Safety of Ocrelizumab in Multiple Sclerosis: Updated Analysis in Patients with Relapsing and Progressive Multiple Sclerosis



SL Hauser,¹ L Kappos,² X Montalban,³ C Chognot,⁴ N Pasquarelli,⁴ K Kadner,⁴ B El Azzouzi,⁴ A Pradhan,⁵ E Incera,⁶ JS Wolinsky⁷

¹UCSF Weill Institute for Neurosciences, University of California San Francisco, San Francisco, CA, USA; ²Research Center for Clinical Neuroimmunology and Neuroscience, University Hospital Basel, University of Basel, Basel, Switzerland; ³Department of Neurology-Neuroimmunology, Multiple Sclerosis Centre of Catalonia, Vall d'Hebron University Hospital, Barcelona, Spain; ⁴F. Hoffmann-La Roche Ltd, Basel, Switzerland; ⁵Genentech, Inc., South San Francisco, CA, USA; ⁶IQVIA Solutions, Inc., Courbevoie, France; ⁷Department of Neurology, McGovern Medical School, The University of Texas Health Science Center at Houston (UTHealth), Houston, TX, USA

STUDY AIM

To examine safety outcomes of patients continuously treated with OCR in clinical trials over a 10-year period (up to November 2022)

CONCLUSIONS

Over a 10-year follow-up period in clinical trials, OCR continues to exhibit a stable and favourable safety profile

- AE rates in all RMS and PMS populations remained consistent with the rates observed during the CTP
- Rates of SIs remained low and stable over time in both RMS and PMS populations irrespective of IgG levels
- AEs leading to treatment withdrawal remained infrequent and were not driven by serious infections

AE, adverse event; CTP, controlled-treatment period; IgG, immunoglobulin G; OCR, ocrelizumab; PMS, progressive multiple sclerosis; RMS, relapsing multiple sclerosis, SI, serious infections

RESULTS



- As of November 2022, 6,155 patients with MS received OCR across 13 clinical trials (all-exposure population), amounting to 28,269 PY of exposure
- Approximately 60% of patients (3,677) received at least 8 doses (~4 years of treatment)
- As of March 2023, over 300,000 patients with MS (amounting to >750,000 PY of exposure) had started OCR globally

Over 10 Years of Continuous OCR Treatment, the Overall Safety Profile Remained Consistent

