A Retrospective Analysis of Creatinine-Based Kidney Function With and Without Sex Assigned at Birth Among Transgender Adults

Sarah K. Fadich, PA-C1, Alin Kalayjian, BS2, Dina N. Greene, PhD3,4, and Lauren R. Cirrincione, PharmD, MPH2

Abstract

Background: Clinicians use sex-based kidney function estimating equations, but the appropriate sex modifier for transgender adults undergoing hormone therapy (HT) is undetermined. Objectives: Compare median estimated creatinine clearance (eCrCL; Cockcroft-Gault) and estimated glomerular filtration rates (eGFRs; Modification of Diet in Renal Disease [MDRD] and Chronic Kidney Disease Epidemiology Study [CKD-EPI]) before and during HT when estimated with and without sex assigned at birth. Methods: Single-system retrospective cohort study of transgender adults (2007-2017) prescribed ≥90 days HT (index date = first order) and measured serum creatinine ≤6 months pre-index date (baseline) and ≤12 months post-index date. We grouped patients based on testosterone or estrogen treatment and compared eCrCL and eGFRs at baseline up to 6-12 months post-index date using equations based on sex assigned at birth (female or male modifier in testosterone or estrogen groups, respectively) or gender identity (male or female modifier in testosterone or estrogen groups, respectively). We used Wilcoxon signed-rank tests (Bonferroni correction) for all comparisons. Results: In total, 29 (median age 26 years, follow-up 259 days) and 41 patients (29 years, 250 days) were prescribed testosterone or estrogen, respectively. In the testosterone group, the maximum eCrCL and eGFR changes based on sex assigned at birth were −14%, P = 0.0181; −18%; P = 0.0009, respectively, and based on gender identity were +5%, P > 0.025 and +11%, P = 0.0094, respectively. In the estrogen group, eCrCL or eGFRs based on sex assigned at birth did not change from baseline but based on gender identity were −17%, P < 0.0001 and −26%, P < 0.0001, respectively. Conclusion and Relevance: Female-based equations may underestimate kidney function in transgender adults undergoing testosterone or estrogen treatment. Prospective cohort studies are needed to confirm the clinical significance of these findings.

Keywords

hormone therapy, transgender adults, estimated creatinine clearance, serum creatinine, estimated glomerular filtration rate

Introduction

Approximately 1 million adults are transgender in the United States,1 and this population is growing worldwide.2 Transgender people, individuals whose gender identity differs from their sex assigned at birth, may take testosterone or estrogen treatment to align secondary sex characteristics with their gender identity.3 Although hormone therapy is associated with improved quality of life among transgender adults,4 its impact on interpreting sex-based clinical estimates, specifically creatinine-based kidney function estimating equations,5 is unclear.

In vivo and population-based cohort data describe conflicting effects of sex hormones on kidney function.6 Although it is unclear whether hormone therapy influences kidney function directly, it typically causes marked physiologic and body composition changes within months after initiation among transgender adults.7 Increased or decreased percent lean muscle mass, and corresponding increases or decreases in serum creatinine concentrations, may alter...
Because estimating equations include a sex-based modifier to account for average body composition differences between sexes, clinicians have recommended using equations based on gender identity, rather than one’s sex assigned at birth, among transgender adults undergoing hormone therapy. Despite this recommendation, no investigators have studied the impact of switching between male and female estimating equations on longitudinal kidney function estimates before and during hormone therapy.

Because pharmacists and prescribing clinicians use creatinine-based kidney function estimating equations to guide dose adjustment of several drugs cleared by the kidneys, we conducted a short-term preliminary investigation to compare estimated creatinine clearance (eCrCL) and estimated glomerular filtration rates (eGFR) before and during hormone therapy among transgender adults. Our primary objective was to compare changes in eCrCL and eGFR based on sex assigned at birth before and during hormone therapy. Our secondary objective was to compare changes from baseline when estimated based on gender identity. We hypothesized that when using equations based on sex assigned at birth during hormone therapy, testosterone or estrogen treatment would be associated with decreased or increased estimates, respectively, and the extent of these changes would be decreased when using equations based on gender identity.

