	Once-week	ly injections
EFG IV: n=84 PBO: n=83	EFG IV: n=145	EFG IV: n=55 EFG PH20 SC: n=55 Rollover	EFG PH20 SC: n=179	Stage A, EFG PH20 SC: n=322 Stage B, EFG PH20 SC: n=111 PBO: 110	EFG IV: n=86 PBO: n=45	EFG PH20 SC: n=147 PBO: n=75	EFG PH20 SC: n=123 PBO: n=60
26 weeks	≤3 years	10 weeks	≤3 years	≤12 weeks (Stage A) ≤48 weeks (Stage B)	24 weeks	30 weeks	≤60 weeks
	EFG IV  ≤3 cycles of 4 once-weekly infusions  Rollover  EFG IV: n=84 PBO: n=83	ADAPT  3 3, OLE  EFG IV 10 mg/kg  ≤3 cycles of 4 once-weekly infusions infusions  EFG IV: n=84 PBO: n=83  EFG IV: n=145	ADAPT  ADAPT+  ADAPT-SC  3 3, OLE 3  EFG IV 10 mg/kg or EFG PH20 SC 1000 mg  ≤3 cycles of 4 once-weekly infusions infusions  Rollover  EFG IV: n=84 PBO: n=83  EFG IV: n=145  EFG IV: n=145  EFG IV: n=55 EFG PH20 SC: n=55 Rollover	ADAPT         ADAPT+         ADAPT-SC         ADAPT-SC+           3         3, OLE         3         3, OLE           EFG IV 10 mg/kg or EFG PH20 SC 1000 mg           ≤3 cycles of 4 once-weekly infusions         ≤19 cycles of 4 once-weekly administrations         ≤11 cycles of 4 once-weekly infusions           Rollover         Rollover         EFG IV: n=55 EFG PH20 SC: n=55 EFG PH20 SC: n=55 Rollover         EFG PH20 SC: n=179	ADAPT         ADAPT+         ADAPT-SC         ADAPT-SC+         ADHERE           3         3, OLE         3         3, OLE         2           EFG IV 10 mg/kg or EFG PH20 SC 1000 mg         EFG PH20 SC 1000 mg           ≤3 cycles of 4 once-weekly infusions         ≤19 cycles of 4 once-weekly administrations         ≤11 cycles of 4 once-weekly injections         Once-weekly injections           EFG IV: n=84 PBO: n=83         EFG IV: n=145         EFG IV: n=55 EFG PH20 SC: n=55 Rollover         EFG PH20 SC: n=111 PBO: 110           26 weeks         ≤3 years         10 weeks         ≤3 years         ≤12 weeks (Stage A)	ADAPT   ADAPT+   ADAPT-SC   ADAPT-SC+   ADHERE   ADVANCE     3   3, OLE   3   3, OLE   2   3     EFG IV 10 mg/kg   or	ADAPT         ADAPT+         ADAPT-SC         ADAPT-SC+         ADHERE         ADVANCE         ADDRESS           3         3, OLE         3         3, OLE         2         3         3           EFG IV 10 mg/kg or EFG PH20 SC 1000 mg         EFG IV 10 mg/kg         EFG IV 10 mg/kg         EFG PH20 SC (2000 mg on 2000 mg on 2000 mg           ≤3 cycles of 4 once-weekly infusions infusions infusions infusions         ≤11 cycles of 4 once-weekly injections         Once-weekly injections         4 once-weekly injections, then once-weekly injections         Once-weekly injections         Once-weekly injections         Once-weekly injections         Once-weekly injections         EFG IV: n=86         PBO: n=75         EFG IV: n=85         EFG IV: n=86         PBO: n=75         EFG IV: n=86

aNot all participants enrolled in these studies received vaccines targeting COVID-19. bSARS-CoV-2-lgG-RBD, -S, and -N antibodies were assessed using an ELISA assay. For COVID-19 vaccinations consisting of multiple injections (ie, injections within 45 days of each other), only blood samples before the first injection and after the last injection were selected. One sample was collected if postvaccination time points (≥4 wk post Vacc at maximum vaccine effect and Max EFG PD) coincided with each other. Samples were collected from the first cycle that occurred after the vaccination.

