A Decline in Glomerular Filtration Rate Rather than Renal Arterial Stenotic Lesions, per se, Predicts Cardiovascular-Renal Events in Type 2 Diabetic Patients

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Background: In diabetic patients with renal artery arteriosclerosis (RAAS), the factors associated with a greater risk for cardiovascular-renal events (CVREs) remain unclear: the decline in estimated glomerular filtration rate (eGFR) caused by RAAS or the advance of arteriosclerosis that causes RAAS. Hence, the features to determine which best predicts the onset of CVREs in such patients were compared.

Methods and Results: The renal arteries of 162 type 2 diabetes patients were assessed by using magnetic resonance angiography (RAAS diagnosed as arteriosclerotic stenosis ≥50%) and they were studied longitudinally over 7 years. The influence of the presence/absence of RAAS, a decline in eGFR, clinical factors, surrogate arteriosclerotic markers and ischemic markers on patient's CVREs were assessed. A Cox regression analysis showed the detection of RAAS to be an independent risk factor for CVREs (bilateral RAAS was an extremely strong risk factor for the development of CVREs within 1,000 days), as was the decline in eGFR in a logistic regression analysis; the latter being a more powerful risk factor for CVREs. A multiple regression analysis revealed angiopoietin-2, a marker of ischemia, to be a risk factor for the decline in eGFR.

Conclusions: A decline in renal function but not the renal arterial stenotic lesion itself appears to be associated with an increased incidence of CVREs in type 2 diabetic patients with RAAS. (Circ J 2013; 77: 2816–2822)

Key Words: Angiopoietin-2; Cardiovascular-renal event; Estimated glomerular filtration rate; Ischemic renal injury; Renal artery arteriosclerosis
Methods
This research is a longitudinal cohort study over 7 years. The participants of this study were type 2 diabetic patients whose physician suspected the existence of severe arteriosclerosis causing RAAS (Supplementary File 1-1A). The primary end-point of this study is an initial development of CVRE onset (in patients that suffered a CVRE) or the end of the 7-year follow-up period (in patients that did not experience a CVRE). The influence of the presence/absence of RAAS, a decline in eGFR, clinical factors and surrogate arteriosclerotic markers on patient’s CVREs were assessed.

The presence or absence of RAAS was confirmed by magnetic resonance imaging (MRI) using gadolinium contrast media in 200 diabetic out-patients that attended our hospital and who were enrolled into this longitudinal study between January and March 2003 (baseline). The subjects, who were registered into this study, were followed up as outpatients of our hospital or our associated hospital. The number of subjects was established based on a power analysis (Supplementary File 1-1B). Patients whose MRA images showed stenosis of 50% or more in the renal arteries,4 as assessed by at least 3 professionals (including a radiologist, cardiologist and nephrologist), were diagnosed with RAAS. A flowchart showing the number of patients included and that of the patients excluded during the follow-up period is provided in Supplementary File 1-1C. Ultimately, a total of 162 patients who met the aforementioned criteria were followed for 7 years, to determine whether they suffered a CVRE in that period.

The CVREs considered in this study consisted of death (from any cause), cerebral infarction, cerebral hemorrhage, transient cerebral ischemia, asymptomatic cerebrovascular disorder, myocardial infarction, angina pectoris, heart failure, initiation of dialysis, a doubling of serum creatinine, leg amputation and the implementation of revascularization in any organ. Only the initial event was counted even if a patient subsequently suffered additional CVREs. In diabetes, there are many cases of asymptomatic CVREs, so, with asymptomatic CVREs whose onset period is unclear, we designated, as the date of onset, the date when the primary physician was able to confirm CVREs (Supplementary File 1-1D). Chest X-rays, ECGs and MRIs of the brain were obtained at baseline for all patients and similar tests were performed when the onset of a CVRE was suspected. These tests were also performed at the end of the study, even in the absence of symptoms, and if a brain MRI revealed any abnormalities in asymptomatic cases, such as an increased size of infarction foci (as assessed by a radiologist), the patients were diagnosed as having suffered an asymptomatic cerebrovascular event. Each CVRE was diagnosed by more than 1 medical specialist from the corresponding area following established diagnostic guidelines.10,11 The authors were not involved at any stage of these diagnostic processes.

