



Association between T2-related co-morbidities and effectiveness of biologics in severe asthma

Michael E Wechsler, Ghislaine Scelo, PhD, Désirée E.S. Larenas-Linnemann, Carlos A Torres-Duque, Jorge Maspero, Trung N Tran, Ruth B Murray, Neil Martin, Andrew N Menzies-Gow, Mark Hew, Matthew J Peters, Peter G Gibson, George C Christoff, Todor A Popov, Andréanne Côté, Celine Bergeron, Delbert Dorscheid, J Mark FitzGerald, Kenneth R. Chapman, Louis Philippe Boulet, Mohit Bhutani, Mohsen Sadatsafavi, Libardo Jiménez-Maldonado, Mauricio Duran-Silva, Bellanid Rodriguez, Carlos Andres Celis-Preciado, Diana Jimena Cano-Rosales, Ivan Solarte, Maria Jose Fernandez- Sanchez, Patricia Parada-Tovar, Anna von Bülow, Anne Sofie Bjerrum, Charlotte S Ulrik, Karin Dahl Assing, Linda Makowska Rasmussen, Susanne Hansen, Alan Altraja, Arnaud Bourdin, Camille Taille, Jeremy Charriot, Nicolas Roche, Andriana I Papaioannou, Konstantinos Kostikas, Nikolaos G Papadopoulos, Sundeep Salvi, Deirdre Long, Patrick D Mitchell, Richard Costello, Concetta Sirena, Cristina Cardini, Enrico Heffler, Francesca Puggioni, Giorgio Walter Canonica, Giuseppe Guida, Takashi Iwanaga, Mona Al-Ahmad, Ulises Garcia, Piotr Kuna, João A Fonseca, Riyad Al-Lehebi, Mariko S Koh, Chin Kook Rhee, Borja G Cosio, Luis Perez de Llano, Diahn-Wang Perng, Erick Wan-Chun Huang, Hao-Chien Wang, Ming-Ju Tsai, Bassam Mahboub, Laila Ibraheem Jaber Salameh, David J. Jackson, John Busby, Liam G Heaney, Paul E. Pfeffer, Amanda Grippen Goddard, Eileen Wang, Flavia C.L. Hoyte, Nicholas M Chapman, Rohit Katial, Victoria Carter, Lakmini Bulathsinhala, Neva Eleangovan, Con Ariti, Juntao Lyu, Celeste Porsbjerg, and David B. Price



Rationale

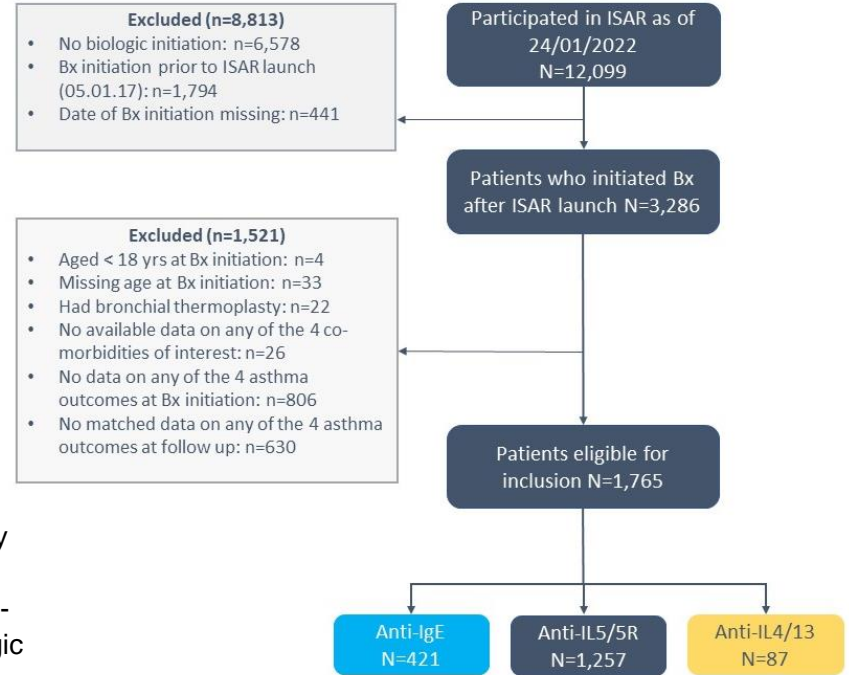
Previous studies investigating comorbidity impact on biologic effectiveness have been relatively small, of short duration, and have not compared biologic classes