#### **SUMMARY**



Participants receiving efgartigimod IV or efgartigimod PH20 SC across multiple indications and dosing schedules mounted antigen-specific IgG responses to COVID-19 immunization, even when total IgG levels were maximally reduced



**Effective humoral immune response to COVID-19 vaccination was not precluded** by efgartigimod IV or efgartigimod PH20 SC treatment, regardless of indication or dosing regimen; similar responses were seen with placebo



Additional data on several different vaccines are being retrospectively analyzed from efgartigimod studies across multiple indications

#### **RESULTS**

#### Table 1. Baseline Demographics of Participants in gMG, CIDP, ITP, and PV Studies Receiving COVID-19 Vaccines<sup>a</sup>

Characteristic	<b>gMG</b> (n=71)	<b>CIDP</b> (n=29)	<b>ITP</b> (n=17)	<b>PV</b> (n=31)
Age, y, mean (SD)	49.0 (14.1)	53.3 (13.2)	53.3 (17.1)	50.1 (11.4)
Age category, n (%)				
18-64 y	58 (81.7)	24 (82.8)	13 (76.5)	29 (93.5)
65-74 y	11 (15.5)	4 (13.8)	3 (17.6)	1 (3.2)
≥75 y	2 (2.8)	1 (3.4)	1 (5.9)	1 (3.2)
Sex at birth, n (%)				
Female	47 (66.2)	10 (34.5)	6 (35.3)	15 (48.4)
BMI (kg/m)				
Mean (SD)	28.6 (7.9) <sup>b</sup>	27.5 (4.6)	27.3 (6.0)	27.5 (4.5)
Range (min, max)	18.0, 64.4	18.6, 36.6	17.5, 41.8	19.5, 41.6
<b>Race,</b> n (%)				
Asian	9 (12.7)	4 (13.8)	2 (11.8)	4 (12.9)
Black or African American	-	-	-	1 (3.2)
White	60 (84.5)	21 (72.4)	15 (88.2)	26 (83.9)
Multiple	2 (2.8)	-	-	-
Not reported	-	4 (13.8)	-	-

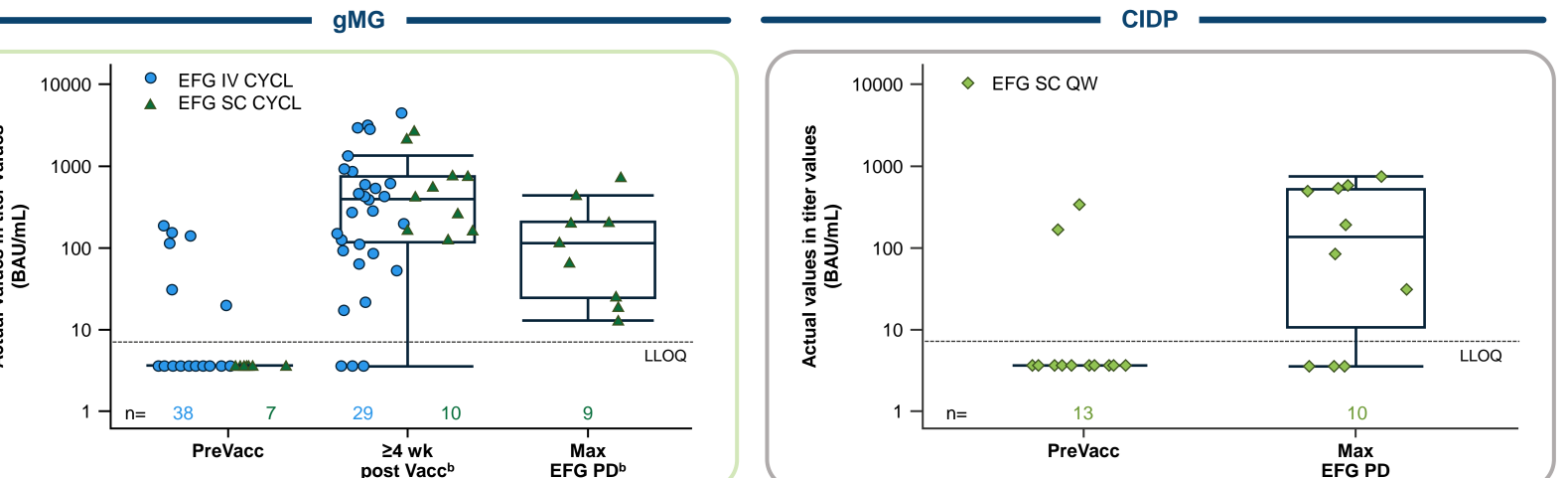
<sup>a</sup>Participant data are included only for those who had a prevaccination titer sample and ≥1 postvaccination titer sample available. <sup>b</sup>BMI data were unavailable for 2 individuals.

