

EVALUATION OF TREATMENT PATTERNS AMONG CROHN'S DISEASE PATIENTS INITIATING BIOLOGICS WITH THREE YEARS OF FOLLOW-UP

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BACKGROUND

Crohn's disease (CD) is a chronic inflammatory disease of the gastrointestinal tract, which is associated with high healthcare resource utilization, and high healthcare costs resulting in estimated medical costs of \$3.48 billion in the United States in 2016.^{1,2}

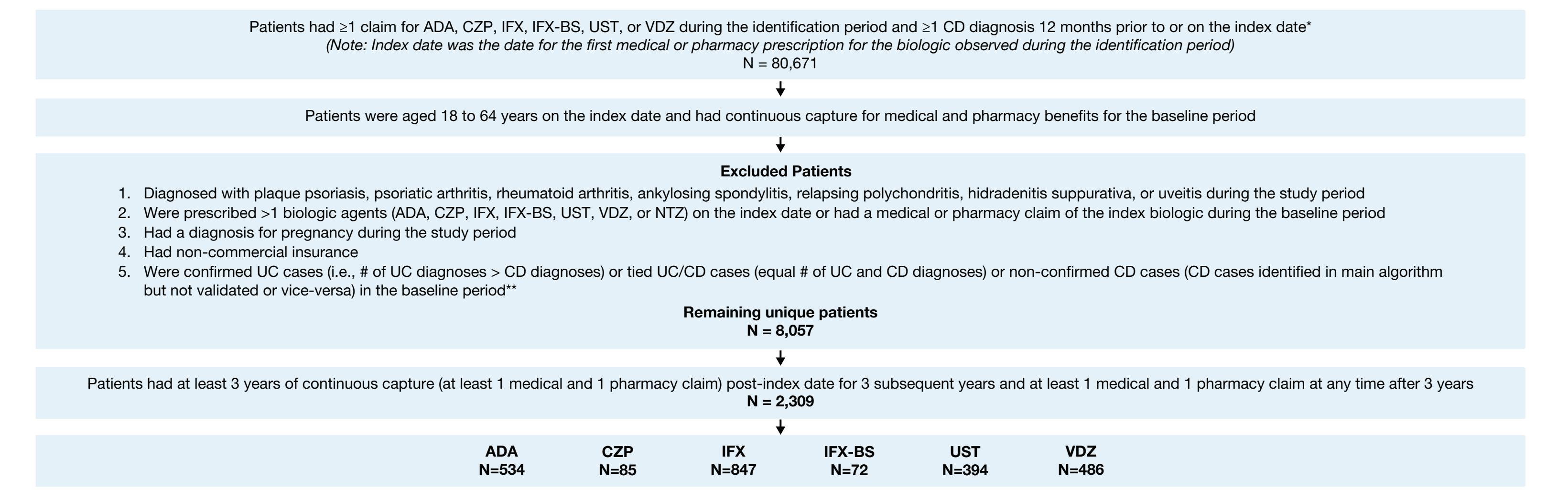
Many new biologic therapies have been recently approved for CD. However, there is limited real-world evidence on treatment patterns, and dose titration among CD patients using biologics with 3 years of follow-up post-initiation of index biologics.

OBJECTIVE

This descriptive study examined biologic treatment patterns among Crohn's Disease (CD) patients initiating biologics with 3 years of follow-up. We examined the CD patients treated with adalimumab (ADA), certolizumab pegol (CZP), infliximab (IFX) and its biosimilar products (IFX-BS), ustekinumab (UST), and vedolizumab (VDZ).

METHODS

Figure 1. Patient Selection Criteria



Baseline and outcome variables

- Baseline variables: age, sex, US geographic region, Charlson comorbidity index (CCI) score, and baseline comorbidities.

- Outcome variables: The following outcomes were examined during the follow-up period.

1. **Persistence:** Proportion of patients who remained on the index biologic without a gap of >60 days for ADA and >120 days for IFX, UST, and VDZ (gapped days were approximately two times of the United States Food and Drug Administration (FDA) labeled maintenance dosing interval) between the run-out date of two consecutive biologic claims were considered persistent to their index biologic.

2. **Switch:** Patients who were administered a non-index biologic during follow-up.

3. **Restart:** Patients who restarted their index biologic (after the gap of >60 days for ADA and >120 days for IFX, UST, and VDZ) during follow-up.