Materials and Methods

Study Design

This was a single-system, multicenter, retrospective cohort study of healthy transgender adults (≥18 years of age) receiving medical care at University of Washington (UW) Medicine in Seattle, Washington, USA. UW Medicine is an integrated health system that includes 4 hospitals and affiliated clinics, plus 12 neighborhood clinics. The University of Washington Institutional Review Board (STUDY00009038) reviewed this project.

Patients

We identified patients between January 1, 2007, and January 31, 2017, with at least one transgender health-related clinical visit based on validated diagnosis codes (International Classification of Diseases Ninth or Tenth Revision, ICD-9 or ICD-10). We extracted demographic, height, weight, clinical laboratory measures (serum creatinine, blood urea nitrogen), concomitant medication orders, and clinical diagnoses within 12 months pre-index and post-index date from the UW Medicine Electronic Data Warehouse, a central repository of electronic medical record data across UW Medicine clinics. Because transgender people may have gender identities outside the binary of man or woman (non-binary), and not all transgender people want or are able to obtain hormone therapy, we grouped patients based on receipt of either testosterone or estrogen treatment to avoid misrepresenting patients’ gender identities within our cohort. Although we approximated gender identity based on hormone therapy for our secondary objective, this approach was meant to align with availability of binary (male or female) kidney function estimating equations and was not intended to exclude or oversimplify identities of nonbinary adults undergoing hormone therapy.

Eligible patients were prescribed testosterone or estrogen treatment for at least 90 days, with the index date set as the first hormone order date, and at least one creatinine measure within 6 months pre-index date (baseline) and within 12 months post-index date. We selected a 90-day threshold for hormone therapy duration to allow our cohort to primarily include patients receiving maintenance doses of hormone therapy. We excluded patients with documented history of chronic kidney disease, dialysis, kidney transplant, HIV infection, or baseline eGFR <60 mL/min/1.73 m² (using the re-expressed 4 variable Modification of Diet in Renal Disease [MDRD] equation and sex assigned at birth) to minimize potential sources of nonhormone-related variability in eCrCL and eGFR. We excluded patients with creatinine measures reported within 90 days of the index date only, as clinicians at our institute typically perform follow-up laboratory monitoring 3 months after hormone therapy initiation in healthy patients. We excluded patients with >12 creatinine measures reported within any 6-month period 1 year pre-index or post-index date as a surrogate marker of kidney instability.

Primary Outcome Measures

Our primary endpoint was the percent difference in median eCrCL 3-6 months (inclusive) and 6-12 months post-index date compared with baseline using the Cockcroft-Gault (C-G) estimating equation, as 1998 Food and Drug Administration (FDA)-issued industry guidance recommended eCrCL for medication dose adjustment in patients with chronic kidney disease, and most drug product prescribing information includes eCrCL-based dosing guidance for medications requiring kidney dose adjustment. We used ideal body weight, actual body weight, or adjusted body weight at baseline for C-G estimates, and we applied the same body weight estimate used at baseline to all weights reported at follow-up to minimize the influence of potential changes in the body weight estimates between time intervals.

We estimated GFR using the MDRD equation and Chronic Kidney Disease Epidemiology Study (CKD-EPI) equation, as 2020 FDA draft industry guidance recommended using eGFR for medication dosing in
pharmacokinetic studies among patients with kidney impairment. For our primary objective, we estimated baseline eCrCL or eGFR using C-G, MDRD, and CKD-EPI equations associated with sex assigned at birth (ie, male-based equations in the estrogen group; female-based equations in the testosterone group). We described the proportion of patients with ≥0.20 mg/dL absolute change in creatinine concentrations and the proportion of patients with ≥25% absolute change in eCrCL and eGFR, as investigators have recommended this threshold for clinically significant decreases in eGFR in the context of acute kidney injury. For our secondary objective, we re-analyzed C-G, MDRD, and CKD-EPI estimates 3-6 months and 6-12 months post-index date using equations associated with gender identity (ie, male-based equations in the testosterone group; female-based equations in the estrogen group) and compared these with baseline estimates using sex assigned at birth.