Adverse event Rate per 100 PY (95% CI)	OPERA (RMS)			All RMS°	ORATORIO (PPMS)			All PMSd	All OCR trials
	CTP ^a (Jul 2015)		CTP + OLE ^b (Nov 2022)	Nov 2022	Nov 2022 CTP ^a (Jul 2015)		CTP + OLE ^b (Nov 2022)	Nov 2022	OCR all-exposure
	IFN β-1a	OCR	OCR	OCR	Placebo	OCR	OCR	OCR	populatione
Total no. of patients	826	825	1,448	4,558	239	486	644	1,597	6,155
Total PY	1,399	1,448	10,798	21,080	729	1,606	4,669	7,190	28,269
Any AEs	296 (287–305)	290 (281–299)	194 (191–196)	227 (225–229)	259 (247–271)	252 (244–260)	223 (219–228)	215 (212–219)	224 (222–226)
AEs leading to withdrawal	3.9 (3.0–5.1)	2.4 (1.6–3.3)	1.3 (1.1–1.5)	1.0 (0.9–1.2)	1.1 (0.5–2.2)	1.2 (0.8–1.9)	1.1 (0.8–1.4)	1.0 (0.8–1.3)	1.0 (0.9–1.2)
Serious AEs	6.3 (5.1–7.8)	5.4 (4.3–6.7)	6.2 (5.8–6.7)	5.7 (5.4–6.0)	12.1 (9.7–14.9)	10.2 (8.7–11.8)	12.7 (11.7–13.8)	10.9 (10.1–11.7)	7.0 (6.7–7.3)
Infections and infestations	67.8 (63.5–72.2)	84.5 (79.9–89.4)	65.9 (64.4–67.4)	66.2 (65.1–67.3)	72.5 (66.5–79.0)	70.8 (66.8–75.0)	70.0 (67.8–72.6)	61.6 (59.8–63.4)	65.1 (64.1–66.0)
Serious infections ^f	1.8 (1.2–2.6)	0.8 (0.4–1.5)	1.7 (1.5–2.0)	1.5 (1.3–1.7)	3.0 (1.9–4.6)	2.7 (2.0–3.7)	4.4 (3.8–5.0)	3.7 (3.3–4.2)	2.1 (1.9–2.2)
IRRs	7.9 (6.5–9.5)	34.9 (31.9–38.1)	11.2 (10.5–11.8)	23.2 (22.6–23.9)	20.3 (17.2–23.8)	31.0 (28.3–33.9)	16.6 (15.5–17.8)	16.8 (15.9–17.8)	21.6 (21.1–22.2)
Malignancies ^{g,h}	0.1 (0.0–0.5)	0.3 (0.1–0.7)	0.4 (0.3–0.6)	0.4 (0.3–0.4)	0.3 (0.0–1.0)	0.9 (0.5–1.5)	1.0 (0.7–1.3)	0.9 (0.7–1.2)	0.5 (0.4–0.6)
Deaths	0.1 (0.0–0.5)	0.1 (0.0–0.4)	0.1 (0.0–0.2)	0.1 (0.1–0.2)	0.4 (0.1–1.2)	0.3 (0.1–0.6)	0.5 (0.3–0.7)	0.4 (0.3–0.6)	0.2 (0.1–0.2)

COVID-19 related AEs were excluded from this analysis, but patients continued to contribute to the incidence of all other AEs (see Supplementary Material for COVID-19 analysis). AEs were classified according to MedDRA versions 18.0, 18.1, 22.1 and 24.1. Multiple occurrences of the same AE in one patients are counted multiple times, except for malignancies. [®]Data as of April–July 2015; [®]Includes patients who received any dose of OCR during the CTP and associated OLE periods of the Phase III studies plus VELOCE, CHORDS, CASTING, OBOE, ENSEMBLE, LIBERTO, CHIMES and OLERO (data as of November 2022); [®]Includes patients with PMS who received any dose of OCR during the CTP and associated OLE periods of OBOE, CONSONANCE and OLERO (data as of November 2022); [®]Includes patients who received any dose of OCR during the CTP and associated OLE periods of OBOE, CONSONANCE and OLERO (data as of November 2022); [®]Includes patients who received any dose of OCR during the CTP and associated OLE periods of OBOE, CONSONANCE and OLERO (data as of November 2022); [®]Includes patients who received any dose of OCR during the CTP and associated OLE periods of OBOE, CONSONANCE and OLERO (data as of November 2022); [®]Includes patients with PMS who received any dose of OCR during the CTP and associated OLE periods of OBOE, CONSONANCE and OLERO (data as of November 2022); [®]Includes patients with PMS who received any dose of OCR during the CTP and associated OLE periods of OBOE, CONSONANCE and OLERO (data as of November 2022); [®]Includes patients with PMS who received any dose of OCR during the CTP and associated OLE periods of OBOE, CONSONANCE and OLERO (data as of November 2022); [®]Includes patients with PMS who received any dose of OCR during the CTP and associated OLE periods of OBOE, CONSONANCE and OLERO (data as of November 2022); [®]Includes patients who received any dose of OCR during the CTP and associated OLE periods of OBOE, CONSONANCE and OLERO (data as of November 2022); [®]Includes patients who received any dose



