Successful revascularization versus medical therapy in diabetic patients with stable right coronary artery chronic total occlusion: a retrospective cohort study

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Abstract

Background: The territory of the right coronary artery (RCA) is smaller than that of the left anterior descending artery. Previous studies have reported conflicting results when considering whether stable RCA-chronic total occlusion (CTO) should be reopened. The coexistence of diabetic and coronary artery diseases represents a severe situation. Therefore, we aimed to determine if stable RCA-CTO in diabetic patients was necessary to be reopened. To our knowledge, no studies have focused on this topic to date.

Methods: We enrolled diabetic patients with RCA-CTO who had clinical presentations of symptomatic stable angina or silent ischemia. RCA-CTO was treated with either successful revascularization (the CTO-SR group) or medical therapy (the CTO-MT group). The primary endpoint was all-cause death. Both Cox regression and propensity score matching analyses were used. Sensitivity analysis was performed based on subgroup populations and relevant baseline variables.

Results: A total of 943 patients were included: 443 (46.98%) patients in the CTO-MT group and 500 (53.02%) patients in the CTO-SR group. After a mid-term follow-up (CTO-SR: 48 months; CTO-MT: 42 months), we found that CTO-SR was superior to CTO-MT in terms of all-cause death (adjusted hazard ratio [HR] [model 1]: 0.429, 95% confidence interval [CI] 0.269–0.682; adjusted HR [model 2]: 0.445, 95% CI 0.278–0.714). The superiority of CTO-SR was consistent for cardiac death, possible/definite cardiac death, repeat revascularization, target vessel revascularization (TVR) and repeat nonfatal myocardial infarction. Subgroup analysis confirmed the mortality benefit of CTO-SR by percutaneous coronary intervention (the successful CTO-PCI subgroup, 309 patients in total). While CTO-SR by coronary artery bypass grafting (the CTO-CABG subgroup, 191 patients in total) offered patients more benefit from repeat revascularization and TVR than that offered by successful CTO-PCI.

Conclusions: For stable RCA-CTO patients with diabetes, successful revascularization offered patients more clinical benefits than medical therapy. CTO-CABG might be a more recommended way to accomplish revascularization.

Trial registration This study was not registered in an open access database

Keywords: Chronic total occlusion, Diabetes, Revascularization, Medical therapy

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Background
The incidence of chronic total occlusion (CTO) is reported to reach 18.2–52% [1–4] in patients undergoing coronary angiography. Right coronary artery (RCA) CTO has been observed in approximately 38–50% [1, 5–7] of the entire CTO population. Although CTO revascularization was observed to have an acceptable success rate and to gain more clinical benefits from successful procedures [8, 9], the RCA-CTO patients who are treated with revascularization, either by percutaneous coronary intervention (PCI) or coronary artery bypass grafting (CABG), are fewer than those patients with left anterior descending branch (LAD) CTO [1, 3, 4, 10]. We speculated that the reason for this discrepancy might partially because the territory of the RCA is smaller than that of the LAD and partially because the conflicting debates on the question of whether the successful reperfusion of RCA-CTO offers patients clinical benefits. Safley and colleagues [5] enrolled the entire CTO population and reported that successful PCI for RCA-CTO did not offer patient survival benefits, which was also confirmed by other experts [11]. However, this claim was doubted by Mitomo and colleagues [6], who observed that successful RCA-CTO PCI was related to a lower cardiac mortality than that related to failed CTO-PCI. Interestingly, we observed that the percentages of diabetic patients in the three studies were different: 21%, 21.6% and 45.2%. These results implied that diabetic RCA-CTO patients might be different. Moreover, Migliorini and colleagues [10] demonstrated that the existence of RCA CTO is a significant risk predictor of death in patients with unprotected left main coronary artery disease who have undergone PCI, which indicated that RCA-CTO should be reopened in patients with severe coronary atherosclerosis in other vessels to prevent adverse events in cases of occlusion in a non-CTO epicardial artery.

Diabetes mellitus is observed in 34–40% CTO patients [12, 13]. The coexistence of diabetes is often associated with a severe coronary situation, which is characterized by endothelial cell dysfunction, microcirculation disorders, high-grade coronary atherosclerosis, a rapid progression of atherosclerosis and poor clinical outcomes [14–18]. Previous studies demonstrated clinical benefits after successful CTO revascularization [12, 19, 20]. However, to our knowledge, no study has focused on the diabetic RCA-CTO population.

Thus, in the present study, we focused on diabetic RCA-CTO patients who had clinical manifestations of stable angina or silent ischemia. We attempted to determine whether successful revascularization offers a clinical benefit when compared with conservative medical therapy.

Methods
Study design and population
The present study was a retrospective cohort study that was performed at Beijing Anzhen Hospital (Beijing, China). From January 2007 to December 2017, a population of 2502 stable patients with one main stem CTO were selected. In this study, we enrolled patients who had only RCA-CTO. The inclusion criteria were as follows: (1) diabetic patients with only one main stem CTO that was diagnosed by angiography and (2) patients with RCA-CTO (American Heart Association segment maps 1, 2 and 3) who had stable clinical presentations of symptomatic stable angina or silent ischemia. We excluded those patients with (1) a prior CABG history; (2) left main coronary artery disease (stenoses ≥50%); (3) a history of myocardial infarctions (MIs) due to a non-CTO artery within 30 days; and (4) patients with cancer or other diseases that may have confounded the endpoints.

