Performance versus Risk Factor-Based Approaches to Coronary Artery Disease Screening in Waitlisted Kidney Transplant Candidates

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Coronary artery disease · Screening · Kidney transplantation · Six-minute walk test

Abstract

Introduction: Current screening algorithms for coronary artery disease (CAD) before kidney transplantation result in many tests but few interventions. Objective: The aim of this study was to study the utility of 6-minute walk test (6MWT), an office-based test of cardiorespiratory fitness, for risk stratification in this setting. Methods: We enrolled 360 patients who are near the top of the kidney transplant waitlist at our institution. All patients underwent CAD evaluation irrespective of 6MWT results. We examined the association between 6MWT and time to CAD-related events (defined as cardiac death, revascularization, nonfatal myocardial infarction, and removal from the waitlist for CAD), treating noncardiac death and waitlist removal for non-CAD reasons as competing events. Results: The 6MWT-based approach designated approximately 45% of patients as “low risk,” whereas a risk factor- or symptom-based approach designated 14 and 81% of patients as “low risk,” respectively. The 6MWT-based approach was not significantly associated with CAD-related events within 1 year (subproportional hazard ratio [sHR] 1.00 [0.90–1.11] per 50 m) but was significantly associated with competing events (sHR 0.70 [0.66–0.75] per 50 m). In a companion analysis, removing waitlist status from consideration, 6MWT result was associated with the development of CAD-related events (sHR 0.92 [0.84–1.00] per 50 m). Conclusions: The 6MWT designates fewer patients as high risk and in need of further testing (compared to risk factor-based approaches), but its utility as a pure CAD risk stratification tool is modulated by the background waitlist removal rate. CAD screening before kidney transplant should be tailored according to a patient’s actual chance of receiving a transplant.

Introduction

Kidney transplant, the preferred treatment for advanced chronic kidney disease (CKD) and end-stage kidney disease (ESKD), is an elective surgical procedure. Screening for coronary artery disease (CAD) prior to transplant has been and remains a key part of the pre-transplant evaluation [1]. The optimal, evidence-based approach to screening asymptomatic patients, however, is far from clear.
Recent advances have questioned the utility of CAD screening, especially as revascularization of stable, asymptomatic CAD lesions has not been shown to improve outcomes in multiple patient populations [2, 3]. However, in practice, pretransplant screening remains ubiquitous [4]. A rational first step may be to reduce the number of patients screened, starting with risk stratification that appropriately focuses testing on a subset of high-risk patients while shielding low-risk patients from unnecessary testing. Current guidelines [5, 6] risk-stratify patients based on traditional CAD risk factors. However, these risk factors “rule in” most kidney transplant candidates for testing. By definition, 100% of kidney transplant candidates have advanced CKD, a CAD risk factor. Furthermore, most patients with CKD have additional CAD risk factors. In a multicenter cohort of kidney transplant recipients, diabetes mellitus, dyslipidemia, smoking, and CAD or CAD-equivalents were present at 26, 36, 35, and 14%, respectively [7]. In another cohort, 92% of transplant candidates qualified for cardiac testing under risk factor-based guidelines [8]. Despite this high “rule in” rate, the intervention rate was only 9.5% in a study of Medicare beneficiaries with ESKD [9].

Risk stratification might otherwise be accomplished by physical performance testing. Ability to achieve ≥4 metabolic equivalent of tasks (METs) is an early branch point in the screening algorithm for elective, non-cardiac surgeries in the 2014 ACC/AHA guideline [10]. Two studies to date have considered physical performance in risk stratification prior to kidney transplantation. In one, self-reported physical performance was not sensitive for need for revascularization in kidney transplant candidates [8]. Such a finding may have been due to the absence of a standard questionnaire, recall bias, accommodation to impaired performance, or a tendency to overstate health. A second study examined the utility of aerobic capacity, or peak oxygen intake (pVO₂), obtained through formal cardiopulmonary exercise testing (CPET) [11]. At a threshold of 17 mL/kg/min, sensitivity of pVO₂ in predicting cardiovascular events and all-cause mortality was 79–94% in kidney transplant candidates [11]. However, measuring pVO₂ still requires specialized equipment and personnel, and abnormal results need to be “followed up” with formal CAD testing; the addition of CPET could therefore increase, rather than decrease, the overall test burden and health-care utilization of kidney transplant candidates.