The presence or absence of RAAS [presented as RAAS (+) and RAAS (−), respectively] was examined by contrast MRA, and the eGFR, age, blood glucose, blood pressure and lipids, as well as the presence or absence of obesity, smoking, left ventricular hypertrophy (LVH) on ECG, and diabetic retinopathies were also evaluated at baseline. In addition, other parameters measured included the intima-media thickness (IMT), brachial-ankle pulse wave velocity (baPWV), adiponectin, interleukin-6 (IL-6), monocyte-chemoattractant protein (MCP-1), interleukin-1β, interferon-inducible protein (IP-10), vascular endothelial growth factor (VEGF), angiotensin-2, atrial natriuretic peptide (ANP) and brain natriuretic peptide (BNP), as well as the decline in eGFR from the baseline to the time of CVRE onset (in patients that suffered a CVRE) or until the end of the 7-year follow-up period (in patients that did not experience a CVRE). The influence of these factors on the onset of CVREs during the 7 years was studied, and we assessed if they constituted risk factors for RAAS.

This study received approval from the Ethics Committee at the Tohoku University Hospital, and all patients provided their informed written consent before any tests were performed.

Measurements
The methods used to measure adiponectin, IL-6, MCP-1, IP-10 and VEGF have been described previously.12 Angiopoietin-2 was determined using the commercially available Human Angiopoietin-2 Immunoassay Kit (Human R&D Systems Inc, MN, USA). The measuring method of IMT13 and baPWV14 adopted in this study is shown in Supplementary File 1-1E. The equation for calculating the eGFR is described in Supplementary File 1-1F.

Statistical Analyses
Numerical values with a normal distribution are expressed as the mean±SEM, and those that did not entirely conform to a normal distribution are expressed as the median (range). The t-test was used to perform inter-group comparisons of normally distributed numeric measurements and the Mann-Whitney U-test was used for those measurements that were not distributed normally. The χ2-test was used to compare the prevalence rate of diabetic retinopathy, rate of smoking, CVRE incidence, RAAS prevalence rate and other factors between groups.

In addition, in this longitudinal study, 3 types of analyses were performed: analysis that uses CVREs as the dependent variable and only those factors that could be identified at the baseline as independent variables constitutes a prospective analysis, so a Cox regression analysis was used. In contrast, because analysis that added the percentage change of eGFR to the independent factor becomes a retrospective analysis, a logistic regression analysis was used. A multiple regression analysis was used for items where dependent variables were in numerical form (eg, percentage changes of eGFR and angiopoietin-2 levels, etc.), rather than categories such as “presence or absence of CVREs” (Supplementary File 1-1G).

The method of selecting the factors that should be analyzed as independent factors in the Cox regression analysis is shown in Supplementary File 1-1H.

Results
The baseline clinical characteristics of all 162 patients were analyzed as a whole and, in addition, these parameters were compared between RAAS (+) (n=36) and RAAS (−) (n=126) patients (Table S2). Compared with the RAAS (−) patients, RAAS (+) patients had lower hemoglobin A1c (HbA1c), eGFR and high-density lipoprotein cholesterol (HDL-C) levels, and higher systolic blood pressure (SBP), heart rate (HR), IMT, baPWV, diabetic retinopathy prevalence and smoking rate, as well as higher adiponectin, IL-6, MCP-1, IP-10, VEGF and angiopoietin-2 levels. With regards to drug administration, the proportion of patients that received each type of treatment was 88.9% RAAS (+) and 64.3% RAAS (−) (P=0.86) for rennin angiotensin system inhibitors (RASIs); 97.2% and 57.2% (P=0.36) for calcium channel blockers (CCBs); 83.3% and 33.3% (P=0.42) for other anti-hypertensive agents; 80.6% and 38.8% (P=0.37) for insulin therapy; 66.7% and 77.8% (P=0.79) for oral hypoglycemic agents; 58.3% and 42.9% (P=0.89) for statins; and 80.6% and 26.2%
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patients suffered RAAS at baseline (P<0.01). Furthermore, while the eGFR (ml · min$^{-1}$ · 1.73 m$^{-2}$) decreased by 52.3 ± 2.3% in the CVRE (+) patients (from 64.2 ± 5.1 to 29.6 ± 2.3 ml · min$^{-1}$ · 1.73 m$^{-2}$), it decreased less in the CVRE (–) patients (16.4 ± 1.5%: from 83.6 ± 2.4 to 69.7 ± 2.2 ml · min$^{-1}$ · 1.73 m$^{-2}$). Hence, the CVRE (+) patients had a lower baseline eGFR (P<0.001) and a greater decline in eGFR (P<0.001) than those in the CVRE (–) Group.