Patients were assigned to the medical therapy (CTO-MT) group or to the successful revascularization (CTO-SR) group according to their final treatment strategy of the CTO vessel. Of note, the CTO-SR group enrolled patients who were treated with CTO-CABG or with successful CTO-PCI. The CTO-MT group enrolled patients who were managed by failed CTO-CABG, failed CTO-PCI or initial medical therapy of the CTO vessel. The definition of successful CTO-PCI was obtaining a residual stenosis of <20% and a thrombolysis in myocardial infarction (TIMI) flow ≥2 [21] after implantation of a drug eluting stent to the CTO vessel. Failed CTO-CABG was defined as performing grafts to other arteries but not the RCA territory.

An optimal medical treatment was recommended to both groups of patients who were managed by SR and MT. To reiterate, an antiplatelet therapy involving the use of aspirin and/or clopidogrel (ticagrelor), a lipid lowering therapy, an antianginal therapy and other secondary prevention therapies were all recommended to be prescribed for the patients as needed.

The strategies of MT or revascularization (CABG or PCI) were both alternative methods for treating a CTO vessel, and the final strategy decision was made based on the preferences of both the physicians and the patients. During the PCI procedure, the uses of contemporary techniques, such as microcatheters, bilateral injections and retrograde approaches, among others, were left to the discretion of the operator.

Definitions of variables and clinical endpoints
The definition of RCA-CTO was a TIMI flow grade of 0 within the RCA vessel with a duration of at least 3 months [22]. The occlusion interval was calculated from the last episode of myocardial infarction, from
the first episode of stable angina or from a previous angiogram. For those patients who exhibited no clinical symptoms, we artificially identified them as meeting our inclusion criteria. Diabetes mellitus was defined as a previous diagnosis of diabetes before hospitalization or a new diagnosis with either fasting blood glucose levels ≥ 7.0 mmol/L or a glucose level at 2 h after a meal of ≥ 11.1 mmol/L on more than two occasions [23]. A prior MI was confirmed according to the electrocardiogram, the ultrasonic cardiogram or the records from a hospital information system. An angiographic stenosis > 50% was considered to indicate a diseased vessel. Patients with a CTO vessel that represented the only diseased vessel were diagnosed with single-vessel disease. Systolic heart failure consisted of only heart failure with a reduced ejection fraction (HFrEF) and heart failure with a mid-range ejection fraction, which was defined as a left ventricular ejection fraction (LVEF) < 50% with either the presence of dyspnea or equivalent symptoms, while diastolic heart failure exhibited a preserved ejection fraction (HFpEF) [24]. Both right dominance and codominance were considered to be right dominance.

We predefined the primary clinical endpoint as all-cause death, which was mortality due to any reason. Other clinical outcomes included cardiac death, repeat nonfatal MI, repeat revascularization and target vessel revascularization (TVR). Cardiac death was defined as the following by the Academic Research Consortium (ARC) [25]: any death that is related to a cardiac reason, an unwitnessed mortality or an unknown cause of mortality. Noncardiac death was defined as death from a certain noncardiac cause. Probable/definite cardiac death was defined as death of certain (myocardial infarction, heart failure, etc.) or probable (sudden death) causes. The definition of a repeat MI was determined by following the third universal definition of MI [26]: a complicated consideration of ischemic symptoms, electrocardiograms and myocardial damage biomarkers. In this study, we collected data from only repeat nonfatal MI patients. The definition of repeat revascularization was an unplanned revascularization to the target vessel (RCA) or to the nontarget vessel. TVR was defined as any surgical bypass or percutaneous intervention of the target vessel (RCA).

### Data collection and follow-up

The data collection was managed by experienced raters who were trained beforehand in order to ensure accuracy. The following items were selected for analysis: age, gender, clinical history (hypertension, dyslipidaemia, prior MI and systolic heart failure, among others), inspection information, medical data and angiogram information. For the angiogram data, experienced, interventional physicians rescanned the cine angiograms by using standard definitions in order to reduce bias.

A minimum follow-up period of 12 months was predefined. A pre-designed chart including all of the follow-up items was applied. The follow-up procedure was performed by experienced investigators who were blind to the patients’ assigned groups. A phone call was the preferred follow-up method. For patients who had records of re-hospitalization in the Beijing Anzhen Hospital, noteworthy information was also obtained from the hospital information system. In the present study, we attempted to minimise the crossover between the groups. Thus, for patients who changed their treatment strategy during the follow-up, we ended the follow-up process at the moment of the treatment change. The follow-up information was evaluated by an adjudication board (Shuzheng Lyv, Hong Liu and Fei Yuan) who were blind to the patients’ assigned groups. The certifications of the clinical endpoints were also based on the decisions of the board.