#### Table 2. First Documented COVID-19 Vaccination Received by Participants in gMG, CIDP, ITP, and PV Studies<sup>a,b</sup>

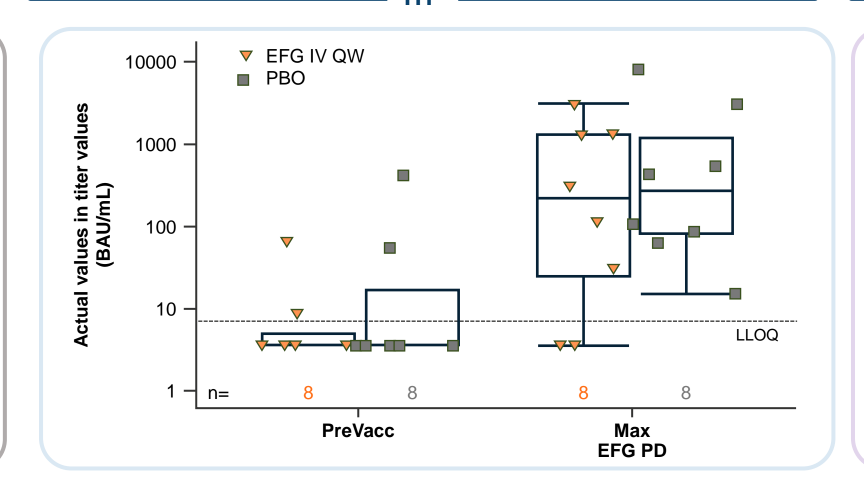
	gMG <sup>c</sup>	CIDPc	ITP		PV	
First COVID-19 Vaccine,d n (%)	EFG IV/SC (n=45)	EFG SC (n=13)	EFG IV (n=8)	<b>PBO</b> (n=8)	<b>EFG SC</b> (n=16)	<b>PBO</b> (n=8)
Pfizer-BioNTech	33 (73.3)	5 (38.5)	5 (62.5)	3 (37.5)	9 (56.3)	5 (62.5)
Unknown	5 (11.1)	2 (15.4)	1 (12.5)	1 (12.5)	4 (25.0)	-
Spikevax (Moderna)	4 (8.9)	2 (15.4)	-	1 (12.5)	2 (12.5)	1 (12.5)
Janssen	1 (2.2)	2 (15.4)	-	1 (12.5)	1 (6.3)	-
Oxford-AstraZeneca	-	1 (7.7)	1 (12.5)	-	-	1 (12.5)
Sputnik V	2 (4.4)	-	-	-	-	1 (12.5)
Sinovac	-	-	-	2 (25.0)	-	-
Sinopharm	-	1 (7.7)	1 (12.5)	-	-	-