Statistical Analysis

We compared the percent change in median eCrCL (C-G) and eGFR (MDRD and CKD-EPI) at 3-6 months post-index date to baseline and 6-12 months post-index date to baseline within each hormone treatment group using Wilcoxon signed-rank tests due to small sample sizes. We summarized continuous variables using median and interquartile range (IQR) or ranges. We used frequencies and percentages to summarize categorical variables. A 2-sided P value <0.025 (Bonferroni correction for multiple testing) was considered statistically significant. We used SAS software version 9.2 (SAS Institute Inc., Cary, North Carolina).

Results

Baseline Patient Characteristics

A total of 989 patients had at least one clinical visit for transgender health-related medical care (Figure 1). We analyzed 70 patients, 29 of whom were prescribed testosterone treatment (Table 1). The median follow-up duration in the testosterone group was 259 days (range: 101-357 days). Most patients were prescribed injectable testosterone weekly (n = 20 cypionate or enanthate, median dose: 60.0 mg [32.5-95.0 mg], 6 patients had unknown dosing). Three patients were prescribed 2-4 mg/24-hour testosterone patches daily. Five patients were prescribed topical testosterone gel preparations (median daily dose: 30.5 mg [21.4-43.8 mg]). One patient was prescribed an unknown testosterone preparation.

Forty-one patients were prescribed estrogen treatment (Table 1). The median follow-up duration in the estrogen

Figure 1. Flow diagram of cohort selection. Abbreviations: eGFR, estimated glomerular filtration rate; ICD-9 or ICD-10, International Classification of Diseases Ninth or Tenth Revision; MDRD, Modification of Diet in Renal Disease; UWM, University of Washington Medicine health system.
Most patients were prescribed oral estradiol tablets (n = 21, median total daily amount: 4 mg [2-4 mg] divided as once, twice, or thrice daily doses). Eight patients were prescribed transdermal estradiol patches (median weekly amount: 150 μg [100-200 μg] delivered via 1-3 patches applied up to twice weekly). Eleven patients were prescribed injectable estradiol (cypionate or valerate, median weekly dose: 10.0 mg [6.6-15.0 mg], 4 patients had unknown dosing). One patient was prescribed topical estrogen gel (dose unknown). Thirty-eight (90.1%) patients were prescribed oral spironolactone (median total daily amount: 200 mg [100-200 mg] divided as once or twice daily doses). Eleven patients were prescribed adjunctive antiandrogenic agents ( dutasteride, finasteride) or progestogens (micronized progesterone or medroxyprogesterone acetate).

Laboratory Values and Body Composition Measures at Baseline and Follow-up

In the testosterone group, median body weight and body mass index were lower at 3-6 months and 6-12 months compared with baseline, but this was not statistically significant (Table 2). Median creatinine concentrations at 3-6 months and 6-12 months were statistically increased compared with baseline (P = 0.0020 and P = 0.0006, respectively). Seven (24.1%) patients had ≥0.20 mg/dL increased creatinine concentration at either 3-6 months or 6-12 months compared with baseline (data not shown).

In the estrogen group, median body weight and body mass index were similar at 3-6 months and 6-12 months compared with baseline (Table 2). Median creatinine concentrations were similar at 3-6 months and 6-12 months compared with baseline. Eight (19.5%) patients had ≥0.20 mg/dL absolute change in creatinine concentrations at either 3-6 months or 6-12 months, 4 of whom had increased values compared with baseline (data not shown).

Percent Changes in eCrCL and eGFR Using Estimating Equations Based on Sex Assigned at Birth

Testosterone group. Using female-based equations at baseline and follow-up in the testosterone group, C-G estimates decreased from baseline, although this was statistically
Table 2. Longitudinal Changes in Laboratory and Physiologic Values During the First Year of Hormone Therapy in Transgender Adults.