### Statistical analysis

The continuous variables are presented as the means ± SDs (normal distribution) or medians with interquartile ranges (skewed distribution). Comparisons were performed using Student’s t-test or the Mann–Whitney U test, where appropriate. The categorical items are presented as numbers and percentages and were analyzed using the Chi-square test or Fisher’s exact test. Events per 1000 patient-years and survival curves using the Kaplan–Meier method were applied for all of the endpoints. A univariate Cox proportional hazard regression model was used to calculate the unadjusted hazard ratios (HRs). After the univariable analysis, a multivariable Cox regression model was performed. The potential adjusted factors were selected according to both the univariate Cox regression model (via the analysis of all of the variables listed in Table 1 and by using a threshold of P ≤ 0.2) and the relevant clinical implications. The adjusted HR (model 1) was calculated by utilizing the variables that exhibited statistical significance, as demonstrated by the univariate Cox regression: age, chronic kidney disease (CKD), chronic obstructive pulmonary disease (COPD)/asthma, prior MI, systolic heart failure (HF), LVEF, regional wall motion abnormality (RWMA), single-vessel disease, triple-vessel disease and syntax scores. Afterwards, we added further potential clinically relevant factors to calculate the adjusted HR (model 2): sex, peripheral vascular disease (PVD) and HbA1c. To further evaluate the difference between the CTO-SR and CTO-MT groups, we performed a propensity score-matched analysis. First, a propensity score utilizing a logistic regression model, which absorbed all of the variables listed in Table 1 (except for the retrograde
Table 1  Baseline characteristics (total population n = 943)

| Clinical characteristics | CTO-MT (n = 443) | CTO-SR (n = 500) | P value |
|--------------------------|------------------|------------------|---------|
| Age (years)              | 60.65 ± 10.58    | 59.97 ± 8.78     | 0.285   |
| Male                     | 322 (72.7)       | 392 (78.4)       | 0.041   |
| Hypertension             | 308 (69.5)       | 337 (67.4)       | 0.483   |
| Dyslipidemia             | 152 (34.3)       | 139 (27.8)       | 0.031   |
| PVD                      | 18 (4.1)         | 15 (3.0)         | 0.375   |
| Prior MI                 | 234 (52.8)       | 285 (57.0)       | 0.198   |
| Prior PCI                | 90 (20.3)        | 72 (14.4)        | 0.016   |
| Prior stroke             | 35 (7.9)         | 22 (4.4)         | 0.024   |
| Heart failure            | 125 (28.2)       | 149 (29.8)       | 0.593   |
| Systolic heart failure   | 52 (11.7)        | 63 (12.6)        | 0.686   |
| Diastolic heart failure  | 73 (16.5)        | 86 (17.2)        | 0.768   |
| CKD                      | 15 (3.4)         | 10 (2.0)         | 0.186   |
| COPD/asthma              | 4 (0.9)          | 4 (0.8)          | 0.864   |
| Hyperuricemia            | 97 (21.9)        | 86 (17.2)        | 0.069   |
| Smoking                  | 216 (48.8)       | 266 (53.2)       | 0.173   |
| Drinking                 | 74 (16.7)        | 91 (18.2)        | 0.546   |
| BMI (kg/m²)              | 26.46 ± 3.31     | 26.67 ± 3.02     | 0.304   |
| LVEF (%)                 | 61.00 (55.00–66.50) | 60.00 (55.00–66.00) | 0.153 |
| RWMA                     | 144 (32.5)       | 160 (32.0)       | 0.868   |
| Fasting blood glucose(mmol/L) | 7.20 (6.25–9.31) | 7.41 (6.13–9.26) | 0.727   |
| HbA1c (%)b               | 7.3 (6.7–8.0)    | 7.3 (6.9–7.9)    | 0.753   |
| Medical treatment        |                 |                 |         |
| Aspirin                  | 428 (96.8)       | 429 (98.4)       | 0.114   |
| P2Y12 inhibitor          | 396 (89.6)       | 410 (82.3)       | 0.001   |
| Statin                   | 422 (95.5)       | 462 (92.8)       | 0.080   |
| Nitrites                 | 262 (59.3)       | 152 (30.5)       | 0.000   |
| Beta-blocker             | 341 (77.1)       | 405 (81.3)       | 0.114   |
| CCB                      | 123 (27.8)       | 123 (24.7)       | 0.276   |
| ACEI/ARB                 | 246 (55.7)       | 272 (54.6)       | 0.750   |
| Insulin                  | 160 (36.1)       | 202 (40.4)       | 0.177   |
| Sulfonylureas            | 64 (14.4)        | 66 (13.2)        | 0.579   |
| Glindide                 | 20 (4.5)         | 18 (3.6)         | 0.476   |
| Biguanides               | 152 (34.3)       | 195 (39.0)       | 0.136   |
| Thiazolidinediones       | 48 (10.8)        | 54 (10.8)        | 0.986   |
| Alpha-glucosidase inhibitor | 159 (35.9)    | 151 (30.2)       | 0.063   |
| Angiographic characteristics |             |                 |         |
| Dominance artery (right) | 396 (89.4)       | 488 (97.6)       | 0.000   |
| Number of diseased vessels |             |                 |         |
| 1                        | 115 (26.0)       | 94 (18.8)        | 0.008   |
| 2                        | 151 (34.1)       | 201 (40.2)       | 0.053   |
| 3                        | 177 (40.0)       | 205 (41.0)       | 0.744   |
| Syntax score a           | 20.00 (13.00–27.00) | 20.00 (17.00–23.00) | 0.400 |
| Rentrop grade ≥ 2 a      | 309 (82.4)       | 368 (88.0)       | 0.025   |
| Abrupt stump a           | 188 (50.1)       | 205 (49.0)       | 0.759   |
| Calcification a          | 73 (19.5)        | 80 (19.1)        | 0.907   |
| Bending ≥ 45°a           | 327 (87.2)       | 361 (86.4)       | 0.729   |
| CTO length ≥ 20 mm a     | 278 (74.1)       | 308 (73.7)       | 0.886   |
approach), was calculated. The pairs were then matched at a 1:1 ratio by utilizing a nearest-neighbor matching method (caliper value = 0.02). We assessed the balance of the variables through absolute standardized differences (ASDs). ASDs < 10.0% revealed a relatively small imbalance. The baseline items of the propensity-matched population were then reanalyzed using the same methods as previously described. The clinical endpoints were also reanalyzed with the Kaplan–Meier method and the log-rank tests. The univariate Cox proportional hazard regression model was applied to calculate the HRs. To evaluate whether right coronary dominance affects clinical endpoints, an additional analysis of patients with only right dominance was also performed between the CTO-SR and CTO-MT groups.