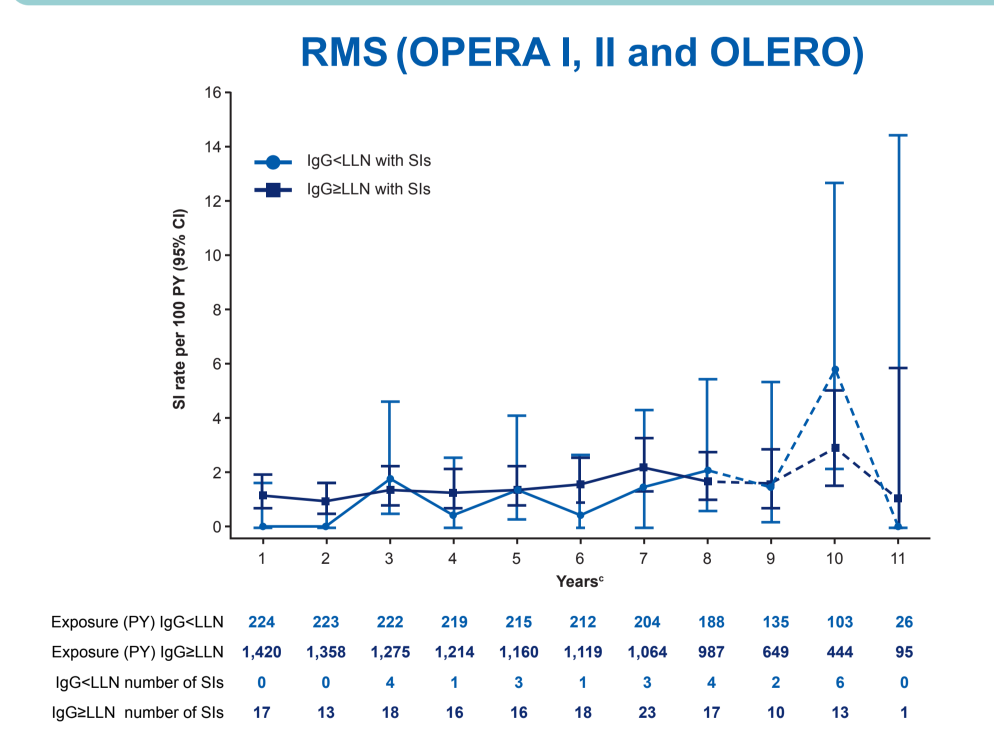
- Cumulative AE and SAE incidence rates remained consistent with the rates observed during the CTP
- The cumulative standardised incidence rates of (a) all malignancies and (b) female breast cancer remained within the range reported in real-world registries^{1,2} (see Supplementary Material)
- Withdrawal due to AEs was infrequent and did not increase over time

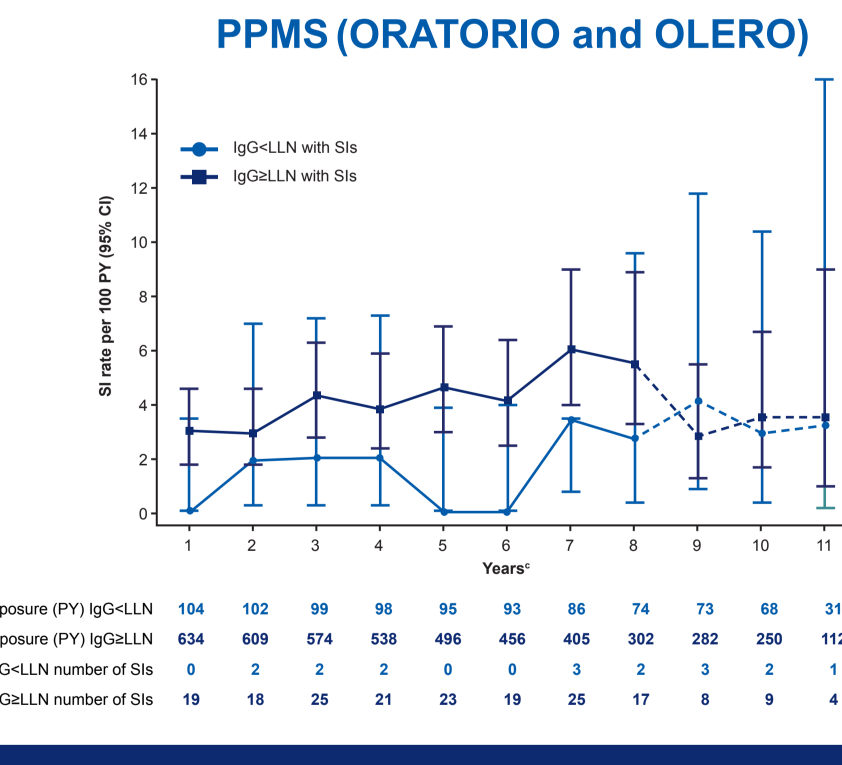
AE, adverse event; CTP, controlled treatment period; SAE, serious adverse event.