The 6-minute walk test (6MWT) is a promising office-based tool that approximates pVO₂ and could serve as a risk-stratification tool suitable for screening. The prognostic value of 6MWT has been demonstrated in numerous cardiopulmonary and vascular conditions, including heart failure, chronic obstructive pulmonary disease, pulmonary arterial hypertension, and peripheral arterial disease [12, 13]. The 6MWT correlates directly with pVO₂ and METs (r = 0.55–0.70) [12], including in patients with CKD [14]. Compared to self-report, the 6MWT is less prone to bias. Compared to pVO₂, 6MWT results are obtainable in routine clinical settings.

At our institution, we incorporated the 6MWT into the routine evaluation for medically complex deceased donor kidney transplant candidates nearing the top of our waitlist of 1,800–1,900 patients [15]. Organ shortage is acute in our area, and wait-time to a deceased donor kidney may exceed 10 years [16]. We therefore focused CAD assessment on patients who have survived to the top of waitlist. Our transplant team used 6MWT results to inform counseling on frailty, transplant candidacy, perioperative risk, and need for prehabilitation. Our team did not use 6MWT results to inform need for CAD testing or retesting, but adhered to our center-specific protocol outlined below.

We examined outcomes and CAD test results in our patients who underwent the 6MWT. Our main questions (and objectives) were as follows:

1. Does the 6MWT-based approach predict CAD-related events, and how does it perform compared to history- or risk factor-based approaches?
2. Does the 6MWT-based approach correlate with CAD test results, and how does it perform compared to history- or risk factor-based approaches?
3. How does the 6MWT-based approach perform compared to noninvasive CAD test results in predicting CAD-related events within 1 year of testing?

**Methods**

**Cohort Assembly**

We included all patients evaluated through our program’s Transplant Readiness Assessment Clinic from May 2017, when we implemented the 6MWT, through April 2019, to allow at least 1 year of follow-up. The operational details of our deceased donor transplant candidate waitlist management strategy have been published previously [15, 16] (see online suppl. S1; for all online suppl. material, see www.karger.com/doi/10.1159/000516158 for details).

**CAD Testing**

CAD testing schedule adhered to our pre-established clinical algorithm (online suppl. S2). Specifically, our protocol required coronary angiography as the first-line test in patients with diabetic kidney disease and high-risk features [17], and revascularization of any flow-limiting stenoses was detected.
Six-Minute Walk Testing

Experienced personnel from an affiliated Veterans Affairs Medical Center’s Exercise Training Unit trained 3 transplant nurse coordinators to perform the 6MWT, who conducted them at initial and follow-up visits. The training personnel performed within-subject measurements periodically against the coordinators’ measurements for quality assurance. Coordinators instructed patients to walk as fast as they could, at a safe pace, with their usual walking aid/prosthesis and/or with family assistance in the case of sensory impairment, in a quiet hallway between 2 marked objects delineating a distance of 100 feet over 6 min. Breaks and early stops were permitted and included in the 6 min. Nonambulatory patients received a 6MWT result of 0, and the reason for nonambulation was recorded. When a patient had multiple visits and multiple 6MWT results, we included the first result. Symptoms during testing were recorded; they were categorized as cardiac (chest pain, dyspnea, unexplained upper gastrointestinal symptoms with exertion, and fatigue [early stops due to lack of stamina]) or noncardiac (noncardiac pain and neuromuscular weakness). The tests were performed on the same day as the patient’s clinic appointment and generally scheduled on nondialysis days, although we did not collect that information.

Risk Stratification Approaches

We analyzed 6MWT results in multiple ways: (1) as a continuous variable; (2) in a binary fashion using 400 m as a cut-off, based on prior studies demonstrating that 400 m corresponds to 4 METs [18–20]; and (3) designating 6MWT result ≥400 m and no cardiac symptoms provoked by 6MWT as low risk.

We defined self-reported cardiac symptoms as chest pain, dyspnea, or unexplained upper gastrointestinal symptom, elicited through history and review of systems. Of the risk factor guidelines, we chose the 2012 American Heart Association/American College of Cardiology Foundation Scientific Statement endorsed by the American Society of Transplantation (hereafter referred to as “AHA/ACCF criteria”) [6], reported to be the most commonly used in a recent survey of kidney transplant professionals in the United States [4].

Outcomes

Our primary outcome was a CAD-related event, a composite of cardiac death, nonfatal myocardial infarction (defined as type 1 or 2 per the universal definition) [21], coronary revascularization (urgent and nonurgent), or removal from waitlist for advanced CAD. We adjusted the analyses for competing events, including death from a noncardiac cause and waitlist removal for non-CAD-related reasons.