The rate of RASI use by the RAAS (+) patients was no different from that of the RAAS (–) group, nor were there differences in the % change in eGFR between RASI monotherapy (“mono”) and concomitant therapy (“dual”) in both RAAS (+) and RAAS (–) groups. However, both therapies provoked a greater % decline in eGFR in the RAAS (+) as opposed to the respective therapy in the RAAS (–) patients (P<0.01, Table S7). When the patients were subdivided into LVH (+) and LVH (–) groups, the LVH (+) group showed a greater incidence of CVREs, % decline of eGFR and prevalence of RAAS than the LVH (–) group (Table S8).

There was a lower incidence of CVREs and a smaller decline in eGFR in the RAAS (–) group than in the RAAS (+) group. Moreover, patients with bilateral RAAS had a greater incidence rate of CVREs than those with unilateral RAAS. The changes

(P=0.34) for anti-platelet agents (Table S4).

CVREs occurred in 24.7% (40 patients) of all the participants during the 7-year study period. When analyzed according to the presence or absence of RAAS, 75.0% (27 patients) of RAAS (+) patients and 10.3% (13 patients) of RAAS (–) patients developed CVREs, reflecting the significantly higher occurrence of these events in RAAS (+) patients (P<0.001). There were more RAAS (+) patients than RAAS (–) patients when CVREs were considered by type, except for the category “others” (Table S5). Among the RAAS (+) patients, the incidence of CVREs was concentrated in the first half of the observation period (1–4 years), while patients in the RAAS (–) group developed CVREs with equal frequency each year (Table S6).

In terms of function of the development of CVREs during the 7-year follow-up period, the patients were subdivided into 2 groups: CVRE (–) (n=122) and CVRE (+) (n=40). When the baseline clinical characteristics of these groups were compared (Table S3), age, SBP, IMT, baPWV, ANP, BNP, MCP-1, IP-10, angiopoietin-2, presence of LVH and RAAS (+) prevalence were all higher in the CVRE (+) Group, while HbA1c, eGFR and HDL-C levels were lower. RAAS (+) patients accounted for as many as 67.5% (26/40) of the patients in the CVRE (+) Group, whereas only 7.4% (9/122) of CVRE (–) patients accounted for as many as 67.5% (26/40) of the patients in the CVRE (+) Group, whereas only 7.4% (9/122) of CVRE (–) patients.
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Figure. Probability of not experiencing a CVRE over time. Compared with RAAS (-) patients, the probability of not experiencing CVREs continued to decrease sharply over time in the RAAS (+) patients, with a particularly sharp decrease during the first 1,000 days (A). The probability decreased even in unilateral RAAS patients, although patients with RAAS in both renal arteries (bilateral) experienced an extremely sharp decrease in the probability of not experiencing CVREs, with all the patients having developed CVRE within 1,000 days (B), indicating that bilateral RAAS carries an extremely high risk for CVRE onset. CVRE, cardiovascular-renal events; RAAS, renal artery arteriosclerosis.

Cox regression analyses were performed using CVREs as the dependent variable, and SBP, IMT, BNP, RAAS and LVH as the independent variables (Supplementary File 1-1H) (Table 1). The factor that influenced CVREs most strongly was RAAS, with a hazard ratio of 3.53 (95% CI: 2.36–5.27) and P<0.001, followed by SBP with a hazard ratio of 1.02 (95% CI: 1.00–1.03) and P=0.012; these 2 variables were significant independent risk factors for CVREs. By contrast, another Cox regression analysis that used CVREs as the dependent variable, and MCP-1, IP-10, urinary 8-hydroxydeoxyguanosine (8-OHdG), and angiopoietin-2 as the independent variables (Supplementary File 1-1H) only identified angiopoietin-2 to be a significant independent risk factor (P<0.001, Table 2). We therefore conducted a Cox regression analysis, using RAAS and angiopoietin-2 as the independent variables. The results showed that both RAAS and angiopoietin-2 were independent factors for CVREs (Table S10). Of these, RAAS had the largest hazard ratio of 1.84 (95% CI: 1.18–2.85).
A logistic regression analysis using the presence or absence of CVREs as a dependent variable identified the percentage change of eGFR (P=0.001) and angiotensin-2 (P=0.001) as independent risk factors, while RAAS was not even close to being a significant factor (P=0.678). Accordingly, the decline in eGFR was identified as a risk factor, while RAAS no longer represented a risk factor for the occurrence of CVREs (Table 3). This showed that RAAS and the percentage change of eGFR were strongly related (Table S11), and that the percentage change of eGFR was even more strongly related to CVREs (Table 3). Intima-media thickness, which is an indicator for systemic arteriosclerosis, can be considered to be closely related to RAAS (Table S11). Because the systemic vascular disorders and the renal damage progress appears concomitantly in diabetes mellitus, and given that RAAS is directly related to both conditions, it is assumed that RAAS in diabetic patients might serve as a potent predictor of CVREs. Angiotensin-2 was also identified as an independent risk factor of RAAS (Table S12). However, the percentage change of eGFR and IMT were stronger risk factors of RAAS than angiotensin-2 (Table S13). Furthermore, eGFR, angiotensin-2, and RAAS were very closely related each other (Tables S14-S15).

