Sex Based Disparities in Cost Related Medication Nonadherence in Older Adult Patients with Type 2 Diabetes from the All of Us Database

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BACKGROUND

CRITICAL BURDEN

Type 2 Diabetes (T2DM) affects nearly 30% of all adults over the age of 65 in the U.S. -- effective medication adherence is crucial to prevent serious complications.

COST BARRIERS

High medication costs hinder adherence, leading to poorer outcomes; understanding this barrier is essential for improving outcomes.

SEX DISPARITIES

Investigating sex-based disparities in cost-related medication nonadherence (CRN) can inform targeted interventions

To examine the association of sex and CRN in patients over the age of 65 who are prescribed medication for the treatment of T2DM

METHODS

- Data from the All of Us Research Program, a diverse biomedical database funded by the NIH
- A total of **6,450 participants aged 65 years or older** and diagnosed with **T2DM** with a **prescription for medication** used to treat T2DM within 90 days of diagnosis
- Independent variable was sex at birth
 - Male vs Female
- Outcome variables were responses to seven questions on CRN
 - 1. Was there any time when you needed prescription medicines but did not get it because you could not afford it?
- 2. Did you skip medication doses to save money?
- 3. Did you take less medicine to save money?
- 4. Did you delay filling a prescription to save money?
- 5. Did you ask your doctor for a lower cost medication to save money?
- 6. Did you buy prescription drugs from another country to save money?
- 7. Did you use alternative therapies to save money?
- Covariates:
- Sociodemographic factors (Age at time of survey, Income, Race/Ethnicity, Education, Insurance Status)
- **BMI** (encoded as categorical variable)
- Comorbidities (Charlson Comorbidity Index, Alcohol Use Disorder, Substance Use Disorder, Cigarette Use History)
- Medication Use History (Biguanide, DPP-4 Inhibitor, GLP-1 Agonist, SGLT-2 Inhibitor, Sulfonylurea, Thiazolidinediones, Alpha-glucosidase inhibitors, Bile Acid Sequestrants, Dopamine-2 Agonists, Meglitinides, Insulins and Analogues)

Statistical Analysis

- Chi-square and Analysis of Variance (ANOVA) were used to test the baseline characteristics of the sample grouped by sex
- Multivariable Logistic Regression Models were used to estimate the association between sex and CRN, adjusting for all covariates. All analyses were performed on the All of Us Researcher Workbench Platform using *sklearn* in Python v3.10.12

RESULTS

Baseline Descriptive Characteristics of the

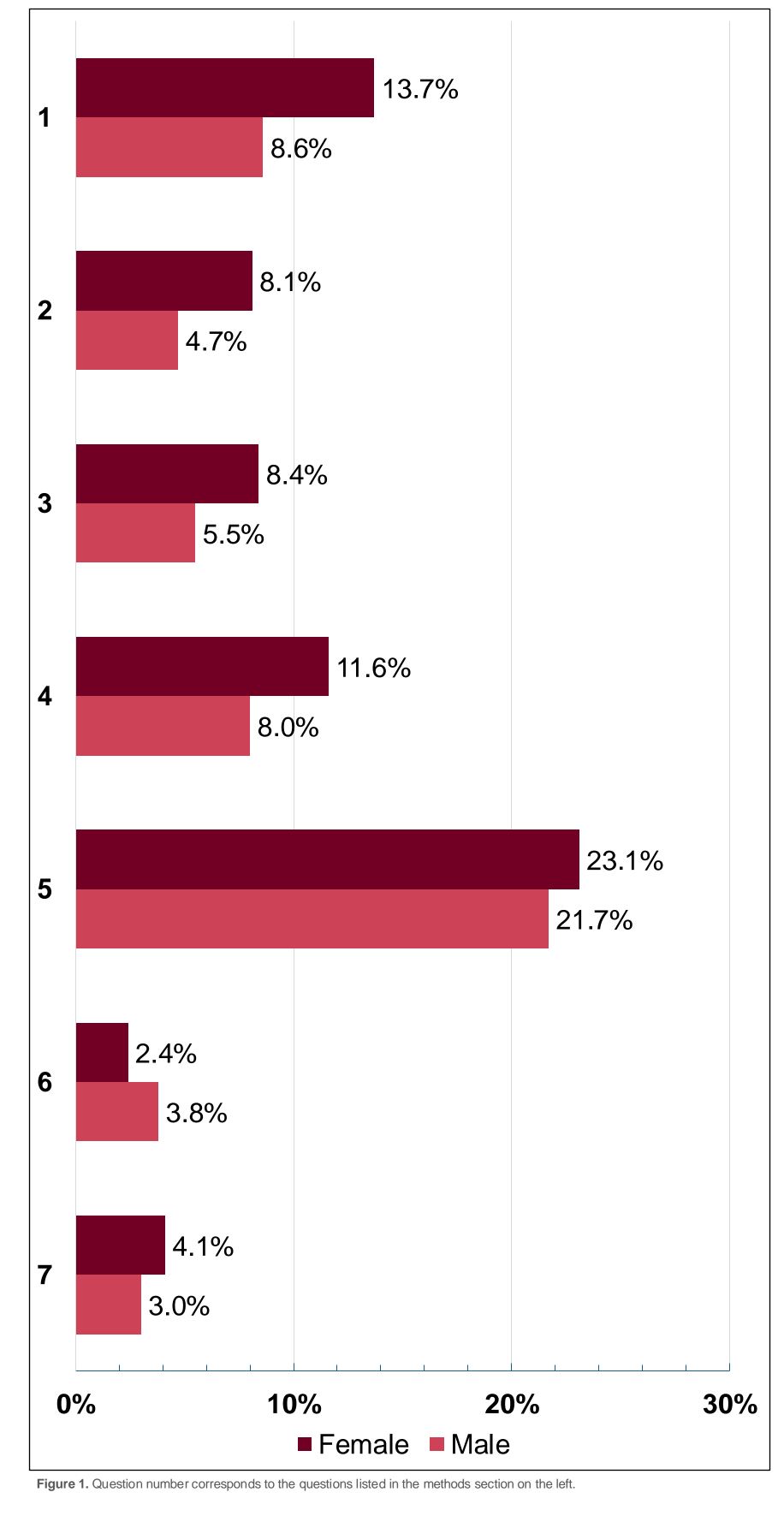
Sample by Sex (N=6450).