4. **Discontinuation:** Patients who were not persistent to their index biologic were classified as discontinuers.

- **Discontinuation without restart/switch:** Patients who were not administered/prescribed any biologics after the discontinuation date (defined as the run-out date of the last index medication claim or the switch date, whichever occurred first) without any restart or switch until the end of follow-up.

CONCLUSION

During the 3 years of follow-up, unadjusted persistence was highest in the UST cohort. In addition, the UST cohort had the numerically lowest proportion of patients with dose titration but the numerically highest proportion of patients with dose-reduction. The IFX cohort had the numerically highest proportion of patients with >100% dose escalation and UST had the lowest.

RESULTS

• A total of 2,309 CD patients were identified for the study, of which 534 [23.1%] were treated with ADA, 85 [3.7%] with CZP, 847 [36.7%] with IFX, 72 [3.1%] with IFX-BS, 394 [17.1%] with UST, and 486 [21.1%] with VDZ.

• Patients on UST and VDZ were slightly older and had slightly higher CCI score (Table 1).

• Common comorbidities among CD patients included anemia, hypertension, anxiety, depression, fatigue, obesity, and hyperlipidemia (Table 1).

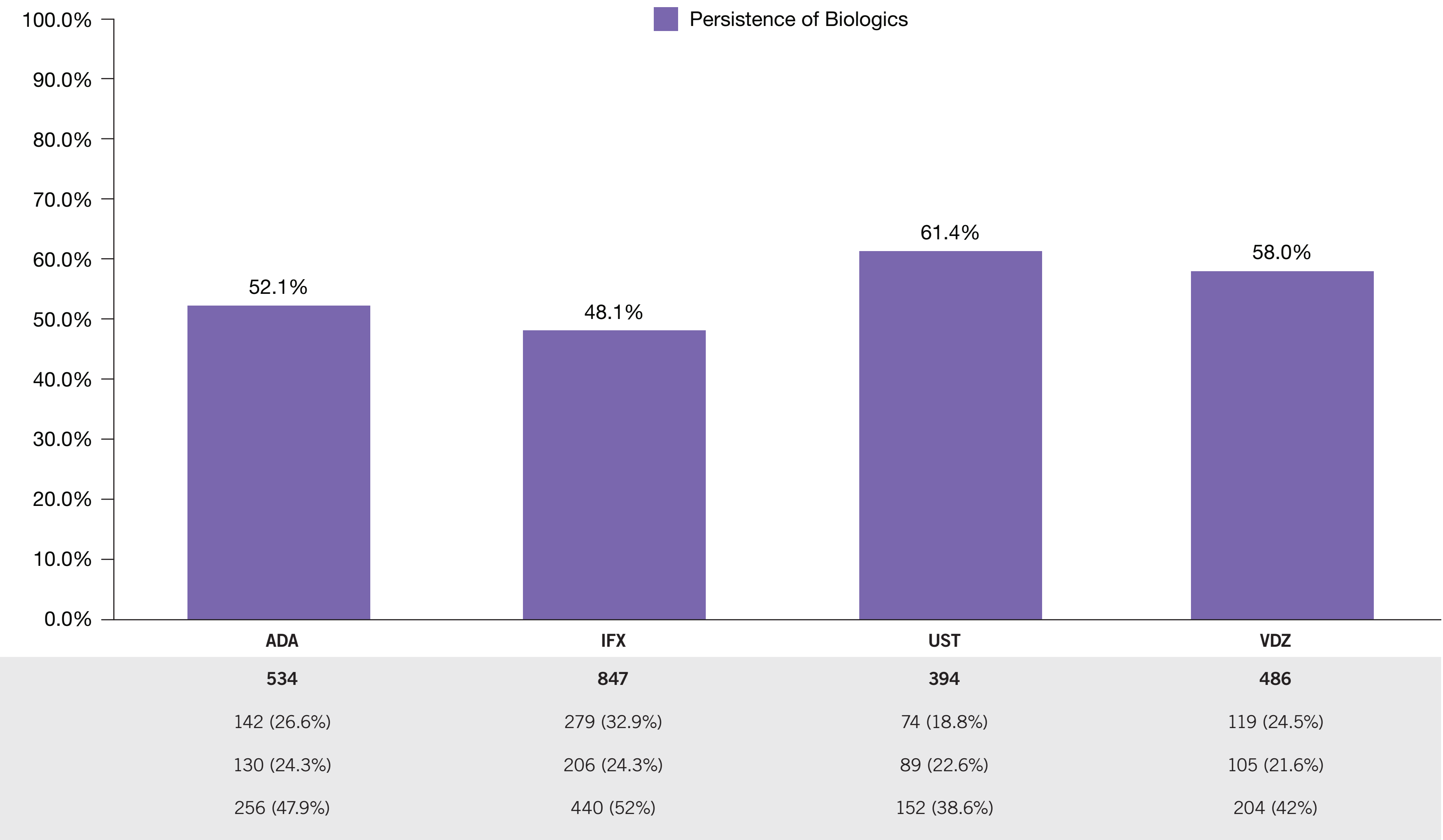
Table 1. Descriptive Baseline Characteristics Among Crohn's Disease Patients