Aim

To determine the association between T2-related comorbidities and biologic effectiveness in adults with severe asthma (SA)

Methods

This cohort study used ISAR data (n=21 countries, 2017-2022) to quantify pre- to post-biologic change for four outcomes (annual asthma exacerbation rate, % predicted FEV1 (ppFEV1), asthma control, and long-term oral corticosteroid daily dose [LTOCS]) in patients with/without allergic rhinitis (AR), chronic rhinosinusitis +/- nasal polyps (CRS+/-NP), NP, or eczema/atopic dermatitis (AD).



Irrespective of T2 comorbidities all groups showed improvement but greater in those with CRS+/- NP

Asthma-related outcome	Allergic rhinitis		Chronic rhinosinusitis		Nasal polyposis		Eczema/atopic dermatitis	
	Ever N=761	Never N=493	Ever N=968	Never N=748	Ever N=636	Never N=1120	Ever N=243	Never N=1510
Exacerbation rates: mean (SD)	N=559	N=363	N=719	N=541	N=463	N=818	N=189	N=1092
Pre-biologics	2.24 (2.34)	2.16 (2.23)	2.65 (2.77)	3.37 (3.74)	2.88 (3.02)	3.05 (3.40)	1.97 (2.00)	3.15 (3.39)
Post-biologics	0.65 (1.21)	0.65 (1.04)	0.75 (1.25)	1.13 (1.62)	0.77 (1.21)	1.01 (1.55)	0.72 (1.35)	0.96 (0.46)
Change	-1.59 (2.54)	-1.51 (2.33)	-1.89 (2.74)	-2.24 (3.51)	-2.11 (2.82)	-2.04 (3.30)	-1.25 (2.30)	-2.19 (3.22)
p-value*	<0.001	<0.001	<0.001	<0.001	<0.001	<0.001	<0.001	<0.001
ppFEV₁: mean (SD)	N=313	N=267	N=493	N=386	N=306	N=573	N=101	N=776
Pre-biologics	76.4 (21.7)	72.2 (23.3)	75.8 (22.5)	71.0 (22.6)	76.4 (22.1)	72.2 (22.9)	73.9 (22.5)	73.6 (22.7)
Post-biologics	80.1 (22.6)	76.6 (23.2)	79.5 (23.3)	73.0 (22.1)	79.7 (23.0)	75.1 (22.8)	75.6 (21.7)	76.8 (23.1)
Change	+3.7 (17.9)	+4.4 (16.0)	+3.8 (17.1)	+2.0 (17.1)	+3.3 (17.1)	+2.9 (17.1)	+1.7 (13.7)	+3.1 (17.5)
p-value*	<0.001	<0.001	<0.001	0.023	0.001	<0.001	0.210	<0.001
Asthma control: % of uncontrolled/ partly controlled/well controlled	N=430	N=237	N=570	N=450	N=414	N=629	N=118	N=923
Pre-biologics	65.6/22.6/11.9	57.8/23.2/19.0	65.8/21.2/13.0	69.6/18.9/11.6	65.2/21.3/13.5	70.3/18.6/11.1	71.2/19.5/9.3	67.8/19.7/12.5
Post-biologics	25.6/31.9/42.6	27.0/29.1/43.9	30.2/26.5/43.3	42.4/25.3/32.2	29.5/24.9/45.7	39.6/27.2/33.2	39.0/33.1/28.0	35.2/25.4/39.4
p-value*	<0.001	<0.001	<0.001	<0.001	<0.001	<0.001	<0.001	<0.001
LTOCS								
Users, n (%)	283 (37.2)	202 (41.0)	445 (46.0)	383 (51.2)	312 (49.1)	543 (48.5)	243 (33.3)	772 (51.1)
LTOCS: mean daily dose in users pre-biologics (SD)	N=128	N=74	N=243	N=262	N=196	N=332	N=42	N=485
Pre-biologics	13.2 (10.9)	15.5 (15.4)	12.2 (10.0)	13.2 (10.6)	12.0 (9.3)	13.1 (10.7)	10.5 (10.1)	12.8 (10.2)
Post-biologics	11.7 (9.9)	13.9 (14.7)	10.5 (9.5)	11.0 (10.1)	9.8 (8.3)	11.4 (10.4)	8.8 (9.0)	10.9 (9.8)
Change	-1.4 (7.6)	-1.6 (11.7)	-1.7 (6.9)	-2.2 (7.6)	-2.2 (7.2)	-1.7 (7.1)	-1.7 (8.9)	-1.9 (7.0)
p-value*	0.020	0.204	<0.001	<0.001	<0.001	<0.001	0.116	<0.001

