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The relationship between acute kidney injury and chronic kidney disease in patients with Type 2 Diabetes: an observational cohort study

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Significance Statement

There is currently a limited understanding of the interplay between AKI and CKD in people with type 2 diabetes and how this compares to the non-diabetic population. Through development of an algorithm which can be applied to routinely collected biochemistry data, this study has quantified the risk of AKI in patients with diabetes and how this relates to CKD. These findings have both important epidemiological and clinical implications demonstrating that the risk of AKI and associated adverse outcomes in this population of patients is currently underestimated. Increasing awareness may allow for implementation of simple interventions which prevent the occurrence of AKI thereby improving patient outcomes.

Abstract

Background

Type 2 diabetes is one of the leading causes of chronic kidney disease (CKD) and an independent risk factor for Acute Kidney Injury (AKI). This study aims to evaluate rates of AKI and how this relates to CKD status and further renal function decline in patients with and without type 2 diabetes using electronic healthcare records.

Methods

Study design was a retrospective cohort study. The negative-binomial model for counts with follow-up time as offset, adjusted for sex and age was used to evaluate AKI rates in people with and without diabetes depending on CKD status. A mixed effect linear model adjusted for demographic characteristics and co-morbidities was developed to evaluate decline in glomerular filtration rate (GFR) before and after an AKI event depending on diabetes and CKD status.

Results

The cohort was formed of 16700 participants with a median follow-up of 8.2 years. 9417 of these had type 2 diabetes and 7283 had no diabetes. 48.6% (N=4580) of participants with diabetes developed AKI compared to 17.2% (N=1257) of controls. 46.3% (N=4359) of those with diabetes had CKD vs 17.1% (N=1251) of controls. In the absence of CKD, AKI rate was five times higher in people with diabetes than controls (121.5 vs 24.6 per 1000 person-years, Rate Ratio RR=4.9, 95% CI 4.4-5.5), whereas for people with CKD, rate of AKI was twice higher in people with diabetes than controls (384.8 vs 180.0 per 1000 person-years, RR=2.1, 95% CI 1.9-2.4 after CKD date and 109.3 vs 47.4 per 1000 person-years, RR=2.3, 95% CI 1.8-3.0 prior to CKD). Fall in eGFR slope before AKI was steeper in people with diabetes compared to those without diabetes. After AKI episodes, loss of eGFR became steeper in people without diabetes, but did not increase in those with diabetes and pre-existing CKD.

Conclusion

Rates of AKI are significantly higher in patients with diabetes compared to patients without diabetes, and this remains true for individuals with pre-existing CKD.

Keywords: acute kidney injury; chronic kidney disease; type 2 diabetes; epidemiology; incidence.

The relationship between AKI and CKD in patients with T2DM



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1 Introduction

Type 2 diabetes is one of the leading causes of chronic kidney disease (CKD) and end-stage renal disease worldwide ¹. A large proportion of patients who develop CKD experience prior episodes of acute kidney injury (AKI), with evidence suggesting that kidney function does not fully recover following the AKI event¹. Moreover, CKD is a well-known risk factor for AKI, with recent studies suggesting that there is a considerable overlap between the pathophysiology underlying the two conditions ². However the relationship is likely to be complex and remains poorly understood.

8 Type 2 Diabetes (T2D) has been reported as an independent risk factor for AKI in previous observational 9 studies ^{3,4} and progressive decline in kidney function has also been well described in this population ¹. Both AKI and CKD have been identified as risk factors for cardiovascular disease ⁵, which is the most 10 11 frequent complication in T2D. Despite the increased access to routinely collected health care data, there are few observational studies evaluating the risk of AKI in people with T2D ^{6,7}, and even fewer 12 13 simultaneously investigating AKI and CKD in this population ¹. As a result, there is a limited 14 understanding of the interplay between AKI and CKD in people with T2D and how this compares to the non-diabetic population.⁸ Previously, quantification of AKI from routine health care data was limited to 15 the use of hospitalization and death using International Classification of Diseases (ICD) coding ⁶. More 16 17 recently, the Kidney Disease Improving Global Outcome (KDIGO) definition for AKI based on changes in serum creatinine (SCr) has been universally adopted which has enabled a more uniform approach ^{9,10}. 18 19 However, this approach comes with its challenges, which mainly relate to the application of the KDIGO 20 definition. In clinical practice AKI can only be identified when previous tests within a time window are 21 available for comparison, which may not be the case when blood testing is infrequent. To overcome 22 this, various time windows to define baseline creatinine have been proposed, including the use of both prior and post index values ¹¹⁻¹⁶. Despite the numerous definitions, the variation in the intensity of blood 23 sampling may still lead to misclassification between AKI and CKD¹⁰. This highlights the importance of 24

accurate definitions for both AKI and CKD that can be used in database studies to help understand the
 contribution of AKI to CKD and CKD progression, as well as the risk of developing AKI in patients with
 CKD.

The aim of this study was to develop an algorithm to examine rates of AKI in patients with and without T2D depending on CKD status using routinely collected healthcare data, and to investigate whether the association between AKI on GFR decline is different in people with T2D compared to people without diabetes.