| Median parameter | Testosterone group, n = 29a | Baselineb | 3-6 months | 6-12 months | Estrogen group, n = 41a | Baselineb | 3-6 months | 6-12 months |
|------------------|-----------------------------|-----------|------------|-------------|-------------------------|-----------|------------|-------------|
| Weight, kg       | 78.7 (66.1-89.8)            | 68.5 (62.1-87.0), n = 9 | 74.5 (65.9-90.5), n = 22 | 75.3 (67.0-105.6) | 75.3 (64.1-114.3), n = 29 | 72.7 (67.6-103.0), n = 37 |
| Δ from baseline  | –0.7 (−3.2 to 1.2)         | –0.9 (−2.4 to 0.0) | –0.3 (−3.2 to 3.2) | 0.2 (−3.8 to 5.4) |
| P valuec         | 0.5703                     | 0.0286     | 0.9329     | 0.4537       |
| BMI, kg/m²       | 29.4 (25.0-33.7)            | 27.5 (23.5-28.5), n = 9 | 28.6 (25.2-34.0), n = 22 | 24.1 (21.1-32.7) | 25.2 (21.3-34.1), n = 29 | 24.4 (22.2-33.5), n = 37 |
| Δ from baseline  | –0.3 (−1.3 to 0.4)         | –0.3 (−0.9 to 0.0) | –0.1 (−1.0 to 1.1) | 0.1 (−1.3 to 1.8) |
| P valuec         | 0.4961                     | 0.0286     | 0.9412     | 0.4537       |
| Serum creatinine, mg/dL | 0.74 (0.64-0.82) | 0.81 (0.77-0.86), n = 17 | 0.84 (0.74-0.94), n = 20 | 0.83 (0.76-0.94) | 0.87 (0.81-0.92), n = 26 | 0.81 (0.74-0.95), n = 29 |
| Δ from baseline  | 0.05 (0.02 to 0.14)        | 0.12 (0.00 to 0.18) | –0.03 (−0.11 to 0.04) | 0.00 (−0.09 to 0.06) |
| P valuec         | 0.0020                     | 0.0006     | 0.5245     | 0.8911       |
| BUN, mg/dL       | 10.0 (9.0-13.0)             | 9.0 (8.0-11.0), n = 17 | 12.0 (10.0-14.0), n = 20 | 12.0 (10.0-14.0) | 13.5 (10.0-16.0), n = 26 | 12.0 (10.0-14.0), n = 29 |
| Δ from baseline  | –1.0 (−3.0 to 1.0)         | 1.0 (−1.0 to 2.5) | 0.0 (−1.5 to 4.0) | 0.0 (−2.5 to 1.5) |
| P valuec         | 0.1537                     | 0.1161     | 0.3403     | 0.6465       |

Abbreviations: BMI, body mass index; BUN, blood urea nitrogen.

aData presented as median (interquartile range). For all parameters, each participant had at least one value reported at 3-6 months only, 6-12 months only, or at both 3-6 months and 6-12 months.

bBaseline values were measured within 6 months before hormone therapy initiation.

cWithin-group comparisons at 3-6 months versus baseline and at 6-12 months versus baseline using Wilcoxon signed-rank test (with Bonferroni correction for multiple comparisons).
significant at 6-12 months only (Figure 2a, gray plots): baseline, 120 (97-143) mL/min; 3-6 months (n = 5), 94 (93-114) mL/min, −3%, $P = 0.3125$; 6-12 months (n = 15), 93 (83-119) mL/min, −14%, $P = 0.0181$. Median MDRD and CKD-EPI estimates statistically decreased at both 3-6 months (n = 17) and 6-12 months (n = 20) compared with baseline (Figure 2a, gray plots): MDRD at baseline, 99 (83-120) mL/min/1.73 m$^2$; 3-6 months, 89 (81-96) mL/min/1.73 m$^2$, −7%, $P = 0.0013$; 6-12 months, 80 (71-100) mL/min/1.73 m$^2$, −18%, $P = 0.0006$; CKD-EPI at baseline, 116 (97-124) mL/min/1.73 m$^2$; 3-6 months, 105 (94-112) mL/min/1.73 m$^2$, −7%, $P = 0.0046$; 6-12 months, 94 (83-113) mL/min/1.73 m$^2$, −9%, $P = 0.0009$. Seven (24.1%) patients had ≥25% decrease in C-G, MDRD, or CKD-EPI estimates at 3-6 months or 6-12 months (data not shown).