Our secondary outcome was CAD test results, performed within 6 months of the 6MWT and without intervening revascularization or a CAD-related event. We defined a positive myocardial perfusion scan as moderate or severe reversible ischemia, and a positive stress echocardiogram as a stress-induced wall motion abnormality. We defined a positive coronary angiogram as ≥70% stenosis or fractional flow reserve ≤0.80 in a major epicardial vessel.

Blinding

In the choice of whether or not to perform CAD testing and which test to perform, we did not consider the 6MWT result, but rather adhered to our center protocol (online suppl. S2). The consulting cardiologists who performed and read the CAD tests were all blinded to the 6MWT results. A nephrologist (DJW) and a cardiologist (HA) performed independent adjudications of primary and secondary outcomes and resolved any discrepancy via a consensus format. Both were blinded to the exposure (6MWT result).

The transplant team was not blinded to the 6MWT result. The 6MWT formed a part of the patient’s global physical function assessment (in conjunction with other tests including the sit-to-stand tests and SF36 questionnaire) and used it to inform counseling on frailty, transplant candidacy, perioperative risk, and need for prehabilitation. However, we used no threshold of 6MWT to remove a patient from the waitlist or bar a patient from transplant. Further details are in our prior works [15].

Analysis

For objectives #1 and #3, wherein we examined the association between a 6MWT-based approach and CAD-related events, we applied the Fine and Grey subdistribution hazards models to examine time to a CAD-related event as the primary outcome, accounting for the competing events of noncardiac death and waitlist removal for non-CAD-related reasons. Using Kaplan-Meier product limit estimates, we built a time-dependent receiver operating characteristic curve (ROC(t)) and calculated the 1-year area-under-the-curve (AUC(t)) for the predictor in question. The ROC(t) is a statistic generalizing the traditional ROC, developed for binary outcomes, to time-to-event analyses, by allowing the AUC to vary over time. We chose to compute the AUC(t) at 1 year, as most of our patients were within 1 year of a kidney transplant at the time of their 6MWT. We tested for interaction between the 6MWT and prespecified subgroups based on age and presence or absence of diabetes mellitus, CAD, prior CAD testing, and noncardiac factors affecting the 6MWT (including lung disease, severe neuropathy or other neurological deficit, lower extremity amputations, active orthopedic issues, assistive walking device use, or noncardiac pain).

For objective #2, we used logistic regression models to examine the association between the 6MWT and CAD test results. We tested for interaction between the 6MWT and the modality of CAD test (invasive vs. noninvasive).

Companion Analysis

In light of the high competing event rate, we performed a companion analysis in which we examined time to CAD-related event removing waitlist status change as a competing event in the subset of patients on whom we were able to obtain follow-up information. Here, we defined a CAD-related event as cardiac death, nonfatal myocardial infarction, or coronary revascularization. The only competing event was noncardiac death.

We used SAS Enterprise version 7.3 (Cary, NC, USA).

Results

Baseline Characteristics

Over the study period, we performed the 6MWT in 360 patients through TRAC (main cohort and cohort for objective #1). To address the association between the 6MWT and CAD testing (objective #2), we obtained 196 unique 6MWT-cardiac ischemia testing pairs from the
main cohort. To compare the 6MWT-based approach with noninvasive cardiac ischemia testing, we repeated the survival analysis in 89 patients who had noninvasive testing within 6 months of the 6MWT testing and without intervening CAD-related event or revascularization (objective #3). Figure 1 depicts details of cohort assembly.

Table 1 outlines the baseline characteristics of the main cohort. Patients with a 6MWT result ≥400 m, compared to those with a 6MWT result <400 m, were younger, more likely to be male, less likely to have had a prior cardiac evaluation, and less likely to have atherosclerotic disease, diabetes mellitus, or noncardiac factors affecting the 6MWT. There was no appreciable difference in self-reported cardiac symptoms, but patients with 6MWT results ≥400 m were significantly less likely to have cardiac symptoms during the 6MWT or meet the AHA/ACCF criteria for cardiac screening. Online suppl. S3 outlines the baseline characteristics of cohort #2 and #3 – the overall covariate distribution was similar to that of the main cohort, but due to the smaller sample sizes some of the differences were no longer statistically significant.

Patients with a 6MWT result ≥400 m had a significantly longer follow-up time in our cohort, a lower likelihood of removal from the waitlist, and a higher likelihood of receiving a kidney transplant during follow-up, but the CAD test rate was comparable in both groups (Table 2).