Exacerbation rates reduce (reduction shown in red) with biologics for all, regardless of presence of T2 comorbidity

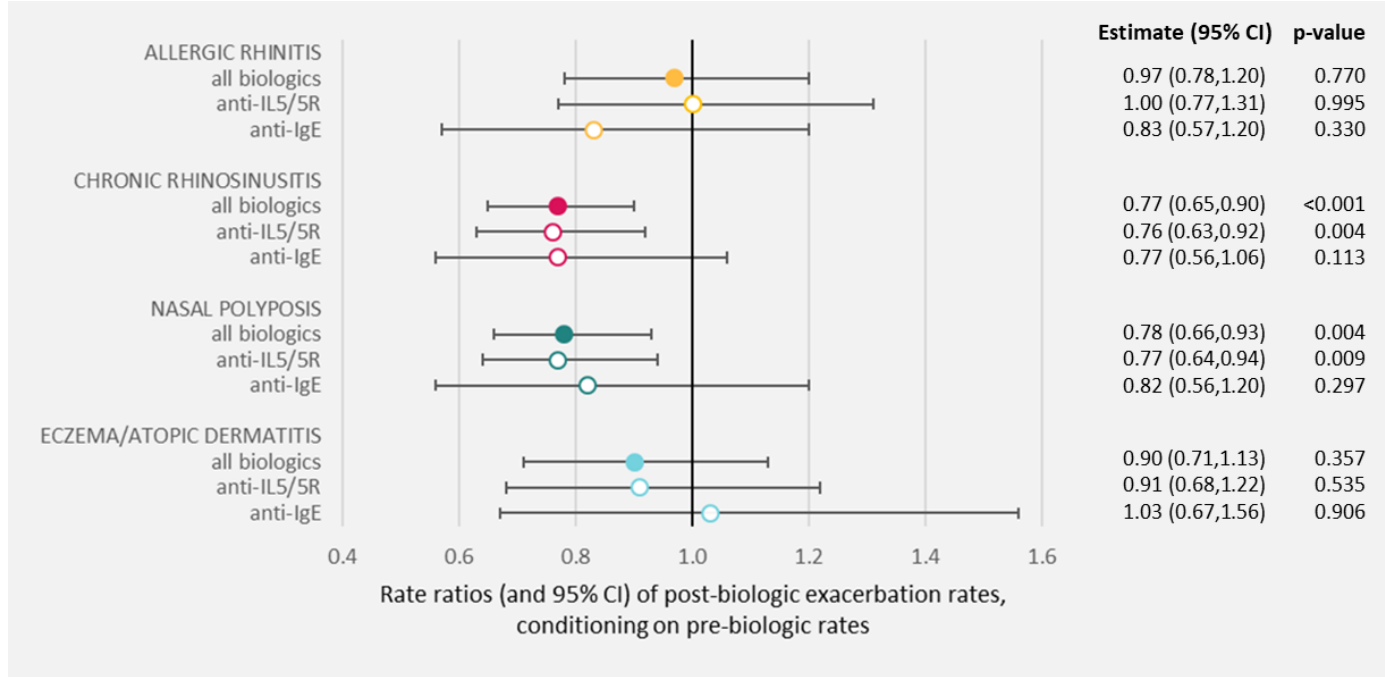
ppFEV₁ increases (increase shown in red) for all following biologic initiation, irrespective of comorbidity status