32

33 Methods

34 Study population

35 The design is a retrospective cohort study of people from the Tayside region of Scotland (n = 402641 on 36 1 January 2012) which represents about 8% of the Scottish population. People with and without type 2 37 diabetes that were matched by age, sex and general practice were recruited in the Genetic of Diabetes 38 Audit and Research in Tayside Study (GoDARTS) from December 1998 to October 2012 which includes either at diabetes or eye screening clinics or through their GP¹⁷. About 50% of the patients with T2D at 39 that time from Tayside region were recruited into GoDARTS⁸. Participants attended a clinic at 40 41 recruitment, where a serum sample was collected to allow a number of routine biochemical measures 42 to be measured. Recruitment was treated as the baseline for this study with participants being followed 43 up until May 2017 using comprehensive electronic records. 44 The current study includes participants from GoDARTS with type 2 diabetes at baseline to form the 45 diabetic group and patients with no diabetes to form the control group. To allow for an accurate estimation of AKI rate in patients without diabetes, patients from GoDARTS who develop diabetes later 46 47 during the follow-up time were not included in the study. Also, patients without SCr measures on or 48 after recruitment were not included. For the eGFR slope analysis, patients with three or more SCr values

with at least one-year gap between the first and last measure prior to the first AKI episode (if applicable)
and three or more SCr measures after the AKI episode with at least 90 days gap between the first and
last of these measures were included. Patients with an AKI event prior to analysis were excluded.

52

53 Datasets and variables

54 The GoDARTS study was linked through an individual-specific anonymised identifier to the following 55 clinical datasets: information on diabetes including type of diabetes and date of diagnosis was acquired 56 from the Scottish Care Information – Diabetes Collaboration (SCI-DC) Diabetes Summary and 57 Longitudinal data ¹⁸. SCr values were obtained from the laboratory biochemistry system, comprising of 58 SCr measures from both primary and secondary care. The Scottish Renal Registry was used to identify patients receiving renal replacement therapy (RRT) and date of therapy initiation ¹⁹. The Scottish 59 60 Morbidity Records 01 (SMR01) for hospital admission was used to evaluate patient comorbidities 61 including coronary artery disease, congestive heart failure, peripheral vascular disease, cerebrovascular 62 disease and liver disease based on ICD-10 codes at admissions prior to recruitment. The community 63 prescribing data was used to assess whether the patient have been prescribed any of the following 64 classes of anti-hypertensive drugs: diuretics, angiotensin converting enzyme (ACE) Inhibitors, angiotensin receptor blockers (ARBs), beta-blockers and calcium channel blockers ²⁰. The demographics 65 66 dataset was used to determine participant sex and date of birth which was used to calculate age at 67 recruitment. Patients who had moved out of Tayside health board were treated as lost to follow-up. The 68 Community Health Index death dataset (CHI - the NHS Scotland population register) was used to obtain 69 date of death. Follow-up time was defined as the time from recruitment until May 2017, or date of RRT, 70 or date of death, or date the patient moved out of Tayside health board, which ever occurred first.

71

72

73 Development of an algorithm to identify AKI episode from serum creatinine tests

74 SCr measures from Jan 1988 to May 2017 were used in the analysis; measures obtained after initiation 75 of RRT were not included. All assays in the region are done in the same regional laboratory, and SCr 76 measures were adjusted for changes in assays over time. AKI was defined based on the KDIGO criteria⁹. 77 As testing was infrequent with large time gaps in some patients, leading to a lack of baseline being 78 calculated, we developed an algorithm to calculate baseline creatinine incorporating both prior and 79 post index creatinine measurements in the definition of baseline (Table 1). Severity of AKI (Stages 1-3, Table 1) was defined using KDIGO criteria⁹. To identify AKI episodes, SCr that were within seven days 80 81 apart were grouped into single episodes of care. Within the episode of care, a 1.2-fold increase in 82 creatinine from baseline was used to evaluate SCr values measured before and after each Scr value 83 flagged as AKI case in order to assess AKI initiation and recovery and determine the start and the end of the AKI episode ²¹. Furthermore, if two AKI episodes were within seven days apart then the two 84 85 episodes and SCr values in-between were grouped into one AKI episode²¹. The length of AKI episode 86 was calculated based on start and end dates of the AKI episode and was used to assess whether AKI had progressed to Acute Kidney Disease (AKD), defined as an AKI lasting more than seven days ²². The 87 88 highest AKI stage within the episode was used to define the stage of the AKI episode.

89

90 Estimated Glomerular filtration rate (eGFR) and CKD status

The CKD-EPI formula was used to estimate glomerular filtration rate (eGFR) from serum creatinine ²³.
Development of CKD was defined according to the CKD-KDIGO guideline as eGFR < 60 ml/min per
1.73m² present on at least two occasions at least 90 days apart ²³. To avoid misclassification between
AKI and CKD, eGFR values contained within AKI episodes were first removed from the longitudinal data.
The variation in the intensity of blood sampling, led to eGFR estimates either too distant (in healthy
individuals) or too dense over time (in sicker patients). As a result a median smoother was applied to the

97 remaining eGFR values based on a set of rules derived from the CKD-KDIGO definition as follows; for
98 each date of index blood test, three eGFR baseline values were calculated using the median eGFR for the
99 period 365 to 91 days prior to the index date, then 7 days prior to 7 days after index, and 91 to 365 days
100 after index date respectively. CKD diagnostic date was established when at least two of the three
101 medians were below 60 ml/min per 1.73m² (Table 2).The CKD date was then compared against
102 recruitment date to determine whether participants had prevalent CKD at recruitment or they
103 developed incident CKD during follow-up.