Subgroup analysis was performed between subgroup populations: successful CTO-PCI versus initial CTO-MT, successful CTO-PCI versus failed CTO-PCI, successful CTO-PCI versus CTO-CABG, CTO-CABG versus initial CTO-MT and CTO-CABG versus failed CTO-PCI.

Another subgroup analysis, including sex (male/female), age (< 60 years old/≥ 60 years old), prior MI (yes/no), RWMA (yes/no), HbA1c (< 7.0%/≥ 7.0%), single-vessel disease (yes/no), systolic heart failure (yes/no), Rentrop grade ≥ 2 (yes/no) and syntax score (< 22/≥ 22), was also conducted using a multivariate Cox regression model. The covariates that were applied in this Cox regression model were the same as the items applied in model 2. Of note, we only performed this subgroup analysis for all-cause death and cardiac death.

Statistical analyses were performed using SPSS 24.0 (SPSS Inc., Chicago, Illinois, USA) and Stata 14.0 (Stata, College Station, TX, USA). A two-tailed P-value ≤ 0.05 was considered statistically significant.

Results
Baseline characteristics (total population)
From January 2007 to December 2017, a total of 943 stable RCA-CTO patients with diabetes were consecutively enrolled in the present study (Fig. 1). Of these patients, 443 (46.98%) were managed by MT (the CTO-MT group), and 500 (53.02%) were managed by successful revascularization (the CTO-SR group). The CTO-MT group enrolled patients with initial CTO-MT (n = 233), failed CTO-PCI (n = 191) and failed CTO-CABG (n = 19). The CTO-SR group enrolled patients with successful CTO-PCI (n = 309) and CTO-CABG (n = 191).

Clinical outcomes
In-hospital mortality occurred in four patients, and these patients were all treated with CABG: three patients in the CTO-SR group and one patient in the CTO-MT group.

| Procedural characteristics | CTO-MT (n = 443) | CTO-SR (n = 500) | P value |
|---------------------------|------------------|-----------------|--------|
| Retrograde approach<sup>c</sup> | 3 (1.6) | 41 (13.3) | 0.000 |
| Perforation<sup>c</sup> | 4 (0.9) | 1 (0.2) | 0.193 |
| Pericardial effusion<sup>c</sup> | 0 (0) | 0 (0) | – |
| Emergency surgery<sup>c</sup> | 0 (0) | 0 (0) | – |
| Contrast retention/dissection<sup>c</sup> | 2 (0.5) | 5 (1.0) | 0.457 |
| Thread off<sup>c</sup> | 1 (0.2) | 0 (0) | 0.470 |
| Sudden cardiac arrest<sup>c</sup> | 0 (0) | 1 (0.2) | 1.000 |
| Death during hospitalization<sup>c</sup> | 1 (0.2) | 3 (0.6) | 0.627 |

Values are n (%), mean ± SD or median with interquartile range

PCI percutaneous transluminal coronary intervention, MT medical therapy, CABG coronary artery bypass grafting, PVD peripheral vascular disease, MI myocardial infarction, CTO chronic total occlusion, COPD chronic obstructive pulmonary disease, LVEF left ventricular ejection fraction, BMI body mass index, CCB calcium-channel blocker, ACEI/ARB angiotensin converting enzyme inhibitor/angiotensin-receptor blocker; CTO chronic total occlusion, HF heart failure, RWMA regional wall motion abnormality

<sup>a</sup> Cine angiograms records got from 794 (84.10%) individuals
<sup>b</sup> HbA1c got from 896 (95.02%) individuals
<sup>c</sup> Only patients who were treated with PCI
When analyzing the mid-term clinical endpoints, we excluded these four patients. Procedural complications occurred in seven patients who were treated with failed CTO-PCI (the CTO-MT group): four patients exhibited contrast extravasation (no pericardial tamponade), two patients with coronary artery dissection, and the guidewire broke and remained in the distal iliac artery in one patient. Additionally, complications occurred in seven patients who were managed by successful CTO-PCI: one patient exhibited contrast extravasation (no pericardial tamponade), five patients had contrast retention and one patient experienced sudden cardiac arrest during the procedure.