**Estrogen group.** Using male-based equations at baseline and follow-up in the estrogen group, C-G estimates decreased at 3-6 months (n = 21) and increased at 6-12 months (n = 28), although neither was statistically significant (Figure 2b, white plots): C-G at baseline, 129 (112-153) mL/min; 125 (116-144) mL/min, 5%, $P = 0.2842$; 145 (105-163) mL/min, 0%, $P = 0.6567$. Changes in median MDRD and CKD-EPI estimates were within +5% at 3-6 months (n = 26) and 6-12 months (n = 29) compared with baseline (Figure 2b, white plots): MDRD at baseline, 111 (94-125) mL/min/1.73 m$^2$; 3-6 months, 106 (94-122) mL/min/1.73 m$^2$, −1%, $P = 0.3777$; 6-12 months, 100 (91-111) mL/min/1.73 m$^2$, −4%, $P = 0.2447$; CKD-EPI at baseline, 118 (101-134) mL/min/1.73 m$^2$; 3-6 months, 108 (97-120) mL/min/1.73 m$^2$, −5%, $P = 0.3093$; 6-12 months, 103 (92-116) mL/min/1.73 m$^2$, −4%, $P = 0.3777$. Seven (24.1%) patients had ≥25% decrease in C-G, MDRD, or CKD-EPI estimates at 3-6 months or 6-12 months (data not shown).
shown). From baseline (n

increased serum creatinine concentrations are associated with testosterone treatment. This finding suggests on average, neither testosterone nor estrogen treatment increased eCrCL or eGFR changes within 25% of baseline estimates.

**Discussion**

To our knowledge, this preliminary study is the first to describe eCrCL or eGFR with or without the estimating equation based on sex assigned at birth in a cohort of transgender adults. Using C-G, MDRD, or CKD-EPI estimating equations based on sex assigned at birth, we observed decreased eCrCL and eGFR estimates 3-6 months and 6-12 months during testosterone treatment (vs baseline) and no change in eCrCL or eGFR estimates 3-6 months and 6-12 months during estrogen treatment. Furthermore, most patients in each treatment group had creatinine concentration changes within 0.2 mg/dL and eCrCL or eGFR changes within 25% of baseline estimates. This finding suggests on average, neither testosterone nor estrogen treatment were likely associated with clinically significant changes in estimated kidney function and aligns with serum creatinine measures and kidney function estimates reported by several retrospective and prospective cohort studies among transgender adults. Our study adds to the body of evidence suggesting that increased serum creatinine concentrations are associated with testosterone treatment.

Assuming our observations in the testosterone group were likely related to nonkidney-based changes in creatinine concentrations, we explored estimating equations based on gender identity during hormone therapy, rather than sex assigned at birth, in an attempt to minimize the percent difference in kidney function estimates between baseline and follow-up. Using male-based C-G, MDRD, and CKD-EPI equations in the testosterone group, we observed percent changes between +13% and +26% at 3-6 months (compared with −9% to −3% based on sex assigned at birth, ie, female-based equations) and between 0% and +5% at 6-12 months (compared with −14% to −9% based on sex assigned at birth). This finding supports the recommendation by Webb et al suggesting clinicians use male-based equations after at least 6 months of testosterone therapy.

We also used female-based estimating equations at follow-up in the estrogen group and observed percent changes between −23% and −12% at 3-6 months (vs 1%-5% using sex assigned at birth, ie, male-based equations) and between −26% and −15% at 6-12 months (compared with 0% using sex assigned at birth). Thus, female-based estimating equations may underestimate eCrCL and eGFR in transgender adults within the first year of estrogen treatment. This finding is unsurprising, as MDRD and CKD-EPI equations include a sex-specific covariate to adjust for average differences in serum creatinine between sexes, and the C-G equation includes an arbitrary female modifier to decrease equation estimates by 15% compared with male-based equations.