104

105 Primary and secondary outcomes

The primary outcome was the number of AKI episodes per person during follow-up, which was used to calculate AKI episode rates per 1000 patients per year (including recurrent events) and AKI rate ratios in people with type 2 diabetes vs non-diabetes depending on CKD status. The secondary outcome was eGFR decline over time calculated as the eGFR slope of the linear regression model per one-year unit increase. Other outcomes were number of patients experiencing AKI during follow-up, length of AKI episodes and AKI stage.

112

113 Statistical methods to analyse AKI rates depending on CKD status

114 Counts and proportions for categorical variables and mean and standard deviation (SD) or median and

115 inter-quartile range (IQR) for quantitative data were used to describe the demographic characteristics.

116 These were reported in people with and without diabetes and by CKD status (no-CKD at recruitment or

- during follow-up, pre-CKD to account for the period prior to CKD development for those that developed
- 118 CKD during follow-up, and post-CKD to include the post- CKD period for those that had CKD at

119 recruitment or developed CKD during follow-up. The difference between two independent proportions

120 were calculated based on Wilson's method ²⁴. The negative-binomial model for counts with log-link and

121 follow-up time as offset was used to analyse the primary outcome and to estimate rates of AKI episodes 122 in cases and controls depending on CKD status. The relationship between the outcome and the 123 explanatory variable (sex, age and diabetes status) was assumed linear via the log-link function ²⁵. Un-124 adjusted AKI rates and rates adjusted for age and sex were provided together with the corresponding 125 rate ratios (RRs) for association. Further adjustment for co-morbidities at recruitment was performed to 126 investigate how much of the effect of diabetes on AKI incidence rates can be explained by pre-existing 127 co-morbidities. The chi-square test was used to investigate the association between diabetes and AKI 128 stage and the non-parametric Mann-Whitney test was used to investigate differences in the length of 129 AKI episodes between the T2D vs control groups. 130 Sensitivity analyses was conducted to evaluate and compare incidence rates for stages 2 and stage 3

AKIs, and AKIs longer than 48 hrs respectively, as well as for AKIs occurring during hospital admission in
people with diabetes vs controls.

133

134 Statistical methods to analyse of longitudinal eGFR data

135 EGFR values measured during AKI episodes were first removed from the data and replaced at the start 136 of the episode with a baseline eGFR calculated as the median eGFR for the seven days prior to the AKI 137 episode if measures were available, otherwise the median eGFR of values measured between 365 and 8 138 days prior to the start of the AKI episode was used. A linear-mixed effect model was used to analyse the 139 association between AKI and eGFR decline from the longitudinal eGFR data. AKI was included into the 140 model as a time-varying factor with three levels: no AKI for patient with no AKI event during the follow-141 up, pre-AKI for patient with an AKI event during follow-up for the period prior to the AKI and post-AKI for the period after the AKI episode ²⁶. To identify significant changes in eGFR slope pre and post AKI 142 143 event and whether these changes differ between people with T2D and controls an interaction term 144 between AKI, T2D status and time was accommodated into the model. Baseline variables such as sex,

145 age (treated as age groups) and presence of cardiovascular diseases were fitted into the model with 146 both fixed intercept and slope. An interaction between these variable and T2D was also included and 147 Akaike Information Criterion (AIC) was used for variable selection. Given the strong interaction effects 148 between AKI, diabetes and CKD status, the analysis was conducted separately for people with no CKD at 149 recruitment and those that had an established CKD diagnosis prior to recruitment. The mixed model was 150 fitted with both random intercept and slope per individual before and after the AKI episode (when 151 applicable), assuming an unstructured covariance matrix for the random effects. 152 Data linkage and analysis was carried out using SAS® 9.4 (SAS Institute Inc., Cary, NC, USA). 153 154 Results

155 The cohort

156 A total of 18306 participants were recruited into the GoDARTS cohort, of which 16700 met the selection 157 criteria. 9417 of patients had type 2 diabetes at recruitment and 7283 did not have diabetes at 158 recruitment nor developed it later and formed the control group. 1606 patients were excluded from the 159 current study, of which 681 had other types of diabetes, 720 developed diabetes after recruitment and 160 205 did not have SCr tests on or after recruitment (Figure 1). Table 3 shows baseline characteristics of 161 the cohort. People within type 2 diabetes were older than controls (66.9 vs 60.8 years old, difference 6.0 162 years, 95% CI 5.7-6.4) and 44.0% were females compared to 51.4% in the control group (difference 163 7.4%, 95% CI 5.8-8.9). People with T2D had a lower eGFR at baseline compared to controls (76.6 vs 84.3, 164 difference 7.7, 95% CI (7.1-8.29), 26.6% of people with T2D had CKD at recruitment compared to only 165 9.1% in the control group (difference 17.5%, 95% CI (16.3-18.6), and there was a higher percentage of 166 people with cardiovascular disease in the diabetic group compared to control (Table 3). The mean (SD) 167 follow-up time from recruitment was 8.2 (3.5) years for people with type 2 diabetes vs (2.4) for controls. 168