Compared with the RAAS (−) patients, the probability that RAAS (+) patients did not develop a CVRE continued to decrease as the study progressed, decreasing particularly sharply over the first 1,000 days (Figure A). Such decreases were even evident in the unilateral RAAS patients, even though it was less severe than in bilateral RAAS patients (Figure B), indicating that bilateral RAAS (+) patients are at high risk of experiencing a CVRE. The differences of the background between patients who developed CVREs during the first 1,000 days after the observation began, and patients who developed CVREs after the first 1,000 days are shown in Table S16. Twenty-three subjects developed CVREs within 1,000 days (Group A), and 17 developed CVREs after 1,000 days (Group B). Compared with Group B subjects, those in Group A had higher HR, IMT, MCP-1, and angiotensin-2 values, and lower HDL-C values. Group A also had a greater decline in eGFR than Group B, and Group A had a higher rate of RAAS (+), especially that of bilateral RAAS (+), than Group B. We therefore conducted a logistic regression analysis, using the onset of CVREs within 1,000 days as the dependent variable, and the HR, IMT, and the percentage change of eGFR, as well as MCP-1, angiotensin-2, HDL-C, and RAAS (+), as the independent variables (Table S17). However, only the percentage change of eGFR was identified as a factor that induced CVREs within 1,000 days, not angiotensin-2 and RAAS (Table S18).

Fortunately, these results did not change, even after eliminating the 5 patients who developed ESRD and after making an analysis.

**Discussion**

Until now, the RAAS (+) lesion itself was believed to be a risk factor for CVREs. We set up the hypothesis that, under conditions where RAAS is not only caused by RAAS (+) itself but also because of peripheral vascular disorders, intra-renal ischemia would increase, tubulointerstitial disorder has progressed, the level of eGFR would drop, and CVREs would increase as a result. In this longitudinal study, we clarified that RAAS (+) and a decline in eGFR are both independent risk factors for CVREs in diabetes mellitus patients; the latter being more powerful than the former. We then revealed that this decline in eGFR became an even stronger risk factor for CVREs than the presence of RAAS (+) itself, with bilateral RAAS (+) being an extremely strong risk factor. In particular, bilateral RAAS (+) was an extremely strong risk factor for developing CVREs within 1,000 days. However, the greater chance of developing CVREs within 1,000 days seemed to be due to the sharp decline in eGFR of the patient with bilateral RAAS (+). In contrast, angiotensin-2, which is an ischemic marker, was identified as a risk factor for the decline in eGFR.