SI Rates Remained Stable with Non-Significant Year-on-Year Variation and Within the Range Reported in Real-World Registries^{3,4}

In Both RMS and PPMS Populations, Longer Exposure to OCR Did Not Lead to an Increased Risk of SIs Regardless of IgG Status (Normal Levels or Levels Below the LLN)

RMSa and PMSb all-exposure population PMS all-exposure PMS all-exposure RMS all-exposure RMS all-exposure PMS all-e



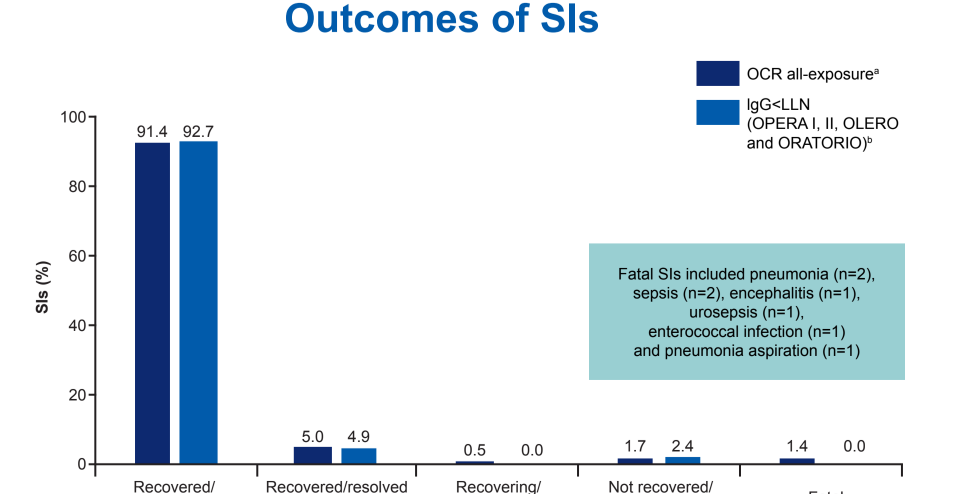


المراجعة الم

- In the RMS and PMS all-exposure populations, UTI and pneumonia were the most commonly reported SIs; this is consistent with incidence rates and patterns observed in real-world studies⁴⁻⁶
- The type, severity, latency and duration of SIs observed during episodes of IgG<LLN were consistent with the overall SIs observed in patients treated with OCR