169 AKI and AKI episodes

170 Table 4 shows summary statistics of SCr measures from recruitment and describes the frequency of AKI 171 in the two groups. A total of 512615 SCr tests were recorded from recruitment; of those 387657 (75.6%) 172 were from patients with type 2 diabetes. The median (IQR) number of SCr measures per individual 173 during the follow-up were 31 (19-51) in type 2 diabetes vs 11 (4-21) in controls. Including post AKI 174 creatinines in order to calculate a baseline value increased the yield of AKI cases from 28306 to 40567. 175 A breakdown of AKI cases identified using the different baseline SCr definitions using pre and post Index 176 SCr measures is shown in Table S1 in the supplementary material. After grouping successive tests into 177 episodes, a total of 13928 AKI episodes were identified from recruitment until end of follow-up. Of these 178 11647 were experienced by patients with diabetes. AKI occurred in 5837 patients representing 48.6% 179 (N=4580) of patients with type 2 diabetes vs 17.2% (N=1257) of controls (difference 31.4%, 95% CI 30.0-180 32.7). More than 50% of patients with diabetes experiencing AKI had recurrent AKI, whereas the 181 majority of patients in the control group with AKI had only one episode of AKI during follow-up (Table 5). 182 Overall 54.2% of AKI episodes lasted no more than two days, a further 26.5% between 2 to 7 days, and 183 the remaining 19.3% of AKI episodes were longer than 7 days resulting in AKD. Less than five AKI/AKD 184 episodes were greater than 90 days, however after inspection it was revealed that these occurred 185 during hospitalisation due to other complications. 76.3% of AKI episodes were stage 1 with the rest 186 being stage 2 or 3. Diabetes was significantly associated with increased AKI episode length (p-value 187 <0.001) but not significantly associated with AKI stage (p-value=0.737).

188

189 AKI and CKD

Figure S1 illustrates the complex interplay between AKI/AKD and CKD and the many trajectories evolving
during the course of the disease. The way AKI initiates and develops can take many forms ranging from
one acute kidney insult which improves rapidly with full recovery within seven days (figure a2) to one or

more acute kidney insults during the course of the disease which progress to AKD requiring more than seven days to resolve. There are also cases when serum creatinine does not fully reverse after an AKI episode leading to the development of CKD (Figure d2). This further shows that, while some patients fully recover following an episode of AKI and never develop CKD (Figure a1-a3), others may experience a rapid kidney decline following an AKI episode (figure d1-d3). At the same time there is also the possibility to develop CKD without prior AKI episodes, and only experience AKI later as superimposed on CKD (figures b1-b3, c1-c3).

200 Table 6 describes the characteristics of people with diabetes vs controls in terms of their sex, age and 201 follow-up time as well as frequency of AKI during follow-up depending on CKD status. 26.6% (N=2504) of 202 people with diabetes had CKD at recruitment and further 19.7% (n=1855) developed the condition 203 during follow-up leading to a total of 46.3% (n=4359) people with CKD in the diabetic group compared 204 to 17.1% (n=1251) in the control group (difference 29.2%, 95% CI 27.8-30.4). In people with diabetes 205 and CKD, 50.3% were female (n=2192) compared to only 38.6% (n=1954) in those without CKD 206 (difference 11.7%, 95% CI 9.7-13.7). Also, people with diabetes developed CKD at a younger age 207 compared to people in the control group (mean age 74.1 vs 77.6 years, difference 3.5 years, 95% Cl 3.0-208 4.0) 66.1% (n=2883) of people with diabetes who developed CKD experienced AKI superimposed on CKD 209 in the diabetic group compared to 45.5% (n=569) in the control group (difference=20.6%, 95% CI 17.5-210 23.8). Additionally, 26.6% (n=493) of people with diabetes who developed CKD after recruitment had at 211 least one episode of AKI prior to development of CKD, the corresponding figure in the control group 212 being 9.9% (n=58, difference 16.7%, 95% CI 13.3-19.8). The proportion of people experiencing AKI was 213 significantly higher in the diabetic group compared to the control group for those patients who did not 214 have CKD at recruitment nor develop it later; 31.7% (n=1602) vs 10.8% (n=651) in the control group 215 (difference 20.9%, 95% CI 19.4-22.4).

216

217 Estimating AKI episode rates in people with and without diabetes

218	Table 6 shows estimates of AKI episode incidence rates and rate ratios for people with diabetes vs
219	control un-adjusted and adjusted for sex and age at recruitment. Regardless of CKD status, adjusted AKI
220	rates were 4.7 times higher in people with diabetes compared to controls (adjusted rate 179.0 vs 38.4
221	per 1000 person-years, RR=4.7, 95% CI 4.3-5.0). In particular, people with diabetes and no CKD
222	experienced AKI at a rate almost five times higher than people with no diabetes (adjusted rate 121.5 vs
223	24.6 per 1000 person-year, RR=4.9, 95% CI 4.4-5.5), whereas in people with CKD rate of AKI for those in
224	the diabetic groups was twice higher than in the corresponding control (adjusted rate 384.8 vs 180.0 per
225	1000 person-year, RR=2.1, 95% CI 1.9-2.4). Similarly, people with diabetes who develop CKD after
226	recruitment experience episodes of AKI at a rate twice higher than those in the control group (adjusted
227	rate 109.3 vs 47.4 per 1000 person-year, RR=2.3, 95% CI 1.8-3.0). It is noteworthy that the AKI rate in
228	people with diabetes in the absence of CKD was very close to AKI rate prior to development of CKD
229	(121.5 vs 109.0 per 1000 person year).
230	Additional model adjustment for other co-morbidities at baseline only partially reduced the association
231	between diabetes and AKI incidence rates (Table S4 in Supplementary material, RR=3.85, 95%CI 3.44-
232	4.32 in people with no CKD at recruitment or during follow-up, and RR=2.01 95%CI 1.82-2.22 in people
233	with CKD at recruitment or during follow-up time).
234	

235 Sensitivity analysis for the AKI rate analysis

The sensitivity analysis conducted to estimate rates for stage 2 and 3 AKIs show consistent results with the main analysis (Tables S2 in the supplementary material). The results show that people with diabetes and no CKD experience stage 2 and 3 AKIs at a rate that is over five time higher than people in the control group (adjusted mean rate 30.6 vs 5.5 per 1000 person-year, RR=5.5, 95% Cl 4.6-6.6) whereas in people with CKD rate of AKI for those in the diabetic groups was twice higher than in the corresponding

control group (adjusted mean rate 76.5 vs 38.9 per 1000 person-year, RR=2.1, 95% Cl 1.8-2.5). Similarly,
analysis of rates of AKIs lasting over 48hrs or AKIs during hospital admission show consistent results with
the main analysis (Table S3 and Table S4).