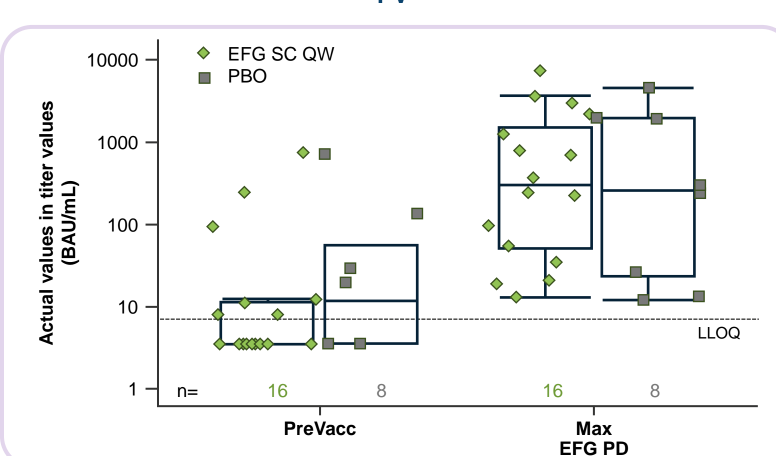
<sup>a</sup>For COVID-19 vaccinations consisting of multiple injections (ie, injections within 45 days of each other), only blood samples before the first injection and after the last injection were selected. bParticipants who had prevaccination titer sample and ≥1 postvaccination titer sample available. Cone participant in the gMG group and 4 participants in the CIDP group treated with placebo received their first COVID-19 vaccination during the study but were excluded from this analysis due to low n values in these groups. Participants who received a COVID-19 vaccination before entering the study are excluded from Table 2.