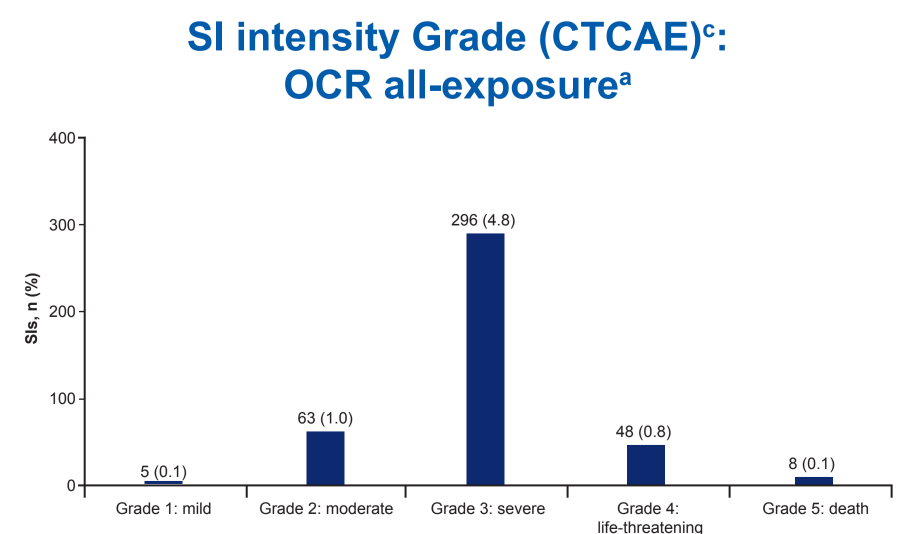
COVID-19 related AEs were excluded from this analysis, but patients continued to contribute to the incidence of all other AEs (see Supplementary Material for COVID-19 analysis). and expected any dose of OCR during the CTP and associated OLE periods of the Phase II and Phase III studies plus VELOCE, CHORDS, CASTING, OBOE, ENSEMBLE, LIBERTO, CHIMES and OLERO (data as of November 2022); black a

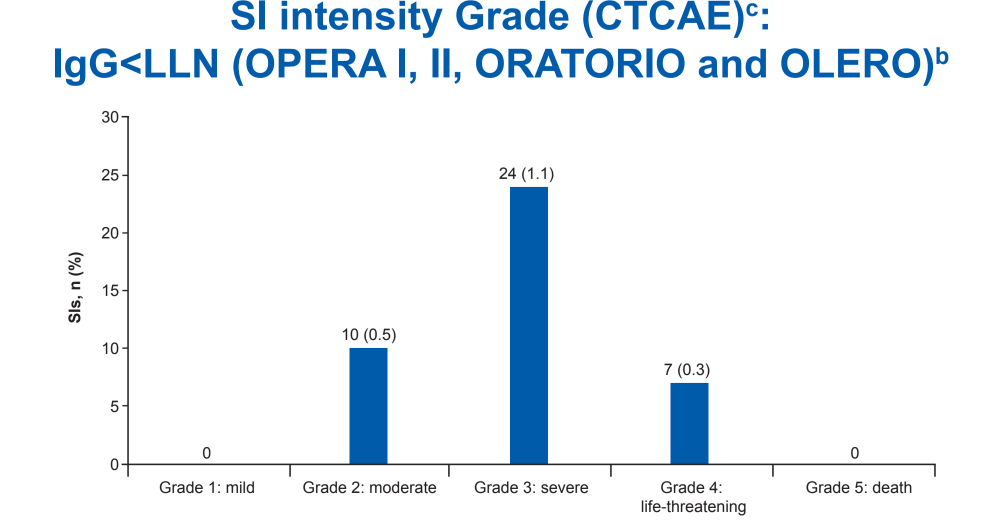
The Majority of SIs Were of Grade 3 Intensity, >90% Had Resolved (Regardless of IgG Status) and Were Not Treatment Limiting