244

245 Estimating the effect of AKI on eGFR slope over time

246 Of the 16700 people included in the initial analysis, there were 3250 people with AKI prior to 247 recruitment which were not included in the eGFR analysis. A further 2558 people did not meet the 248 selection criteria of which 1324 had an AKI post recruitment (738 with T2D and 386 with no diabetes). 249 As a result a total of 10892 people with 279391 SCr measures were included in the eGFR longitudinal 250 data analysis. Of these 5665 had T2D and 5227 were from the control group (Figure 1, Tables S6 and S7 251 in the supplementary material). Of the 10892, 2470 people experienced an AKI during follow-up of 252 which 1859 had T2D and 611 had no diabetes. People with no CKD at recruitment had a significant 253 higher decline in eGFR in the period pre-AKI compared to no-AKI regardless of diabetes status, but rate 254 of decline was significantly higher in people with diabetes (Figure 2, Table S6 in the supplementary 255 material: eGFR slope pre-AKI vs no-AKI =-1.14, 95% CI (-1.24 to -1.03) in people with T2D and -0.29, 256 95%CI (-0.45 to -0.11) in controls, slope difference=-0.85, 95%CI (-1.05 to -0.65)). A further decrease in 257 rate was observed in the control group in the period post-AKI compared to pre-AKI in both T2D and 258 control groups, the increase in rate of decline was only marginally significant in people with T2D (eGFR 259 slope post-AKI vs pre-AKI =-0.29, 95% CI (-0.59 to 0.01)), whereas it was significant in the control group 260 (eGFR slope post-AKI vs pre-AKI =-0.55, 95% CI (-1.08 to -0.03)), however the difference between T2D 261 group and control was not significant (slope difference=0.26, 95%CI (-0.34 to 0.86)). Sex was significantly 262 associated with eGFR with males having a higher mean eGFR than females in people with T2D and lower 263 in control. No change in eGFR slope was observed between male and females in any of the subgroups. 264 An increase in age was associated with a reduction in eGFR at baseline regardless of diabetes status, but

significant differences in eGFR slope among the different age groups were observed only in people with
 T2D. Furthermore, people with peripheral vascular disease and hypertension had a significant further
 decline in eGFR slope regardless of diabetes status.

268 People with CKD at recruitment show a higher rate decline in eGFR in the period pre-AKI compared to 269 no-AKI and this result was significant in the T2D group and marginally significant in the control group, 270 but the difference between the two groups was not significant (Figure 2, Table S7 in the supplementary 271 material: eGFR slope pre-AKI vs no AKI =-0.79, 95% CI (-1.05 to -0.52) in people with T2D and -0.40, 272 95%CI (-0.85 to 0.05) in controls, slope difference=-0.38 (-0.90 to 0.14)). The decline in eGFR rate post 273 AKI compared to pre-AKI did not change in people with T2D diabetes (eGFR slope post AKI vs pre 274 AKI=0.23, 95% CI (-0.24 to 0.71)), whereas AKI was associated with further eGFR decline post AKI 275 compared to pre AKI period in controls (eGFR slope post AKI vs pre AKI =-0.84, 95% CI (-1.73 to 0.06)), 276 with the post-AKI effect being significantly different between T2D and control groups (slope 277 difference=1.07, 95%CI (0.06 to 2.08)). There was no significant eGFR difference between males and 278 females in people with CKD at recruitment regardless of diabetes status. An increase in age was 279 associated with a reduction in eGFR at baseline regardless of diabetes status, and older people with 280 diabetes appeared to have a lower eGFR decline than younger ones. None of the cardiovascular diseases 281 were significantly associated with eGFR at baseline or eGFR slope, however their effect was an 282 important one as reflected in the model AIC and therefore they were retained in the model. 283

284 Discussion

In our study, we have quantified rates of AKI in patients with and without diabetes demonstrating the
extent of the risk. Rates of AKI are significantly higher in patients with type 2 diabetes compared to
those without with a 4.7 fold increase in AKI rate. In people with diabetes and preserved renal function,
rate of AKI is 4.9 fold higher than people without diabetes whereas in people with CKD rate of AKI for

those in the diabetic group is 2 fold higher than in non-diabetics. More than 50% of the patients with
diabetes who develop AKI will suffer from recurrent events. Rates of CKD are also higher in patients with
Type 2 diabetes with 46.3% developing CKD compared to 17.1% in those without diabetes.
Fall in eGFR slope before AKI was steeper in people with diabetes compared to those without diabetes.

After AKI episodes, loss of eGFR became steeper in people without diabetes, but did not increase in
those with diabetes and pre-existing CKD.