## Figure 1. Individual SARS-CoV-2-IgG-RBD Titer Values Box Plots After First Documented Vaccination Across Indications



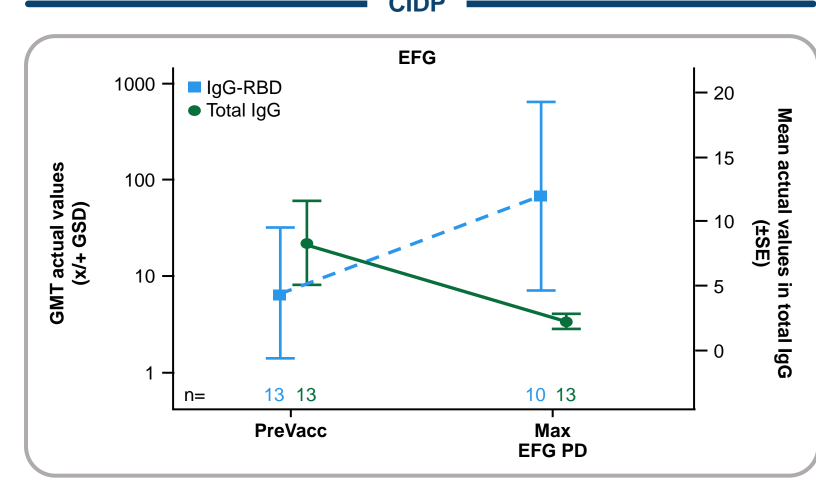


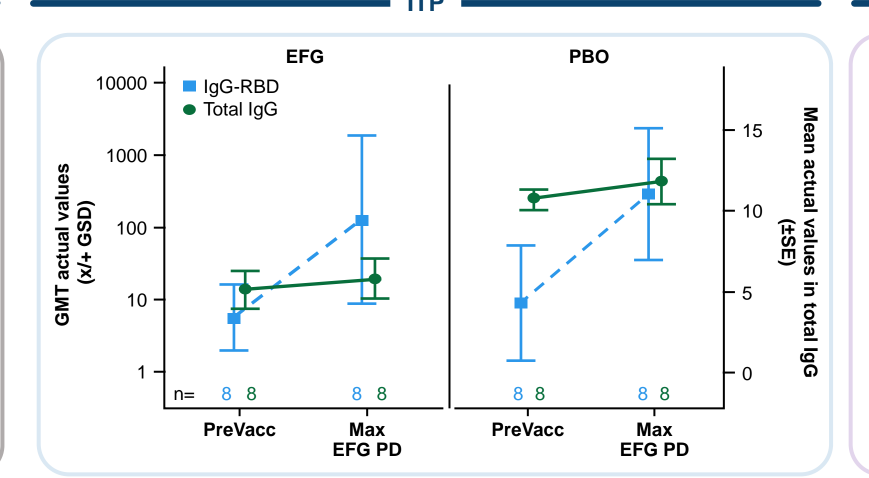


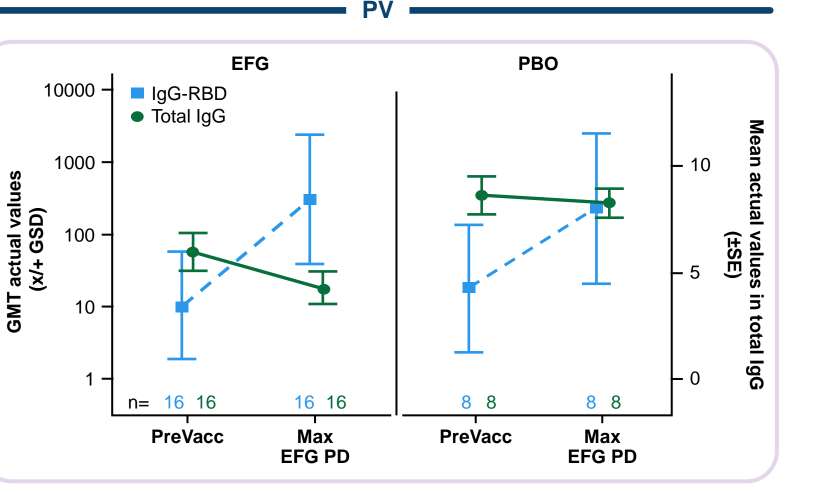


### Figure 2. GMT Actual SARS-CoV-2-IgG-RBD Values and Mean Actual Total IgG Titer Values After First Documented Vaccination Across Indicationsa

## 10000 **│** ■ IgG-RBD Total IgG 1000 -39 13 PreVacc ≥4 wk EFG PDb post Vaccb







<sup>a</sup>For each indication and dosing regimen, only time points with ≥5 samples are presented. <sup>b</sup>If postvaccination time points (≥4 wk post Vacc and Max EFG PD) coincided with each other, the sample is presented at Max EFG PD time point.

#### SARS-CoV-2-IgG-S, SARS-CoV-2-IgG-N, and **SARS-CoV-2-IgG-NEUT Titer Values Across**

- SARS-CoV-2-IgG-S titer values showed a similar trend in response to vaccination as SARS-CoV-2-IgG-RBD titer values
- There was no effect of vaccination on SARS-CoV-2-N titer values (positive SARS-CoV-2-N titer values indicate a previous/resolving COVID-19 infection)12
- SARS-CoV-2-IgG-NEUT titer values increased in response to vaccination, regardless of indication or dosing regimen

BAU, binding antibody units; CIDP, chronic inflammatory demyelinating polyneuropathy; COVID-19, coronavirus disease 2019; EFG, efgartigimod; ELISA, enzyme-linked immunosorbent assay; Fc, fragment crystallizable region; FcRn, neonatal Fc receptor; gMG, generalized myasthenia gravis; GMT, geometric mean titer; Ig, immunoglobulin; ITP, immune thrombocytopenia; IV, intravenous; LLOQ; lower limit of quantification; MG, myasthenia gravis; -N, nucleocapsid protein; -NEUT, neutralizing antibodies; OLE, open-label extension; PBO, placebo; PD, pharmacodynamic; PreVacc, prevaccination; PV, pemphigus vulgaris; QW, once a week; Q2W, every other week; -RBD, receptor-binding domain of S protein; rHuPh20, recombinant human hyaluronidase PH20; -S, spike protein; SARS-CoV-2-IgG, severe acute respiratory syndrome coronavirus 2 specific IgG; SC, subcutaneous.

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