Both a 6MWT result ≥400 m and a 6MWT result ≥400 m plus no symptom during the 6MWT designated 45 and 44% of patients, respectively, as “low risk.” In contrast, only 14% of patients met the AHA/ACCF criteria of “low risk.” Only 19% patients had cardiac symptoms by history, thus 81% patients were designated “low risk” by the self-reported symptom approach.

**Association with a CAD-Related Event**

Over a median follow-up of 499 (IQR 141–716) days, 50 (14%) CAD-related events and 72 (20%) competing events occurred in the main cohort (N = 360) (Table 2). The 6MWT-based approach was not significantly associated with CAD-related events within 1 year but was significantly associated with competing events (Table 3). Self-reported cardiac symptoms were not significantly associated with either CAD-related events or competing events. Not meeting AHA/ACCF criteria, in contrast, was significantly associated with a lower risk of CAD-related events within 1 year (sHR 0.11 [0.02–0.82]). Multivariate adjustment for age, sex, diabetes mellitus, any atherosclerotic disease, and presence of noncardiac factors affecting the 6MWT did not substantially change the results.

In our subgroup analysis, the association between 6MWT and CAD-related events was not significantly modified by age, sex or the presence or absence of diabetes, preexisting CAD, prior CAD testing, or noncardiac factors affecting the 6MWT (Fig. 2). Furthermore, excluding the 17 patients who were nonambulatory did not alter the results noticeably (online suppl. S3).

**Association with Cardiac Ischemia Tests**

Of the 196 unique 6MWT-cardiac testing pairs, 102 (52%) were invasive coronary angiography, while 94 (48%) were noninvasive tests (52 stress echocardiogram, 38 myocardial perfusion scan, and 3 coronary computed tomography angiography). Forty-five (23%) of the tests...
were deemed “positive,” including 34 (33%) coronary angiograms and 11 (11%) positive noninvasive tests. The 6MWT-based approach did not predict positive CAD test results (Table 4, AUC 0.54–0.55), neither did self-reported cardiac symptoms or AHA/ACCF criteria (Table 4, AUC 0.51–0.52). The results did not differ based on the modality of cardiac ischemia testing (noninvasive vs. invasive, p for interaction >0.05, online suppl. S4).
Comparison of the 6MWT-Based Approach with Noninvasive Cardiac Ischemia Tests

Among a subcohort of 89 patients with a 6MWT and a noninvasive CAD test within 6 months of the 6MWT, 8 (9%) CAD-related events and 12 (13%) competing events occurred over a median follow-up of 577 (IQR 424–750) days. Of the 8 CAD-related events, 2 were type-1 MI, 2 were waitlist removal for CAD, 1 was cardiac death, 2 were nonurgent revascularization, and 1 was urgent revascularization. Of the 12 competing events, 8 were waitlist removal for non-CAD reasons and 4 were noncardiac death. Similar to the main analysis, the 6MWT-based approach was not significantly associated with CAD-related events within 1 year but was significantly associated with competing events (Table 5). A negative CAD test was significantly associated with a lower risk of CAD-related events within 1 year (sHR 0.08 [0.02–0.33]).

Association with CAD-Related Events within 1 Year, Regardless of Kidney Transplant Waitlist Status

In the companion analysis, we extended follow-up for CAD-related events beyond waitlist removal, where pos-
sible. Of the 69 patients who were removed from the waitlist, we were unable to obtain any clinical information after waitlist removal in 23 and excluded them, leaving a sub-cohort of 337 patients (Fig. 1). Over a median follow-up of 575 (IQR 391–750) days, 53 CAD-related events and 12 competing events (noncardiac deaths) occurred. Of the 53 CAD-related events, 9 were type-1 MI, 5 were type-2 MI, 13 were cardiac death, 21 were nonurgent revascularization, and 5 were urgent revascularization. A higher 6MWT result was significantly associated with a lower risk of CAD-related event (sHR 0.92 [0.84–1.00] per every 50 m, AUC 0.56, Table 6). A low-risk category, defined as 6MWT ≥400 m or 6MWT ≥400 plus no symp-

toms during 6MWT, was associated with approximately 50% lower risk of CAD-related event, but AUC remained low (Table 6).