295 In comparison to other studies, progressive decline leading to CKD has been well described in people with type 2 diabetes ¹, but AKI in diabetes mellitus have been less investigated ^{6,7,27,28}. Girman et al 296 297 examined 119 966 patients with diabetes and 1 794 516 patients without diabetes from the General 298 Practice Research Database. AKI incidence was markedly higher in their cohort: 198 per 100,000 299 person-years in patients with Type 2 diabetes compared with 27 per 100,000 patients-years among 300 patients without diabetes (crude hazard ratio 8.0, 95% CI 7.4-8.7)⁶. They did not utilise a biochemical 301 definition for AKI relying on clinical coding which can lead to significant under ascertainment ²⁹. In 302 addition, a meta-analysis by James et al showed that the hazards rations for AKI were higher in 303 participants with diabetes compared to those without diabetes at any level of eGFR ³⁰. Once again, the 304 definition for AKI relied on administrative codes in these studies thereby under estimating milder forms 305 of AKI. There are very few studies that have examined AKI and CKD simultaneously and recurrent AKI in this group of patients ³¹. 306

Our results are consistent with existing evidence indicating that diabetes is an independent risk factor for AKI ^{3,6,27}. However, reported AKI rates in people with diabetes vary greatly depending on the population studied (e.g. different specialist settings, age range) and the methods used for AKI identification (e.g. medical history, ICD10 coding or changes in SCr) ¹⁰. Most of the prior studies have reported AKI incidence of new AKI cases within a given time window and therefore estimates relate to number of patients experiencing AKI. The algorithm developed in this study allows quantification of AKI

313 rates based on number of AKI episodes including recurrent AKI. Our findings have important clinical 314 implications. AKI is associated with adverse patient outcomes including increased mortality, future development of CKD and increased length of hospital stays ^{3,32,33}. It therefore places a significant 315 financial burden on healthcare resources ³⁴. In our study over 75% of AKI were Stage 1 reflecting a mild, 316 317 transient increase in serum creatinine. This may be of clinical significance as there is an increasing 318 evidence showing that even mild, transient (lasting less than 24 hours) AKI is associated with poorer 319 long term outcomes ^{35,36} compared to those who do not have AKI. There are currently no effective 320 treatments for AKI once it is established and so earlier detection and prevention is vital. It is, however, important to note that there may be misclassification of chronic decline in renal function in patients 321 322 with diabetes accounting for some of the observed increased rates of AKI. We have shown that rates of 323 AKI are higher in patients with diabetes both with and without CKD with more than half developing 324 recurrent episodes. To our knowledge, there has been no previous work looking at eGFR slopes prior to 325 developing AKI. We found that those who develop AKI with diabetes have a greater decline in eGFR 326 slope prior to developing AKI than those who do not. These findings are expected as a declining kidney 327 would be more susceptible to episodes of AKI. However, it is surprising that there is less additional 328 decline in eGFR in those with diabetes compared to those without following an episode of AKI compared 329 to prior to an AKI episode. It remains unclear what the mechanism underlying AKI is in diabetic patients. 330 A predisposing factor in these patients may be generalised or intrarenal atherosclerosis. In addition, 331 patients with diabetes are likely to have glomerular hyperfiltration which is masking structural renal 332 damage thereby rendering them more susceptible to AKI than those without diabetes due to their 333 reduced repair capacity and so are susceptible to fluctuations in serum creatinine. A further suggested 334 mechanism is that tubular growth in response to hypergylcemia promotes inflammation, senescence, 335 and tubulointerstitial fibrosis which enhance the susceptibility of the diabetic kidney to episodes of AKI 336 ³⁷. It also remains unclear whether prevention of AKI in these patients would prevent or delay

progression of CKD. However, it would seem sensible that these patients are more closely monitoring during intercurrent illnesses with a greater awareness of avoiding high risk medicines such as nonsteroidal anti-inflammatories and aminoglycosides. There is currently a lack of awareness among patients with diabetes of the risk of AKI and so patient education on the importance of hydration may play an important in role. We have also shown that in patients with both hypertension and diabetes, there is an additional decline in eGFR highlighting the importance of blood pressure control in addition to ensuring good glycaemic control in this patient group.

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345 An important strength of the study is the refinement of KDIGO definition enabling a more sensitive 346 estimation of AKI rates which has allowed us to demonstrate the high risk of AKI in patients with 347 diabetes regardless of CKD status. We developed an algorithm to identify AKI episodes from SCr 348 measures. A number of definitions to detect AKI cases based on changes in SCr have been used 349 previously ¹⁶, and NHS England has implemented an algorithm which applies the KDIGO definition to 350 routinely collected SCr tests to automatically produce AKI alerts to support clinical investigations ¹⁵. This 351 algorithm, defines baseline creatinine levels based on SCr one year prior to the index date, potentially 352 leading to undetected AKI when such measurements are not available. The proposed algorithm utilises 353 SCr values both prior and after the index date. Whilst this may not be useful for AKI detection in clinical 354 practice, it may improve AKI detection for epidemiological purpose when applying to routinely collected 355 datasets allowing for a more sensitive estimation of AKI incidence. Our study shows that at least one 356 third of AKI cases remains undetected when baseline creatinine is only based on tests prior to the index 357 date. Previous epidemiological studies of AKI from routinely collected SCr reported AKI cases in isolation, with episodes being defined using either fixed time periods such as 30 days ¹⁴ or admission and 358 discharge dates for hospitalized patients ³¹. The current study is novel through the development of an 359 360 algorithm which examines consecutive SCr measures to detect the start and the end of an AKI episode,