Table 2 shows the clinical endpoints of this study. After a mid-term follow-up (CTO-MT: median: 42.00 months, interquartile range [IQR]: 24.00–78.25 months; CTO-SR: 48 [29.00–90.50] months), the data on all-cause death was successfully obtained from 874 (93.08%) patients: 28.15 all cause death per 1000 patient-years in the CTO-MT group versus 12.17 all cause death per 1000 patient-years in the CTO-SR group (P < 0.001). The univariate analysis (unadjusted HR: 0.423, 95% confidence interval [CI] 0.270–0.663) demonstrated that CTO-SR was superior to CTO-MT for all-cause death. This superiority was also confirmed by the multivariate analysis (adjusted HR [model 1]: 0.429, 95% CI 0.269–0.682; adjusted HR [model 2]: 0.445, 95% CI 0.278–0.714).

When considering the outcome of cardiac death, CTO-SR also demonstrated its superiority in events per 1000 patient-years (22.23 versus 6.90, P < 0.001), the univariate (unadjusted HR: 0.311, 95% CI 0.177–0.547) and multivariate (adjusted HR [model 1]: 0.308, 95% CI 0.172–0.550; adjusted HR [model 2]: 0.307, 95% CI 0.169–0.557) models. So as to probable/definite cardiac death (unadjusted HR: 0.297, 95% CI 0.137–0.644; adjusted HR [model 1]: 0.309, 95% CI 0.138–0.693; adjusted HR [model 2]: 0.341, 95% CI 0.151–0.771). The superiority of patients who were managed by SR was consistent for the clinical endpoints of repeat nonfatal MI (unadjusted HR:
Table 2  Clinical outcomes in all patients

| Event per 1000 patient-years | CTO-MT | CTO-SR | P value |
|-----------------------------|--------|--------|---------|
| All cause death             | 28.05  | 12.17  | 0.000   |
| Unadjusted HR (95% CI)      | 1      | 0.423 (0.270–0.663) | 0.000 |
| Adjusted HR (95% CI) Model 1| 1      | 0.429 (0.269–0.682) | 0.000 |
| Adjusted HR (95% CI) Model 2| 1      | 0.445 (0.278–0.714) | 0.001 |
| Noncardiac death            | 5.82   | 5.28   | 0.080   |
| Unadjusted HR (95% CI)      | 1      | 0.829 (0.370–1.860) | 0.649 |
| Adjusted HR (95% CI) Model 1| 1      | 1.351 (0.538–3.392) | 0.522 |
| Adjusted HR (95% CI) Model 2| 1      | 1.778 (0.684–4.618) | 0.237 |
| Cardiac death               | 22.23  | 6.90   | 0.000   |
| Unadjusted HR (95% CI)      | 1      | 0.311 (0.177–0.547) | 0.000 |
| Adjusted HR (95% CI) Model 1| 1      | 0.308 (0.172–0.550) | 0.000 |
| Adjusted HR (95% CI) Model 2| 1      | 0.307 (0.169–0.557) | 0.000 |
| Probable/definite cardiac death| 12.17 | 3.65   | 0.000   |
| Unadjusted HR (95% CI)      | 1      | 0.297 (0.137–0.644) | 0.002 |
| Adjusted HR (95% CI) Model 1| 1      | 0.309 (0.138–0.693) | 0.040 |
| Adjusted HR (95% CI) Model 2| 1      | 0.341 (0.151–0.771) | 0.010 |

Adjusted covariates (model 1): age, CKD, COPD/asthma, prior MI, systolic HF, LVEF, reginal wall motion abnormality, single vessel disease, triple-vessel disease and syntax scores

Adjusted covariates (model 2): age, sex, PVD, CKD, COPD/asthma, prior MI, systolic HF, LVEF, reginal wall motion abnormality, single vessel disease, triple-vessel disease, syntax scores and HbA1c

HR hazard ratio, CI confidence interval; other abbreviations as in Table 1

Propensity score matched analysis

After a 1:1 propensity matching, 286 patients were included in both groups. The ASDs after matching were all less than 10.0%, except for drinking (see Additional file 1: Figure S1). None of the baseline variables showed significant differences (see Additional file 1: Table S1), thus indicating a good matching balance. For the clinical endpoints, the superiority of CTO-SR was also observed for all-cause death (HR: 0.502, 95% CI 0.291–0.865, P = 0.013), cardiac death (HR: 0.331, 95% CI 0.161–0.681, P = 0.003), repeat revascularization (HR: 0.639, 95% CI 0.440–0.927, P = 0.018) and TVR (HR: 0.574, 95% CI 0.355–0.930, P = 0.024) when compared with CTO-MT (Table 3 and Fig. 3). Regarding repeat nonfatal MI, the trend was also consistent (HR: 0.571, 95% CI 0.277–1.177, P = 0.129).