**Discussion**

In this study, we investigated using the 6MWT to risk-stratify patients with high priority on the kidney transplant waitlist to inform novel screening strategies for ischemic heart disease in this challenging patient population. The patients’ treatment team and the research staff were blinded to the 6MWT results while deciding on di-

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**Table 4.** Concordance between risk stratification and CAD testing in a cohort of 196 patients

| Predictor                                      | N (%) meeting low-risk definition | OR      | AUC  |
|------------------------------------------------|-----------------------------------|---------|------|
| 6MWT result (continuous, per 50 m)             | n/a                               | 0.96 (0.81–1.13) | 0.54 |
| 6MWT result ≥400 m                             | 89 (45)                           | 0.68 (0.35–1.35) | 0.55 |
| 6MWT result ≥400 m + no cardiac symptom during 6MWT | 88 (45)                           | 0.70 (0.35–1.39) | 0.54 |
| No cardiac symptom at rest                     | 185 (94)                          | 1.67 (0.36–7.81) | 0.51 |
| Does not meet AHA/ACCF criteria*               | 20 (10)                           | 0.57 (0.16–2.05) | 0.52 |

None of the results were statistically significant, i.e. all of the 95% confidence intervals for odds ratio crossed 1.0. 6MWT, 6-minute walk test; OR, odds ratio; AUC, area under the curve; CAD, coronary artery disease. * AHA/ACCF criteria: from the 2012 American Heart Association/American College of Cardiology Foundation Scientific Statement endorsed by the American Society of Transplantation, high risk is defined as having CAD or CAD-equivalent, diabetes mellitus, or 2 or more risk factors including hypertension, age (>45 for men or >55 for women), cigarette smoking, or left ventricular hypertrophy. Dyslipidemia and family history were 2 additional risk factors that we did not have in our database.
agnostic testing, waitlist removal, or adjudicating study outcomes. Against a high background of competing events, especially waitlist removal for non-CAD reasons, we did not see an association between the 6MWT and CAD-related events. When we omitted kidney transplant waitlist status in the sensitivity analysis, focusing only on CAD-related events and the competing event of noncardiac death, we saw a modest association between the 6MWT and CAD-related events.

The null finding in our primary analysis is surprising, based on preexisting literature on the 6MWT and a prior study demonstrating the strong association between cardiac events and \( p\text{VO}_2 \), of which the 6MWT is a surrogate, in kidney transplant candidates [11]. As the relation between the 6MWT and \( p\text{VO}_2 \) is linear over 10–20 mL/min/kg [22], a range which encompasses Chakkera et al.’s [11] \( p\text{VO}_2 \) threshold for pretransplant CAD screening, we hypothesized that the 6MWT would perform as well as \( p\text{VO}_2 \). A few explanations are possible. In the \( p\text{VO}_2 \) study, the CAD screening protocol incorporated \( p\text{VO}_2 \); in other words, patients who had a good \( p\text{VO}_2 \) did not undergo additional CAD testing. Such a study design necessarily overestimated the sensitivity and specificity of the index test. Our study was performed blinded to the 6MWT result and we were much more likely to recommend upfront coronary angiography in patients with diabetes mellitus (over 50% in our cohort). We therefore detected, and intervened on, asymptomatic disease at a higher rate.

Another explanation of the null finding is the high event rate for competing event and the strong negative association between the 6MWT and the competing event. The 6MWT, or physical performance, is a snapshot of the patient’s overall physical function and health state, of which cardiorespiratory fitness is a part. In this study, as in our previous study [15], the 6MWT was strongly associated with waitlist removal and death. Such an association bi-
ased the association between the 6MWT and CAD-related events toward the null. Patients with a high 6MWT result were more likely to remain on our transplant waitlist, and any CAD-related events they developed would be captured as an endpoint. In contrast, patients with a low 6MWT result had a 32% probability of waitlist removal (Table 2); any CAD-related event they developed after waitlist removal would not be captured. Indeed, the positive results of our companion analysis appear to support the explanation that the higher waitlist removal rate in patients with low 6MWTs results may have confounded the association between 6MWT and CAD-related events.