361 which can be used to calculate the length of the episode and further assess whether the AKI has 362 resolved quickly or it has progressed to AKD. The grouping of AKI cases into AKI episodes was 363 particularly important to allow an accurate estimate of AKI rates when applied to routinely collected 364 data. Identification of the AKI episode start and end dates was also used in the study to clean the SCr 365 data in order to allow assessment of CKD status and correctly determine the CKD onset date, which 366 represents another strength of the study. Another important strength of the study is the development 367 of a statistical framework for the analysis of the eGFR longitudinal data to evaluate decline in eGFR 368 before and after an AKI event depending on diabetes and CKD status. One of the main limitations of the study relates to the nature of routine healthcare data where blood 369 370 measurements are infrequent, which makes it difficult to calculate baseline creatinine for assessment of 371 AKI. As a result some of the AKI in the longitudinal data might remain undetected leading to 372 misclassification between AKI and progressive CKD. This variation in the intensity of blood sampling may 373 also lead to ascertainment bias in AKI estimation due to more tests that are being performed in sicker 374 patients. In our study, blood tests were performed on average three times more often in people with 375 diabetes than people in the control group. This may partially explain the high AKI rate in people with 376 diabetes compared to controls. It could however be argued that increased testing was performed in 377 response to clinical indication and similarly lack of testing in those who were deemed well. The 378 possibility that the increased AKI is being driven by the increased testing rather than the other way 379 round is diminished by the analysis of more severe (stage 2 and stage 3) AKIs and AKIs lasting more than 380 48hrs, for which a high AKI rate ratio between people with diabetes compared to controls in the 381 absence of CKD were obtained. In addition, diabetes status confers a substantially increased risk for AKI 382 in individuals with pre-existing CKD, where the testing rate is high regardless of diabetes status. These 383 results demonstrates a profoundly increased clinical burden of acute kidney disease in diabetes patients. 384 Another limitation of the study is the potential of selection bias due to the use of consented data from

385 the GoDARTs cohort, a characteristic of most observational studies using consented data, which may 386 lead to AKI rate estimates that are not generalizable. Furthermore calculation of slopes required a 387 number of creatinine measures over a specified time period and so a significant number of patients 388 were excluded from the analysis. This could introduce selection bias which may have affected our 389 findings. However, it is difficult to eliminate this issue when examining eGFR slopes using observational 390 data. 391 In conclusion, we have quantified the risk of AKI in patients with diabetes and its relationship as both a 392 precursor and a consequence of CKD. The risk of AKI in this population of patients is currently 393 underestimated and associated adverse outcomes following AKI are not well understood. Further work 394 to evaluate the pathogenesis for AKI and the risk factors associated with the increased AKI rate in 395 patients with diabetes such as use of medication is required to allow for development and 396 implementation of interventions which both prevent the occurrence of AKI and reduce decline in eGFR 397 thereby improving patient outcomes. 398 399 Acknowledgments

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410

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412 The authors have nothing to disclose

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421 Ethics approval and consent to participate

422 The GoDARTS study was approved by the Tayside Medical Ethics Committee with informed consent

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428

429 Author Contributions

430 S.H. design the study, conducted the data processing and analysis and wrote and revised the

431 manuscript. M.K.S., R.S.Y.K., S.M., A.S.F.D., and E.R.P., contributed to the interpretation of the data and

432	the revision of the manuscript. S.B. and C.N.A.P. contributed to study design, the interpretation of the
433	data and the writing and revision of the manuscript. S.H and C.N.A.P. are guarantors of this work and, as
434	such, had full access to all the data in the study and take responsibility for the integrity of the data and
435	the accuracy of the data analysis.
436	

438 Supplemental Table of Contents

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- 453 Figure S1: Identifying AKI episodes (red circles) from longitudinal serum creatinine test (a1-d1). The
- different trajectories of AKI episodes (red) during the course of the disease (a2-d2) ranging from rapid
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- 457 longitudinal data to ascertain CKD status (a3-d3): first AKI flagged eGFR values are removed (red circles),

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- then median eGFR is calculated based on the remaining eGFR values (black line), which is used to
- 459 determine the date of CKD onset.
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Table 1. Definition of AKI cases, AKI episodes and AKI stages

Definition of AKI cases using the NHS algorithm (one of the three criteria) (19):						
Criterion 1:	Serum creatinine \ge 1.5 times higher than median of all creatinine measures in the 8-365 days prior to index.					
Criterion 2: index.	Serum creatinine \ge 1.5 times higher than the lowest creatinine in the 7 days prior to					
Criterion 3:	Serum creatinine > 26 $\mu mol/L$ higher than the lowest creatinine in the 48 hours prior to index.					
Definition of A	KI cases using the modified algorithm (one of the four criteria):					
Criterion 1:	Serum creatinine \ge 1.5 times higher than median of all creatinine measures in the 8-365 days prior to index					
Criterion 2:	Serum creatinine \ge 1.5 times higher than the lowest creatinine in the 7 days prior to or post index					
Criterion 3:	Serum creatinine > 26μ mol/L higher than the lowest creatinine in the 2 days prior to or post index					
Criterion 4:	Serum creatinine \ge 1.5 times higher than median of all creatinine measures in the 8-365 days post index					
Definition of A	KI episode: grouping AKI cases into episodes					
Step 1:	Serum creatinine tests measured within 7 days were grouped into episodes of care					
Step2:	If any value within an episode of care was flagged as AKI then the whole episode was flagged as AKI					
Step3:	Within each episode of care Serum creatinine values before and after an AKI case that were greater that 1.2 fold increase in baseline were included in the AKI episodes and used to determine the start and the end of the AKI episode					
Step 4	AKI episodes occurring within 7 days further linked to assess AKD					
Step 5:	AKD of length greater than 90 days flagged as potential CKD					
Classification c	riteria for AKI stages (19):					
Stage 1	Rise in creatinine > 26 μ mol/L within 48 h (2 days) or 1.5 \leq index/baseline<2					
Stage 2	2 ≤ index/baseline < 3					
Stage 3	index/baseline ≥ 3					