All the diabetic patients enrolled on this study were suspected of having RAAS and therefore underwent a MRA, from which only renal arterial stenosis ≥50% was considered as significant arteriosclerosis, irrespective of whether it was clinically relevant stenosis. In fact, there were patients with no changes in RASIs renal perfusion scintigraphy, and some without significant changes in pressure between the renal and aortic sides of the stenotic lesion. Accordingly, we assumed that the eGFR decline detected in our patients with RAAS could be attributed to arteriosclerosis progressing at a more peripheral level than in the renal artery and that it had induced RAAS in these patients, rather than this event being due to the RAAS lesion itself. Although the degree of stenosis in the renal artery is not necessarily related to the progress in the decline in eGFR, and no differences in changes in renal function were evident between patients with RAAS treated by angioplasty or pharmacologically, it is thought that the subsequent onset of CVREs in diabetic patients with RAAS is closely related to the low eGFR values. Accordingly, CVREs might occur as a result of the decline in eGFR but not due to the high degree of renal artery stenosis. In other words, in addition to the impaired renal blood flow, provoked by the renal damage directly caused by a reduced renal arterial diameter, diabetic patients with RAAS also suffer impairments in the more peripheral intrarenal microcirculation, both of which would cause more profound ischemic damage to the renal parenchyma and might be reflected in the marked decline in their eGFR. Moreover, the IMT and RAAS (+) were shown to be independent risk factors of the long-term decline in eGFR here, indicating that not only local but also systemic arteriosclerosis is related to a long-term deterioration of renal function. (Table S11) Renal arterial stenotic lesions are probably just one part of the systemic arteriosclerotic lesions. Indeed, RAAS was strongly correlated with cardiovascular markers and RAAS (+) patients frequently developed CVREs as a complication, both here and elsewhere. RAAS is also closely related to arteriosclerosis of the coronary artery, as well as to a significant reduction in cardiac function, which when combined with the reduction in renal function is deemed to make patients prone to heart failure. Indeed, our bilateral RAAS (+) patients were deemed to be especially prone to experience a decline in eGFR, even despite the more aggressive interventions compared to those offered to individuals with unilateral RAAS. Moreover, it appeared that the stronger the decrease in eGFR in a patient, the sooner he/she was likely to suffer a CVRE. Although there were no differences in the incidence of CVREs and in the decline in eGFR between unilateral and bilateral RAAS patients, the time until the onset of CVREs did differ. CVREs are likely to be induced more readily in unilateral RAAS (+) than in RAAS (−) cases, moreover, bilateral RAAS (+) cases were shown to be more susceptible to CVREs, within even a shorter period of time (within ca. 1,000 days). Only the percentage change of eGFR was identified as a factor that induced CVREs within 1,000 days, not angiotensin-2 and RAAS (Supplementary File 1-7). Therefore, a rapid and significant decline in renal function, rather than ischemia, appears to be closely related to rapidly developing CVREs, and a decline in renal function is a factor that directly triggers CVREs. RAAS can be considered to con-
stitute a risk that aggravates survival as it is closely correlated to the decline in eGFR, an index for the progression of renal dysfunction that more strongly correlated to the onset of CVREs than RAAS itself. This decline in eGFR in diabetic patients with RAAS not only reflects the ischemia in the kidney but might also imply that similar arteriosclerosis has occurred in other organs (heart, brain and foot etc.), and accordingly, organ ischemia has advanced. Thus, when a patient experiences a rapid decline in eGFR, not only should RAAS be suspected but also, the aggravation of systemic ischemic lesions should be assessed. We thus believe it important to monitor eGFR in daily clinical practice.

In this study, angiopoietin-2 and VEGF were evaluated as markers of ischemia; it is angiopoietin-2 that is more closely related to CVREs. Angiopoietin-2 is believed to be produced mainly in connection with neovascularization of ischemic lesions. Blood angiopoietin-2 levels rise in line with the decline in renal function and they serve as a predictor of CVREs. Thus, it appears likely that impaired intrarenal blood flow due to arteriosclerosis, as well as renal damage due to renal ischemia caused by sclerotic changes in peripheral vessels (ie, arterioles) rather than by RAAS itself, and the increased angiopoietin-2 levels triggered under these circumstances, might intervene in a complicated manner to cause further cardiovascular/renal disorders. Accordingly, a decline in renal function and elevated angiopoietin-2, as opposed to stenotic lesion itself, might be associated with the heightened incidence of CVREs in type 2 diabetic patients with RAAS (Supplementary File 1-8). By combining the evaluation of albuminuria, which is currently being conducted in diabetes, with the evaluation of the change in eGFR, we can expect to see the advent of higher-precision medical treatment.

One of the limitations of this study is the low number of patients analyzed, with 38 out of the 200 patients having dropped out.

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Disclosure

None.

References


**Supplementary Files**

**Supplementary File 1**

1. Preparation for the study and passage of the study
2. Clinical characteristics of the study subjects, and the comparisons between the RAAS (+) and RAAS (−) groups at baseline or CVREs (+) and CVREs (−) during the 7-year follow up
3. Differences in the incidence of CVREs in the RAAS (+) and the RAAS (−) groups
4. Result of a cox regression analysis that used CVREs as the dependent variable
5. Results of logistic regression analyses using RAAS as a dependent variable
6. Results of multiple regression analyses
7. Baseline characteristics of the subjects who developed CVREs during the first 1,000 days and the independent risk factors that used CVREs were developed during the first 1,000 days
8. Relationship and the pathophysiology of the renal artery injury and CVREs

Please find supplementary file(s); http://dx.doi.org/10.1253/circj.CJ-13-0269