Based on this observation, we contend that any consideration of cardiac risk stratification in the prevalent kidney transplant candidate population must consider the high background of waitlist removal and death for noncardiac reasons. Advances and successes in cardiology, especially in the medical management of CAD, have translated into a declining significance of CAD as a cause of death in both the pretransplant [23, 24] and posttransplant phases of ESKD [25]. In our study, while the 6MWT appears to be associated with CAD-related events in our companion analysis, the effect is overshadowed in the primary analysis by other medical factors and processes. An implication is that any future CAD screening strategy ought to only be applied among patients who have a reasonable chance of surviving and receiving a kidney transplant in the near future. The role of the 6MWT, given its strong association with all-cause mortality on the kidney transplant waitlist [15], may be then to identify such patients, rather than simply to risk-stratify for CAD. Indeed, the ongoing organ shortage means that the probability of dying on or being removed from the kidney waitlist may exceed the probability of receiving a transplant in many parts of the country, particularly for elderly patients [26, 27]. Our findings thus have far ranging implications for the area of CAD screening and waitlist management for kidney transplant candidates.

How does the 6MWT-based approach compare to other approaches of risk stratification? Self-reported symptoms designated the most (81%) patients at low risk, but was no more predictive of CAD-related events than chance (AUC 0.50). The AHA/ACCF criteria, in contrast, only designated 14% of patients as low risk and not needing testing. Meeting the AHA/ACCF criteria was significantly associated with CAD-related events; we suspect that this is at least partly because our CAD testing criteria overlaps largely with the AHA/ACCF criteria, especially as all patients with diabetic kidney disease (also one of the AHA/ACCF risk factors) underwent coronary angiography as a first-line test in our cohort. Negative CAD testing was associated with a lower rate of CAD events with an AUC of 0.70, suggesting moderate discriminatory ability along the lines of what has been reported (reviewed in Hart et al. [1]). However, such an association is clearly confounded, as patients with negative CAD test results were unlikely to be followed up with more invasive testing. Finally, none of the risk stratification approaches, whether the 6MWT-, symptom-, or risk factor-based, correlated well with CAD test results. Overall, all the risk stratification approaches we evaluated in this study have their shortcomings, as described above.

Our study does not address whether the high elective revascularization rate in our patient cohort is appropriate. Our center’s protocol was based on the a small, single-center trial in Lancet from the 1990s, which showed a 5-fold reduction in cardiac events from coronary revascularization over medical treatment in kidney transplant candidates with type 1 diabetes mellitus [28], which our center subsequently validated [17]. Medical treatment in that study consisted of only aspirin and calcium channel blocker, and its extraordinarily high event rate has not been replicated elsewhere. In a recent web-based survey of US kidney transplant programs, a full 26% reported using the coronary angiogram as first-line in dialysis-dependent kidney transplant candidates [4]. Our center practice, although aggressive, therefore accorded with a sizable minority of US transplant programs. In the recently completed, multicenter ISCHEMIA-CKD trial, where contemporary standards of medical management were applied, there was no obvious advantage to revascularization in patients with stable CAD and CKD, including those with ESKD [2]. What is the benefit of revascularization in patients with obstructive CAD on coronary angiography, but no symptoms and, importantly, good physical performance as evidenced by good 6MWT performance? This is an open question that we hope to address with future studies.

This study has several strengths. We assembled a sample diverse in terms of age, sex, race, ethnicity, and duration of ESKD. We conducted analyses accounting for competing risks, methods essential in populations with multiple comorbidities. There are also multiple weaknesses. Because these patients represent those at the “top of the waitlist,” and because our regions has one of the longest kidney waitlists in the country, these results may not generalize to all kidney transplant candidates. Reflecting this long waitlist and the fact that our cohort is created from a “top-of-the-waitlist” clinic, our rate of waitlist removal was quite high over the study period.
Whether 6MWT-based risk stratification will be similarly limited if applied early in the transplant evaluation process, before the patients have accrued comorbidities that place them at risk for waitlist removal and complicate the interpretation of 6MWT, is an open question. Our center’s protocol favors upfront use of invasive coronary angiography rather than noninvasive testing, in contrast with most other kidney transplant programs [4], and hence our rate of revascularization in asymptomatic patients will likely be higher than at other centers, inflating our CAD-related event rate.

In conclusion, we find that all modes of risk stratification we examined, be they performance-, symptom-, or risk factor-based, are limited in a cohort of prevalent waitlisted kidney transplant candidates with high comorbidity burden and likelihood of CAD event. The 6MWT designates fewer patients as high risk and in need of further testing (compared to risk factor-based approaches), but its utility as a pure CAD risk stratification tool is modulated by the background waitlist removal rate.

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Statement of Ethics

The Stanford Institutional Review Board approved this project (protocol #43639) which we conducted in adherence with the Declaration of Helsinki, and waived the need for individual informed consent.

Conflict of Interest Statement

The authors have no conflicts of interest to disclose.

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Author Contributions

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