%of patients with uncontrolled asthma decreases significantly (shown in red) across all groups irrespective of comorbidity status, following biologic initiation

Figures in red show the drop in mean LTOCS dose following biologic initiation, with reduction in dose achieved for all groups irrespective of presence of T2 comorbidities

*Comparing pre- to post-biologics, using paired Wilcoxon test for exacerbations and LTOCS dose, paired t-test for ppFEV₁, and McNemar test (nominal symmetry test) for asthma control.

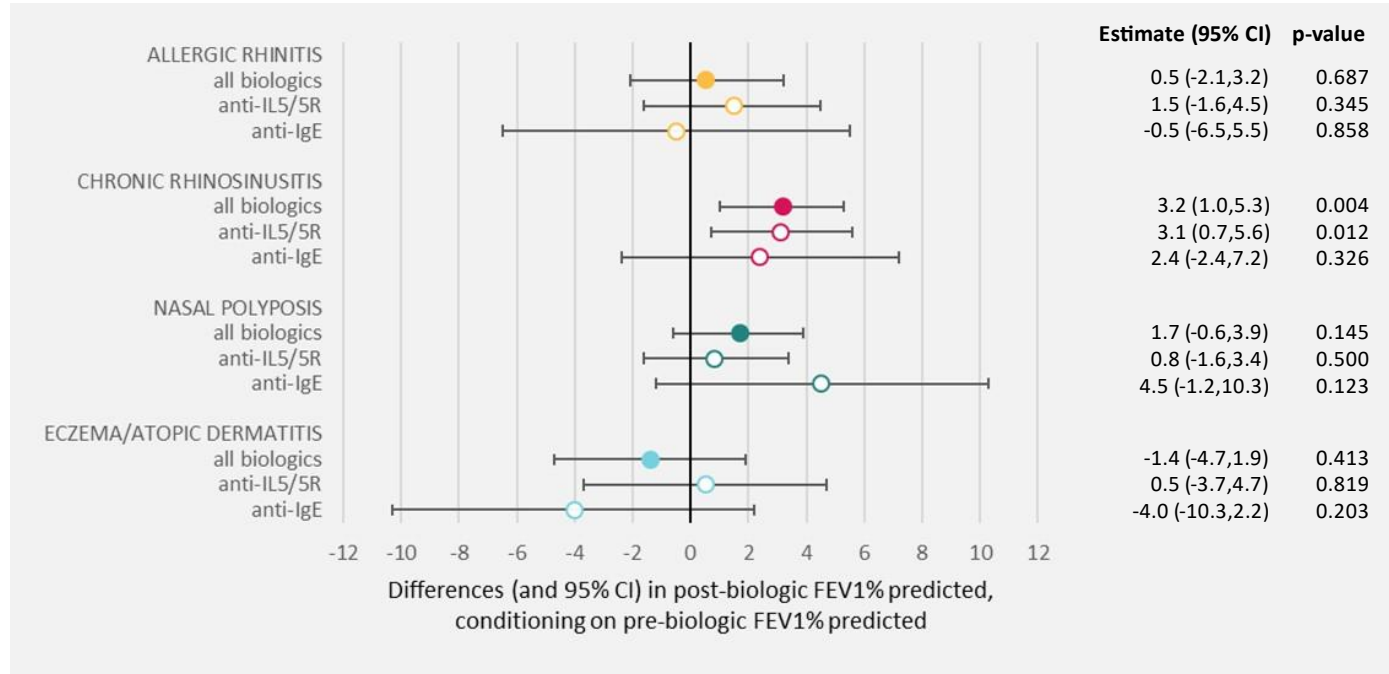
Severe asthma patients with CRS / NP benefit from biologics to a greater extent than patients without: **exacerbations**