Table 1

Table 2

Characteristic Female (n=3375) p-value Age, Median (IQR) 72.00 (68.00,76.00) < 0.0001 74.00 (70.00,78.00) Race and Ethnicity, No. (%) Non-Hispanic White 1950 (60.6) 2382 (77.5) Hispanic/Latino 239 (7.8) 449 (13.9) Non-Hispanic Black 252 (8.2) 651 (20.2) Non-Hispanic Asian < 0.0001 47 (1.5) 34 (1.1) Non-Hispanic Multiple 36 (1.2) 35 (1.1) Other Ethnicity 33 (1.0) 34 (1.1) Non-Hispanic MENA Non-Hispanic NHPI Income, No. (%) <25k 420 (13.7) 826 (25.7) 25k - 50k 664 (20.6) 538 (17.5) < 0.0001 50k - 100k 702 (21.8) 885 (28.8) 100k - 150k 479 (15.6) 275 (8.5) >150k 174 (5.4) 340 (11.1) Education, No. (%) Less than High School or Equiv. 316 (9.8) 152 (4.9) High School or GED 362 (11.8) 535 (16.6) < 0.0001 Some College/Vocational 822 (26.7) 1016 (31.6) College or Advanced Degree 1704 (55.4) 1295 (40.2) Insurance, No. (%) **Employer or Union** 385 (12.5) 436 (13.5) Medicaid 314 (10.2) 359 (11.2) Medicare 759 (23.6) 754 (24.5) Mixed 57 (1.8) 166 (5.4) Purchased 85 (2.6) 57 (1.9) < 0.0001 83 (2.6) Other Insurance 161 (5.2) No Insurance 64 (2.1) 71 (2.2) Employer/Medicare 359 (11.2) 314 (10.2) Medicare/Medicaid 109 (3.5) 278 (8.6) Medicare/Other 53 (1.6) 287 (9.3) Medicare/Purchased 250 (8.1) 271 (8.4) BMI , No. (%) <18.5 NA 18.5 - 25 308 (10.0) 299 (9.3) < 0.0001 25 - 30 966 (31.4) 704 (21.9) 1798 (58.5) 2200 (68.3) Comorbidities, No. (%) Cigarette Use 1648.0 (53.6) < 0.0001 1343.0 (41.7) Alcohol Use Disorder < 0.0001 204 (6.6) 72 (2.2) Substance Use Disorder < 0.0001 158 (4.9) 195 (6.3) Charlson Comorbidity Index 6.00 (5.00,8.00) 6.00 (5.00,8.00) < 0.0001 **Table 1.** BMI – Body Mass Index in kg/m², in compliance with *All of Us* Data Dissemination Policy, categories with <20 participants marked as NA

Figure 1 Prevalence of CRN Responses by Sex (N=6450).