resolving

not resolved





COVID-19 related AEs were excluded from this analysis, but patients continued to contribute to the incidence of all other AEs (see Supplementary Material for COVID-19 analysis). alncludes patients who received any dose of OCR during the CTP and associated OLE periods of the Phase II and Phase III studies plus VELOCE, CHORDS, CASTING, OBOE, ENSEMBLE, LIBERTO, CONSONANCE, CHIMES and OLERO (total N=6,155 patients); biggle-drop method (total N=2,092); crade 1 (mild): Asymptomatic or mild symptoms/clinical or diagnostic observation only/intervention indicated/limiting age-appropriate instrumental ADL; Grade 3 (severe): Severe or medically insignificant but not immediately life-threatening/hospitalisation or prolongation of hospitalisation indicated/disabling/limiting self-care ADL; Grade 4 (life-threatening): Life-threatening): Life-threatening consequences/urgent intervention required; Grade 5 (death): Death related to AE (not applicable for all AEs); percentages are calculated using the number of patients that experienced any number of SIs (counting multiple occurrences once), by the number of patients. ADL, activities of daily living; CTCAE, Common Terminology Criteria for Adverse Events; IgG, immunoglobulin G; LLN: Lower limit of normal [LLN=5.65 (g/L)]; OCR, occelizumab; SI, serious infection.

REFERENCES
 National Institutes of Health (NIH). Surveillance, Epidemiology, and End Results Program. Available from: https://seer.cancer.gov.
 Nørgaard M, et al. Mult Scler Relat Disord 2019;28:81–85.
 Wijnands JMA, et al. J Neurol Neurosurg Psychiatry 2018;89:1050–1056.

4. Knapp R, et al. Mult Scler Relat Disord 2022;68:104245

6. Persson et al. Mult Scler Relat Disord 2020;41(1):101982.

5. Wijnands JMA, et al. Mult Scler 2017;23:1506–1516.

resolved

SL Hauser serves on the Board of Trustees for Neurona and on scientific advisory boards for Alector, Annexon and Accure; and has received travel reimbursement and writing assistance from F. Hoffmann-La Roche Ltd and Novartis for CD20-related meetings and presentations. **L Kappos'** institution (University Hospital Basel) has received the following exclusively for research support: Steering committee, advisory board and consultancy fees (Actelion, Bayer HealthCare, Biogen, Bristol Myers Squibb, Sanofi-Genzyme, Janssen, Japan Tobacco, Merck, Novartis, F. Hoffmann-La Roche Ltd, Sanofi, Santhera, Shionogi and TG Therapeutics); speaker fees (Bayer HealthCare, Biogen, Merck, Novartis, F. Hoffmann-La Roche Ltd, Pfizer, Sanofi, Shire and Teva); license fees for Neurostatus products; and grants (Bayer HealthCare, Biogen, European Union, Innosuisse, Merck, Novartis, F. Hoffmann-La Roche Ltd, Swiss MS Society and Swiss National Research Foundation). **X Montalban** has received speaking honoraria and travel expenses for participation in scientific meetings, has been a steering committee member of clinical trials or participated in advisory boards of clinical trials or participated in advisory boards with AbbVie, Actelion, Alexion, Alexion, Biogen, Bristol Myers Squibb/ Celgene, EMD Serono, F. Hoffmann-La Roche Ltd, Immunic, Janssen, MedDay, Merck, Mylan, NervGen, Novartis, Sandoz, SanofiGenzyme, Teva, TG Therapeutics, ExeMed, MSIF and NMSS. **C Chognot** is a shareholder and employee of F. Hoffmann-La Roche Ltd. **K Kadner** is an employee of F. Hoffmann-La Roche Ltd. **K Kadner** is an employee of F. Hoffmann-La Roche Ltd. **A Pradhan** is an employee of Genentech, Inc. **E Incera** has received consulting fees from F. Hoffmann-La Roche Ltd for statistical assistance, and is an employee of IQVIA Solutions, Inc. **JS Wolinsky** has received personal compensation for consulting, serving on a scientific advisory board, speaking or other activities with Cleveland Clinic Foundation, EMD Serono,

ACKNOWLEDGEMENTS

We would like to thank all patients, their families and the investigators participating in this study. This study is sponsored by F. Hoffmann-La Roche Ltd, Basel, Switzerland. Writing and editorial assistance for this

Hoffmann-La Roche Ltd, Basel, Switzerland.

following URL:

To access this poster and Supplementary Material scan

the QR code or go to the

presentation was provided by Articulate Science, UK, and funded by F.

with sequelae