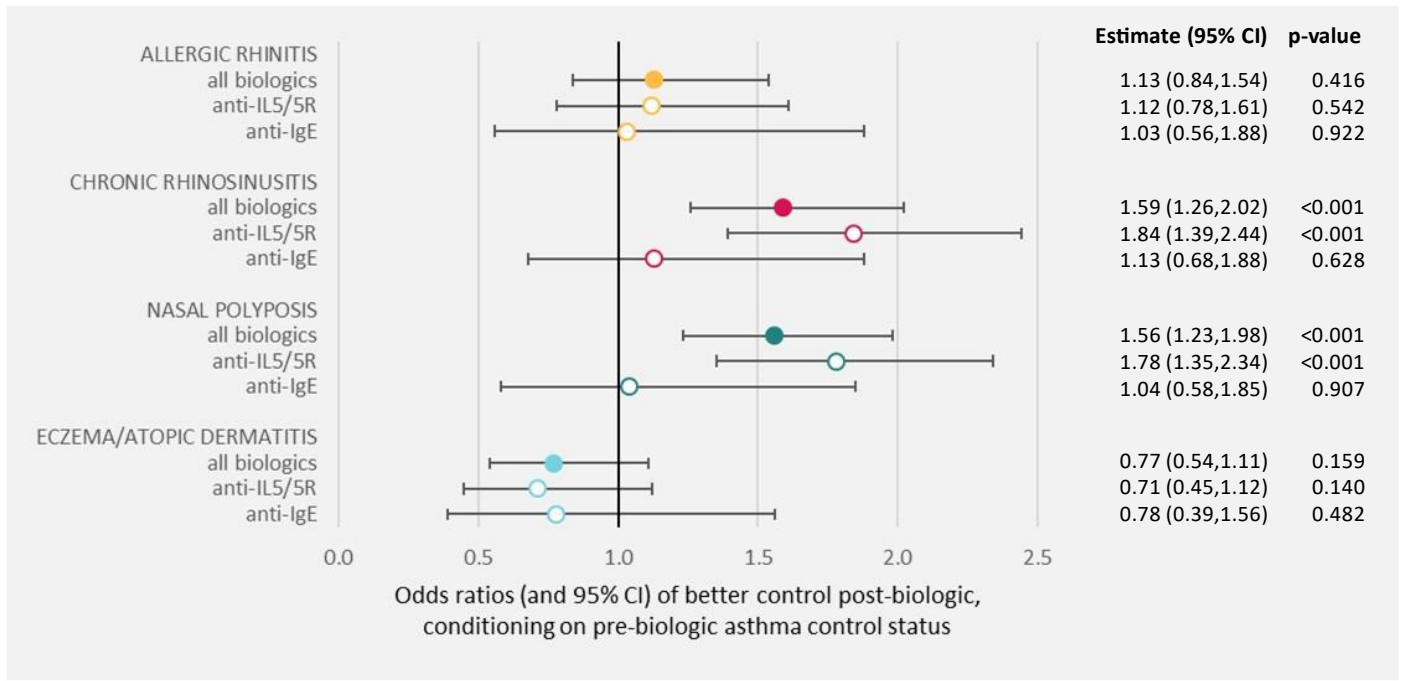
 Biomarker independent

Adjusting for BEC had no impact on the estimate for exacerbations for patients with CRS +/- NP (rate ratio = 0.77, 95% CI: 0.65, 0.91, p=0.002)

Severe asthma patients with CRS / NP benefit from biologics to a greater extent than patients without: lung function