Subgroup analysis

To minimize the influence of the dominance artery, an additional analysis that included only patients with RAC dominance was performed. We found that the superiority of CTO-SR was consistent in all clinical endpoints as in the whole population (see Additional file 1: Table S3, Fig. 4). The baseline characteristics are also provided in Additional file 1: Table S2.

Another subgroup analysis that compared successful CTO-PCI with failed CTO-PCI, initial CTO-MT and CTO-CABG, found that successful CTO-PCI provided patients with a benefit in terms of mortality when compared with initial CTO-MT (all-cause death: adjusted HR 0.498 [0.251–0.987]) and failed CTO-PCI (all-cause death: adjusted HR 0.315 [0.154–0.642]). This benefit can mainly be attributed to cardiac causes (cardiac death: 0.371 [0.154–0.892] and 0.195 [0.082–0.465]; probable/definite cardiac death: 0.463 [0.126–1.701] and 0.151 [0.041–0.563], respectively) (see Additional file 1: Tables S4–S7, Fig. 4). Regarding successful CTO-PCI versus CTO-CABG, CTO-CABG gained more benefit from repeat revascularization (adjusted HR 4.459 [2.044–9.726]) and TVR (21.676 [2.896–162.24]) (see Additional file 1: Tables S8, S9, Fig. 4).

We also compared CTO-CABG with both initial CTO-MT and failed CTO-PCI. Although statistical significance was not observed in all endpoints, CTO-CABG was superior to both subgroups in all-cause death, cardiac death, repeat revascularization and TVR. (see Additional file 1: Table S10–S13, Fig. 4).

Subgroup analysis based on clinically relevant variables showed that the superiority of CTO-SR in all-cause death was consistent in all subgroups, except for patients with single-vessel disease (HR: 1.245 [0.206–7.527]). The same results were also observed in cardiac death (Fig. 5). However, only 8 all-cause deaths and 4 cardiac deaths
occurred in patients with single-vessel disease. Furthermore, no significant difference was observed in patients with/without single-vessel disease (P > 0.05).

**Predictors of survival**

By multivariate analysis (Fig. 6), CTO-SR was a protected predictor of all-cause death (HR: 0.445, 95% CI 0.278–0.714) and cardiac death (HR: 0.307, 95% CI 0.169–0.557). Conversely, increases in syntax score (per 1 score, HR: 1.097, 95% CI 1.043–1.154) and age (per 1 year, HR: 1.044, 95% CI 1.020–1.068) predicted a worse probability for all-cause death. Additionally, an increase in the syntax score (per 1 score, HR: 1.104, 95% CI 1.043–1.169) was also related to a higher incidence of cardiac death.

**Discussion**

**Main discoveries**

To our knowledge, this is the first study that focused on stable RCA-CTO patients with diabetes and that attempted to determine if SR for RCA-CTO offered patients better clinical benefit than that offered by conservative MT. After a mid-term follow-up, our main findings are as follows: (1) CTO-SR offered patients more clinical benefits, which was demonstrated by both the multivariable Cox regression analysis, a propensity score-matched analysis and the subgroup analyses. (2) The subgroup analyses revealed that mortality benefits come from both successful CTO-PCI and CTO-CABG. While, the benefit of repeat revascularization and TVR mainly comes from the CTO-CABG subgroup. Furthermore, CTO-CABG also offered patients more repeat revascularization and TVR benefits when compared with successful CTO-PCI. (3) Syntax score was a harmful predictor of cardiac death. However, neither collateral circulation nor syntax score influenced the survival benefits of CTO-SR. So as to glycemic control (HbA1c).

**Conflicting debates on RCA-CTO treatment strategy**

It is well-known that the territory of the RCA is smaller than that of the LAD. Additionally, good collateral circulation has more often been observed in RCA-CTO [27]. Thus, some experts have argued that successful
Regarding procedural complications, in the present study, we calculated complications mainly from the PCI procedure. The incidence was low: seven for successful CTO-PCI and seven for failed CTO-PCI. After follow-up, we observed only six instances of repeat-revascularization (all were TVR) in those patients (four patients with successful CTO-PCI and two patients with failed CTO-PCI). No death or other adverse events occurred. We thought the influence of procedural complications on midterm clinical outcomes might be slight.

Why should diabetic patients with RCA-CTO be any different?

Previous findings can provide some clues to explain this issue. Prior studies [10, 34] have focused on patients with both RCA-CTO and other severe situations (older age, unprotected left main artery) and reported a higher incidence of mortality or cardiac mortality. A smaller RCA (left coronary dominance) is also a predictor of all-cause death in CTO patients [35]. Thus, the importance of RCA will be more obvious in patients with other severe conditions.