Because female-based estimating equations cause systematically lower kidney function estimates than male-based equations, and because we observed no change in serum creatinine within our estrogen treatment group, the female-based estimating equations likely underestimated kidney function during estrogen treatment. If kidney function is underestimated, then certain medications with kidney-based dose adjustments may be underdosed, although reports of subtherapeutic dosing among transgender adults are lacking. Thus, female-based equations may be inappropriate for medication dosing during the first year of testosterone or estrogen treatment.

Mechanisms underpinning potential hormone therapy–mediated changes in estimated kidney function are unclear. Testosterone treatment increased proinflammatory and profibrotic signaling in animal models of kidney obstruction; however, in humans, low total testosterone concentrations were associated with increased cardiovascular risk among cisgender men with chronic kidney disease. In animal models, estradiol treatment was protective against age-related mechanisms of kidney decline, including diminished nitric oxide synthesis and fibrosis formation. Conversely, large community-based studies among cisgender women observed an association between estrogen-containing medication exposure and increased odds of microalbuminuria, a marker of early kidney disease, and decreased eGFRs. Based on
these data, testosterone or estrogen treatment may be associated with altered kidney function. Prospective, interventional studies using standardized hormone regimens are needed to determine the effect of hormone therapy on kidney function at doses used for gender-affirming medical care.3

Nonkidney-related changes in body composition likely influenced the observed decreases in eCrCL and eGFR in the testosterone group, although this potential influence was slight in the estrogen group. Serum creatinine, an endogenous biomarker of glomerular filtration, is produced from muscle metabolism and it is either increased or decreased, respectively, by increased or decreased muscle mass.29 Thus, it is unclear whether creatinine-based estimating equations reliably estimate kidney function in transgender adults undergoing hormone therapy.5 Because serum creatinine must be at steady state to estimate kidney function accurately,29 prospective reference interval studies using exogenous filtration markers (eg, inulin or iohexol) are needed to determine appropriate kidney function reference intervals for transgender adults undergoing hormone therapy. Furthermore, people who identify as nonbinary may take lower doses of either testosterone or estrogen treatment, with unclear implications on body composition changes and kidney function estimating equations. Future prospective studies, particularly reference interval studies, should include nonbinary patients and develop appropriate reference intervals for these patient populations.

This study has several strengths. Our longitudinal design allowed each patient to serve as their own control before and during hormone therapy. We used validated ICD-9 and ICD-10 codes to identify adults who received transgender health-related services within the UW Medicine system.18,19 However, this small hypothesis-generating study had certain limitations. We used a convenience sample based on electronic medical record data, which may have introduced selection bias that limited generalizability of our findings to transgender adults receiving medical care within the UW Medicine system only. Although we used validated ICD-9 and ICD-10 codes to identify transgender patients, some clinicians document transgender-related medical visits using nonspecific ICD diagnostic codes that are not validated for identifying transgender patients in the electronic medical record (eg, E34.9 endocrine disorder, unspecified).30 We did not have access to clinic notes and were unable to comment on our cohort’s self-reported gender identity. Our analyzable sample was small, limiting our ability to control for potential confounding factors (eg, concomitant medication orders).

We did not have access to measured creatinine clearance (eg, 24-hour urinary creatinine clearance), measured GFRs using exogenous filtration markers (eg, inulin or iohexol), or body composition measures (eg, lean muscle mass), limiting our ability to compare measured (rather than estimated) changes in kidney function during hormone therapy. Because clinicians use creatinine-based kidney function estimates for chronic kidney disease staging, transplant eligibility determination, and dosing of certain medications cleared by the kidney,13,15 future studies should examine the association between altered estimates and these outcomes.

Conclusion and Relevance

Whether sex-based or gender identity-based kidney function estimating equations are accurate for transgender adults remains unclear, although female-based estimating equations may underestimate eCrCL or eGFR among transgender adults undergoing either testosterone or estrogen treatment. Larger prospective studies with measured GFRs are needed to determine the impact of hormone therapy on kidney function in transgender adults during short-term (<1 year) and long-term (≥1 year) treatment. Blanket application female-based estimating equations may underestimate kidney function in transgender adults undergoing hormone therapy.

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ORCID iD

Lauren R. Cirrincione https://orcid.org/0000-0002-9115-1060

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