DISCUSSION

RESULTS & INTERPRETATION

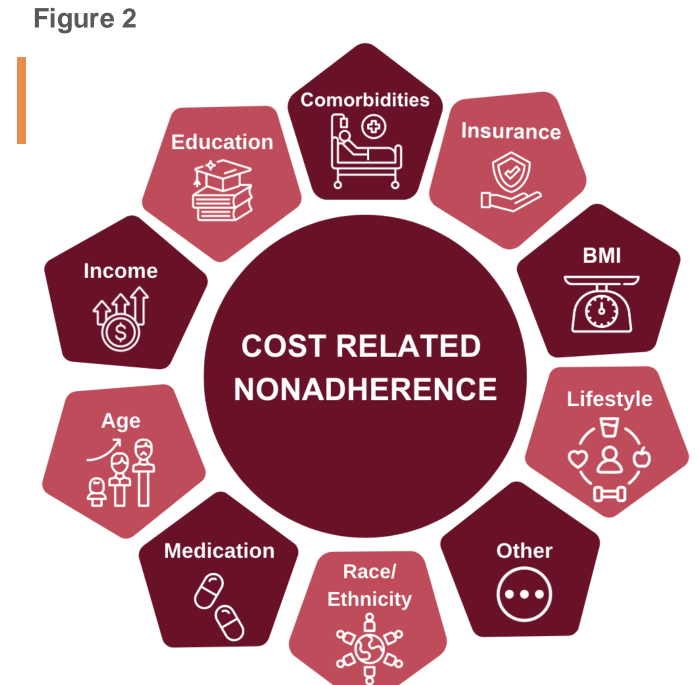
- Older adult **female patients** with diabetes had **higher odds** of the following compared to older adult male patients with diabetes
- Not being able to afford medication.
- Skipping medication to save money.
- Taking less medication to save money.
- Delaying filling medication.
- Older adult **female patients** with diabetes had **lower odds** of the following compared to older adult male patients with diabetes
 - Buying medications from a foreign country to save money.

STUDY STRENGTHS

- Utilizes a large, diverse, representative sample of older adults with diabetes in the United States enhancing the generalizability of the study
- Leverages electronic health record data on comorbidities and medication history combined with responses to survey questions
- Multivariable logistic regression approach allowed for multiple explanatory variable to be explored simultaneously, reducing the effect of confounding factors

STUDY LIMITATIONS

- Limited details on level of medication non-adherence and complications from non-adherence
- Limited data on certain ethnic groups, preventing inclusion of more detailed ethnic categories in regression model
- CRN questions were not specific to diabetes or its medications and may thus represent CRN across a broad range of conditions and medications



INFLUENCING CRN IN OLDER ADULTS.

THERE ARE A VARIETY OF FACTORS

Other factors can include

- Health literacy

- Attitude towards brand medication
- Type of disease - Monitoring of medication therapy
- Awareness and belief in therapy
- Community attitude towards medicine
- Trust in healthcare providers

OR and aOR for Univariable and Multivariable Logistic Regression Model for CRN as a Function of Sex (N=6450)

Medication Barrier	Male (n=3075)	Female (n=3375)	OR (95% CI)	P-value	aOR (95% CI)	p-value
1. Could not afford medication	264 (8.6)	461 (13.7)	1.68 (1.44-1.98)	<0.0001	1.28 (1.07-1.53)	0.0066
2. Skipped medication to save money	144 (4.7)	274 (8.1)	1.8 (1.46-2.21)	<0.0001	1.46 (1.16-1.84)	0.0014
3. Took less medication to save money	168 (5.5)	282 (8.4)	1.58 (1.29-1.92)	<0.0001	1.26 (1.01-1.58)	0.0374
4. Delayed filling medication	245 (8.0)	390 (11.6)	1.51 (1.28-1.78)	<0.0001	1.25 (1.03-1.51)	0.0217
5. Asked for lower cost medication	668 (21.7)	781 (23.1)	1.08 (0.96-1.22)	0.1732	1.04 (0.91-1.19)	0.5768
6. Bought medications from another country	118 (3.8)	80 (2.4)	0.61 (0.46-0.81)	0.0007	0.64 (0.46-0.87)	0.0049
7. Used alternative therapy to save money Table 2. OR – Odds ratio, CI – Confidence interval, aOR – Adjusted Odds Ratio; aOR adjusts	93 (3.0) for sociodemographic factors, B	, ,	1.37 (1.05-1.79)	0.0221	1.06 (0.79-1.43)	0.7056

IMPLICATIONS

PRECISION TAILORED INTERVENTIONS

Figure 2. Variables used in logistic regression model that impact impact cost related nonadherence

Other variables for future research are mentioned in the caption to the right

- Identifying older adults at higher risk of CRN is critical to improve outcomes involving the long-term complications of diabetes.

RESEARCH NEEDS

- Examine the relationship between CRN and outcome measurements such as major adverse cardiac events, emergency room visits, and other complications of diabetes.

FUTURE POLICY

- Future policy should involve addressing the barriers to CRN; interventions can include real-time benefit pricing tools, supplementing brand-name medications with generics, and improving healthcare access and coverage.

ACKNOWLEDGMENT

We acknowledge All of Us participants for their contributions, without whom this research would not have been possible. We also thank the National Institutes of Health's <u>All of Us Research Program</u> for making available the participant data [and/or samples and/or cohort] examined in this study.