When considering diabetes, the SWISSI study [36] demonstrated that diabetes was a predictor of progressive coronary atherosclerosis disease (OR: 19.01, P = 0.026). The DIABETES study [37] reported that 50% of repeat revascularization procedures were due to progression in another vessel that was not previously treated. This finding was also confirmed by other experts [38]. We speculated that RCA-CTO is different in patients with diabetes because it is an inherently severe situation and exhibits a relatively fast progression, even though the RCA area might be smaller than the LAD area. One study [39] examined diabetic patients with a non-infarct-related CTO who suffered a ST-elevation myocardial infarction and demonstrated that CTO increased the risk of mortality when compared with those without CTO, which partially confirmed our speculation. Moreover, diabetic patients suffered a higher risk of any cardiovascular event than the risk in patients without diabetes [17]. From that perspective, the aim of successful revascularization is not only to recover the blood supply of the RCA area but also to support potential collateral circulation to either the LAD or the left circumflex coronary artery (LCX) in the case of an acute coronary occlusion.

Another possible explanation is electronic stability. The VACTO Primary Study [40] demonstrated that CTO was an independent risk factor for predicting the incidence of ventricular arrhythmias. Perfusion therapy can decrease the incidence of fatal ventricular arrhythmias [41]. Thus, electronic stability might partially explain the improvement in survival after successful revascularization.

To summarize, the inherent severity of the coronary artery situation in diabetic patients may make RCA-CTO different in diabetic patients than in the traditional patient

### Table 3 Clinical outcomes in propensity matched population

| Event | CTO-MT | CTO-SR | P value |
|-------|--------|--------|---------|
| All cause death | Event per 1000 patient-years | 30.15 | 15.03 | 0.010 |
| HR (95% CI) | 1 | 0.502 (0.291–0.865) | 0.013 |
| Non-cardiac death | Event per 1000 patient-years | 7.34 | 7.52 | 0.957 |
| HR (95% CI) | 1 | 1.038 (0.422–2.555) | 0.936 |
| Cardio death | Event per 1000 patient-years | 22.82 | 7.52 | 0.001 |
| HR (95% CI) | 1 | 0.331 (0.161–0.681) | 0.003 |
| Probable/definite cardiac death | Event per 1000 patient-years | 11.41 | 6.01 | 0.140 |
| HR (95% CI) | 1 | 0.524 (0.220–1.250) | 0.145 |
| Repeat nonfatal MI | Event per 1000 patient-years | 15.95 | 9.16 | 0.126 |
| HR (95% CI) | 1 | 0.571 (0.277–1.177) | 0.129 |
| Repeat revascularization | Event per 1000 patient-years | 61.79 | 39.08 | 0.012 |
| HR (95% CI) | 1 | 0.639 (0.440–0.927) | 0.018 |
| TVR | Event per 1000 patient-years | 37.70 | 21.45 | 0.018 |
| HR (95% CI) | 1 | 0.574 (0.355–0.930) | 0.024 |

HR hazard ratio, CI confidence interval; other abbreviations as in Table 1

revascularization for RCA-CTO will not benefit patients. Previous studies [5, 11] reported that successful PCI for RCA-CTO did not offer patient survival benefits, which may provide support for this plausible notion. However, Kalandyk and colleagues [28] reported that RCA-CTO increased the incidence of major adverse cardiac cerebrovascular events in patients undergoing CABG, which might mainly due to the smaller number of grafts to the RCA, indirectly indicating that RCA-CTO should be treated with revascularization. Direct positive evidence was also provided by Mitomo and colleagues [6]. However, all of the above studies focused on the entire CTO population. Diabetic status was associated with fast atherosclerosis progression and poor clinical outcomes [18, 29–33]. Under such a special situation, whether revascularization should be performed is unknown. Our study demonstrated that CTO-SR offers this specific patient group both survival benefits and other clinical benefits (lower incidence of repeat revascularization, TVR and repeat nonfatal MI). Moreover, we found that successful CTO-PCI offers patients only a mortality benefit, while CTO-CABG offers patients more clinical benefits than successful CTO-PCI does. This will surely better direct clinical practice.
The highlight of our study was demonstrating the successful revascularization benefits of patients in terms of not only survival items but also other clinical items. Furthermore, we demonstrated that successful CTO-PCI mainly offers patient survival benefits.

Will collateral circulation, coronary severity and glycemic level influence the superiority of successful revascularization?

In the present study, we observed that neither collateral circulation nor syntax score affected the superiority of CTO-SR, and neither did HbA1c.

In terms of collateral circulation, our results were consistent with those from a previous study [41], which enrolled CTO patients with well-developed collateral circulation and demonstrated that revascularization reduced the incidence of cardiac death. Mechanically, although good collateral circulation might offer blood to the occluded area of the RCA, the relationship between collaterals and clinical events remains unclear [30, 42, 43]. Additionally, low diastolic blood pressure [44] and coronary steel [45] can also reduce the protective effects of collaterals. Furthermore, FFR studies [46, 47] observed that collateral circulation affected the evaluation of donor vessel FFR and, hence, the optimal treatment strategy, which might lead to adverse cardiac events. Thus, reperfusion of the RCA-CTO will surely reduce myocardial ischemia as well as cardiac events. On the other hand, diabetic patients were often associated with impaired collateral circulation [23], which makes this reperfusion more essential to perform. In summary, our findings will correct the plausible notion that SR and medical therapy are both acceptable treatments for RCA-CTO patients with well-developed collateral circulation.