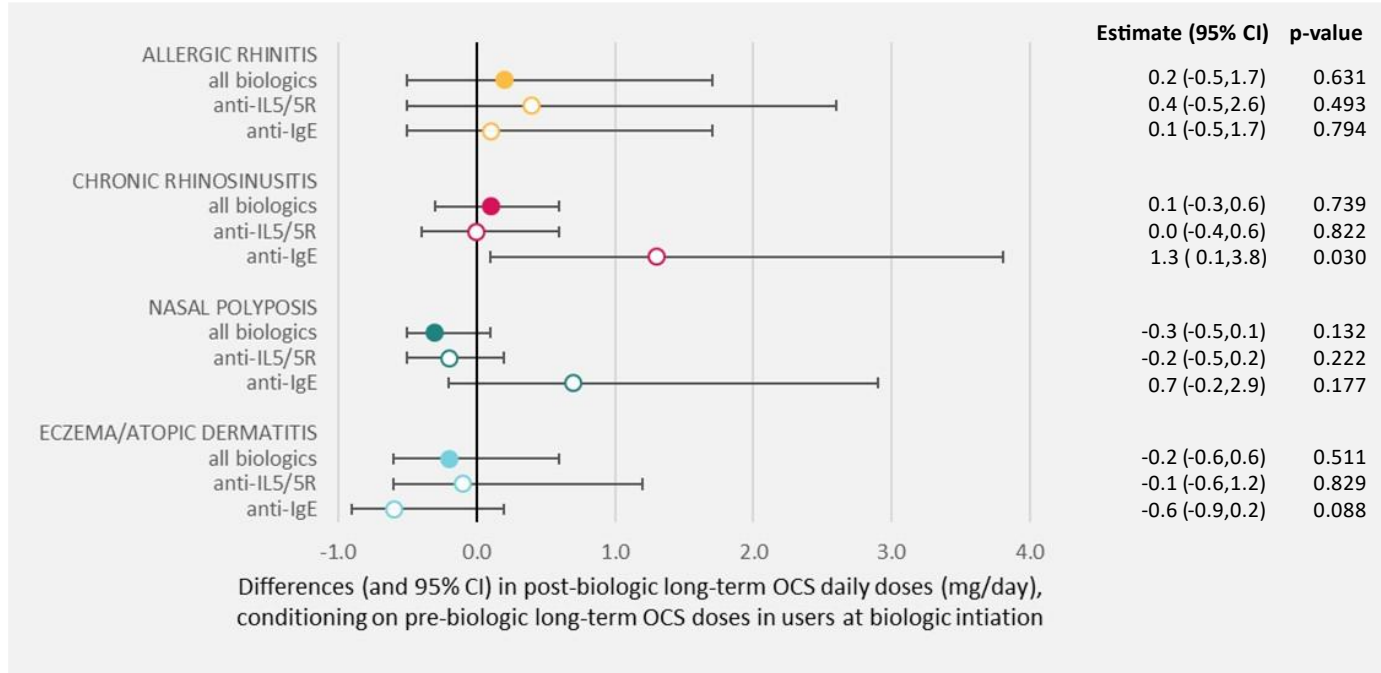
Severe asthma patients with CRS / NP benefit from biologics to a greater extent than patients without: **asthma control**



Biomarker independent

Although attenuated, association for asthma control trends in patients with NP remain when adjusting for BEC (odds ratio=1.37, 95% CI: 1.06-1.77), p=0.015)

No additional benefit observed in terms of LTOCS dose reduction



Association between T2-related co-morbidities and effectiveness of biologics in severe asthma

Where

ISAR Global Study

21 countries + 1765 severe asthma patients



Who



With/without AR, CRS+/-NP, NP, or AD



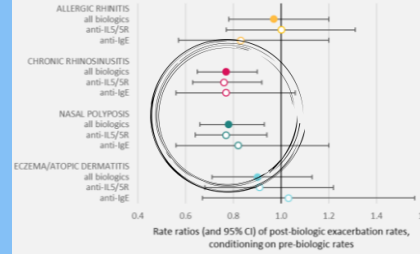
Initiated on anti-IL-5/5R, anti-IgE, or anti-IL-4/13

What

Pre and post biologic change in:
Annual **exacerbation rate**,
% predicted **FEV1**
asthma control
LTOCS daily dose



Results



- Biologics led to **improvements in all four asthma outcomes irrespective of comorbidity status**
- **Comorbid CRS+/-NP : 23% fewer exacerbations, 59% higher odds of better post-biologic control**

- Similar estimates for those with comorbid NP
- **Independent of biomarker profile**
- **AR and AD conversely were not predictive** of treatment effect

Practice change

- ✓ **CRS +/- NP key components in predicting successful treatment with biologics**
- ✓ **Systematic evaluation for comorbidities + multidisciplinary collaboration vital** in achieving optimal outcomes in severe asthma care