Characteristics	ADA (N=534)	IFX (N=847)	UST (N=394)	VDZ (N=486)
Age in years, mean	43.5	43.7	44.0	45.3
Age group, years				
18-34	24.0%	26.4%	25.6%	20.0%
35-54	52.4%	47.2%	48.5%	52.3%
55-64	23.6%	26.3%	25.9%	27.8%
Sex				
Male	45.7%	42.0%	46.7%	43.6%
Female	54.3%	58.0%	53.3%	56.4%
US geographic region				
Northeast	14.0%	19.0%	17.5%	19.5%
North Central	33.1%	33.1%	37.3%	35.2%
South	37.6%	30.7%	30.5%	31.5%
West	15.2%	17.2%	14.7%	13.8%
Index Year				
2016	14.2%	13.2%	7.9%	13.4%
2017	56.7%	55.1%	53.0%	51.4%
2018	29.0%	31.6%	39.1%	35.2%
Charlson Comorbidity Index Score	0.5	0.5	0.7	0.6
Comorbidities				
Anemia	17.4%	16.9%	30.2%	21.0%
Anxiety	14.0%	12.8%	19.8%	17.5%
Atherosclerosis	0.2%	0.5%	0.5%	0.0%
Celiac Disease	1.1%	2.1%	6.1%	2.9%
Cholelithiasis	0.6%	0.0%	0.3%	0.2%
Chronic pain	5.8%	4.7%	8.9%	8.2%
Depression	14.2%	12.4%	18.3%	17.3%
Diabetes	6.2%	6.3%	7.4%	7.2%
Fatigue	7.7%	8.6%	14.2%	11.3%
Fistula	3.4%	6.7%	6.3%	6.0%
Hyperlipidemia	9.2%	9.3%	9.6%	10.7%
Hypertension	18.9%	15.7%	20.3%	17.5%
Obesity	10.1%	9.4%	11.7%	9.9%
Venous Thromboembolism	0.7%	1.5%	2.8%	1.6%

Due to rounding of the percentages, some of the categorical variables might not add up to 100%.

ADA: adalimumab; IFX: infliximab; UST: ustekinumab; VDZ: vedolizumab.

Figure 2. Persistence among Crohn's Disease Patients using Biologics with 3 Years of Follow-Up



A variable discontinuation gap of > 60 days for ADA and >120 days for IFX, UST, and VDZ between the run-out date of two consecutive biologic claims (gap approximately two times the maintenance dosing interval) was used to define the discontinuation.

ADA: adalimumab; IFX: infliximab; UST: ustekinumab; VDZ: vedolizumab.

Note:

• Over the 3 years of follow-up, the proportion of patients with dose titration was highest for IFX (76.5%) and VDZ (74.5%) and lowest for UST (50.8%) (Table 2).

• The highest cases of dose reduction were observed in patients with UST (16.5%) (Table 2).

• The highest cases of dose escalation were observed in patients with IFX (65.9%) followed by VDZ (53.3%) (Table 2).

• The highest cases of >100% of dose escalation were observed in patients with IFX (41.1%) and lowest in patients with UST (8.5%) (Table 2).