When considering coronary severity, we demonstrated that syntax score was a predictive risk factor for mortality. However, the syntax score did not influence the protective effect of CTO-SR. Even though for patients with single-vessel disease, CTO-SR might not offer survival benefit. However, only 209 patients were included in this subgroup, and mortality only occurred in eight patients.
cardiac death: four patients), which reduced the strength of the evidence. Further studies that focus on only a single-vessel RCA-CTO in diabetic patients are needed to verify our results.

Fig. 4 Kaplan–Meier curves for clinical endpoints in subgroup population. a Kaplan–Meier curves for all cause death in patients treated with initial CTO-MT versus successful CTO-PCI versus failed CTO-PCI versus CTO-CABG. b Kaplan–Meier curves for cardiac death

Fig. 5 Subgroup analysis for all-cause death and cardiac death. All HRs were adjusted for age, sex, PVD, CKD, COPD/asthma, prior MI, systolic HF, LVEF, regional wall motion abnormality, single vessel disease, triple-vessel disease, syntax scores and HbA1c (abbreviations as in Table 1)
Some experts observed a connection between glycemic control and clinical outcomes [48]. However, in the present study, we observed that the HbA1c level did not predict mortality. We speculated that the reason might be that the CTO-SR procedure is the main beneficial factor for patients with RCA-CTO. Further studies are still needed to confirm the benefits of glycemic control.

Limitations
The following limitations were present in this study. (1) This study was a retrospective cohort study. Some baseline variables were imbalanced. Although a propensity score-matched analysis was performed to balance any potential bias, the evidence grade is lower than that of a randomized controlled trial. (2) In the present study, cine angiograms could only be obtained from 794 (84.10%) patients. In the multivariable Cox regression analysis, we included syntax scores, which may omit some adverse cardiac events. (3) For the certifications of the clinical endpoints, TVR and repeat nonfatal MI were difficult to confirm. To ensure accuracy, only patients who had records of rehospitalization in the Beijing Anzhen Hospital or who could provide written certificates of diagnosis (by WeChat) were considered as meeting the definitions of TVR or repeat nonfatal MI. This consideration will introduce some bias. (4) We did not evaluate signs of a viable myocardium.

Conclusions
In summary, for stable RCA-CTO patients with diabetes, successful revascularization offered patients more clinical benefits than medical therapy. CTO-CABG might be a more recommended way to accomplish revascularization.

Additional file

Additional file 1: Table S1. Baseline Characteristics in Propensity-Matched Population. Table S2. Baseline Characteristics in right dominance RCA-CTO. Table S3. Baseline Characteristics (Right dominance RCA-CTO). Table S4. Baseline Characteristics (initial CTO-MT versus successful CTO-PCI). Table S5. Baseline Characteristics (Failed CTO-PCI versus successful CTO-PCI). Table S6. Clinical Outcomes (initial CTO-MT versus successful CTO-PCI). Table S7. Clinical Outcomes (Failed CTO-PCI versus successful CTO-PCI). Table S8. Clinical Outcomes (CTO-CABG versus successful CTO-PCI). Table S9. Clinical Outcomes (CTO-CABG versus successful CTO-PCI). Table S10. Clinical Outcomes (Initial CTO-MT versus CTO-CABG). Table S11. Clinical Outcomes (Initial CTO-MT versus CTO-CABG). Table S12. Baseline Characteristics (Failed CTO-PCI versus CTO-CABG). Table S13. Clinical Outcomes (Failed CTO-PCI versus CTO-CABG). Figure S1. Absolute Standard Difference before and after Propensity-Score-Matching.

Abbreviations
CTO: chronic total occlusion; PCI: percutaneous transluminal coronary intervention; MT: medical therapy; MACE: major adverse cardiac event; MI: myocardial infarction; TVR: target vessel revascularization; LAD: left anterior descending coronary artery; LCX: left circumflex artery; RCA: right coronary artery; TIMI: thrombolysis in myocardial infarction; AF: atrial fibrillation; CKD: chronic kidney disease; COPD: chronic obstructive pulmonary disease; LVEF: left ventricular ejection fraction; BMI: body mass index; CCB: calcium-channel blocker; CKD: chronic kidney disease; COPD: chronic obstructive pulmonary disease; LVEF: left ventricular ejection fraction; BMI: body mass index; CCB: calcium-channel blocker;
blocker; ACEI/ARB: angiotensin converting enzyme inhibitor/angiotensin-receptor blocker; HR: hazard ratio; CI: confidence interval.

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Authors’ contributions
YYF and SZL designed and supervised the study. YDF, DW, MZ, MDZ, FX, QMZ and XJG made contribution to data acquisition. HL, FY and SZL made contribution to endpoints adjudication. All authors read and approved the final manuscript.

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Availability of data and materials
The datasets generated and analyzed for this current study are available from the corresponding author upon reasonable request.

Ethics approval and consent to participate
The present study was approved by the Clinical Research Ethics Committee of Beijing Anzhen Hospital, Capital Medical University (No.: 2018008X). Informed consent was exempt by the committee.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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