Limitations

- Due to small sample size for the bio-experienced group, the study could not stratify the results across bio-naïve and bio-experienced patients. This stratification is important to understand the inherent variability between patients in each group. Compared to the bio-naïve group, the bio-experienced patients have an inherent bias introduced from prior biologic treatment exposure and potential non-response or failure that leads to initiation of a new biologic therapy. This needs to be explored in future studies using larger data on these patients, especially in the bio-experienced group.
- Analysis of claims data depends on correct diagnosis, procedure, and drug codes. Coding errors can result in misclassification.
- There are limitations about the operational definitions of various variables. 1) Dose escalation is based on change in units between claims from the same beneficiaries and does not look at exact formulation type. However, it avoids the process of making assumptions regarding doses for HCPCS codes 2) Discontinuation and treatment switch could be due to other reasons such as: insurance coverage, adverse events, or other reasons not related to loss of response.
- There would be potential under-capture of biologics use with claims data, which is primarily used for billing and not research purposes. Furthermore, medications received over the counter or provided as samples by the physician are not observed in claims data.

References

1. Floyd DN, Langman S, Silverman HC, Levesque BG. The economic and quality-of-life burden of Crohn's disease in Europe and the United States, 2000 to 2013: A systematic review. *Dig Dis Sci*. 2015;60(2):299-312.
2. Ganz ML, Sugerman R, Wang R, Hansen BB, Håkansson J. The economic and health-related impact of Crohn's disease in the United States: Evidence from a nationally representative survey. *Inflamm Bowel Dis*. 2016;22(10):1033-1041.
3. Bernstein CN, Blanchard JF, Rawsthorne P, Wajda A. Epidemiology of Crohn's disease and ulcerative colitis in a central Canadian province: a population-based study. *Am J Epidemiol*. 1999 May;150(10):916-24. doi: 10.1093/oxfordjournals.aje.009735. PMID: 10342800.
4. Ondrej C. (2020). Persistence, dose titration, and health care resource utilization among Crohn's disease patients treated with ustekinumab: a real-world analysis in the United States. *Adv Ther*. 2020;37(5):1217-43.

Disclosures

Ruchi Zhao, Zhiye Ding, and Sumita Kachroo are employees of Janssen Scientific Affairs, LLC, and Johnson & Johnson. Parul Gupta, Laurence Gozalo, Robert Bruette, Victor M Johnson, and Keshia Maughn are employees of STATinMED, LLC and supported this study as a paid consultant to Janssen Scientific Affairs.

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Table 2. Dose Titration among Crohn's Disease Patients using Biologics with 3 Years of Follow-Up

Biologics	Eligible Cases	%
ADA Sample Size (N=534)	Eligible Cases	68.7*
Dose Reduction	<50% Dose Reduction	5.2
>50% Dose Reduction	>50% Dose Reduction	3.8
Dose Escalation	Dose Escalation	1.4
<50% Dose Escalation	<50% Dose Escalation	47.4
51-100% Dose Escalation	51-100% Dose Escalation	3.0
>100% Dose Escalation	>100% Dose Escalation	32.2
IFX Sample Size (N=847)	Eligible Cases	76.5*
Dose Reduction	<50% Dose Reduction	11.4
>50% Dose Reduction	>50% Dose Reduction	6.6
Dose Escalation	Dose Escalation	4.8
<50% Dose Escalation	<50% Dose Escalation	65.9
51-100% Dose Escalation	51-100% Dose Escalation	8.3
>100% Dose Escalation	>100% Dose Escalation	16.5
UST Sample Size (N=394)	Eligible Cases	50.8*
Dose Reduction	<50% Dose Reduction	16.5
>50% Dose Reduction	>50% Dose Reduction	0.5
Dose Escalation	Dose Escalation	35.5
<50% Dose Escalation	<50% Dose Escalation	13.0
51-100% Dose Escalation	51-100% Dose Escalation	14.0
>100% Dose Escalation	>100% Dose Escalation	8.5
VDZ Sample Size (N=486)	Eligible Cases	74.5*
Dose Reduction	<50% Dose Reduction	14.6
>50% Dose Reduction	>50% Dose Reduction	8.8
Dose Escalation	Dose Escalation	5.8
<50% Dose Escalation	<50% Dose Escalation	53.3
51-100% Dose Escalation	51-100% Dose Escalation	3.9
>100% Dose Escalation	>100% Dose Escalation	23.2
ADA: adalimumab; IFX: infliximab; UST: ustekinumab; VDZ: vedolizumab.		

*Provided % are of eligible cases with respect to biologic sample size. Other provided % are with respect to eligible cases.

- Over the 3 years of follow-up, the proportion of patients with dose