568 Table 2. Establishing CKD date and CKD status from the longitudinal eGFR data.

Implementat	tion of the median smoother to the eGFR data to ascertain CKD
Step 1:	eGFR values contained within AKI episodes were removed from the data
Step 2:	Calculate the median eGFR for the 91 to 365 days prior to index: <i>Median</i> 91-365d prior.
Step 3:	Calculate the median eGFR for the 7 days prior to 7 days post index: <i>Median</i> _{7d prior-7d po}
Step 4:	Calculate the median eGFR for the 91 to 365 days post index: <i>Median</i> 91-265d post.
Sept 5:	Define <i>Median</i> _{eGFR} as the median of the three medians defined in steps 2-4
Step 6:	CKD date establish when at least two of the medians in steps 2-4 are available and les than 60 ml/min per 1.73m ²
Definition o	f CKD status
No CKD	At recruitment or during follow-up.
Pre-CKD	The period from recruitment until development of CKD, for those people that developed CKD later.
Post-CKD	The period after recruitment, for those that had CKD at recruitment, or post CKD, for those that developed the condition later, until end of follow-up.

584 Figure 1: Flow chart of patient selection in the current study



Figure 2: Visual representation of the eGFR slope estimates in people without AKI (No AKI), prior to the AKI (Pre AKI) and after the AKI event
 (Post AKI) depending on diabetes status and CKD status at recruitment (^aReference group includes: No AKI during follow-up, female, 49 and
 below, no cardiovascular disease; ^bReference group includes: No AKI during follow-up, female, 50 to 64, no cardiovascular disease)



	All patients (N=16700)	T2DM (N=9417)	Control (N=7283)
Sex: Female N (%)	7888 (47.2)	4146 (44.0)	3742 (51.4%)
Age at recruitment: mean (SD)	64.3 (12.5)	66.9 (11.3)	60.8 (13.3)
eGFR at recruitment: mean (SD) ^a	79.9 (19.7)	76.6 (21.0)	84.3 (16.9)
CKD at recruitment: N(%) ^b	3168(18.9)	2503 (26.6)	665 (9.1)
Cardiovascular disease at recruitment			
Coronary Artery Disease (CAD): N (%)	3271 (19.6)	2489 (26.4)	782 (10.7)
Congestive Heart Failure (CHF): N (%)	670 (4.0)	587 (6.2)	83 (1.1)
Peripheral Vascular Disease (PVD): N (%)	636 (3.8)	540 (5.7)	95 (1.3)
Cerebrovascular Disease (CD): N (%)	786 (4.7)	644 (6.8)	142 (1.9)
Hypertension: N (%)	9863 (59.1)	7271 (77.2)	2592 (35.6)

^aeGFR at recruitment was missing for 145 people

^b 2442 additional participants developed CKD during follow-up

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Table 3. Baseline characterises of the cohort broken down by diabetes status

	All patients (N=16700)	Type 2 diabetes (N=9417)	Control (N=7283)
Number of SCr tests	512615	387657	124958
Number of SCr tests per patient: median (IQR)	22 (10-39.5)	31 (19-51)	11 (4-21)
Number of SCr tests flagged as AKI			
Old algorithm (retrospective tests)	28306	24257	4049
Modified algorithm (retrospective and prospective tests)	40567	34469	6098
Number of AKI episodes	13928	11647	2281
Number of SCr tests within AKI episodes	65316	55401	9915
Number of patients with AKI during follow-up	5837	4580 (48.6)	1257 (17.2)
Number of episodes per person: median (IQR)	2 (1-3)	2 (1-3)	1 (1-2)
Length of AKI episode: median (IQR)	3 (1-7)	3 (1-7)	3 (1-6)
AKI episode ≤ 2days	7544 (54.2)	6237 (53.6)	1307 (57.3)
AKI episode> 2days and \leq 7days	3697 (26.5)	3114 (26.7)	583 (25.6)
AKI episode > 7days	2687 (19.3)	2296 (19.7)	392 (17.2)
AKI stages			
Stage 1	10633 (76.3)	8895 (76.4)	1738 (76.2)
Stage2	2285 (16.4)	1901 (16.3)	387 (16.8)
Stage 3	1010 (7.3)	851 (7.3)	159 (7.0)

Table 4. Descriptive statistics showing number of SCr tests from recruitment, number of SCr test flagged by AKI using the NHS England algorithm vs the modified algorithm, number of AKI episodes and number of patients experiencing AKI during the follow-up time as well as characteristics of the AKI episodes in terms of length and severity in the diabetic and control groups.

Patients' groups	Sex Female number (%)	Age at recruitment	Follow-up time (years) Mean (SD)	Number of SCr tests per patient per year: median (IQR)	AKI patients N (%)	Number of AKI episodes
All patients (N=16700)	7888 (47.2)	64.3 (12.5)	8.8 (3.2) [*]	2.6 (1.2-5.2)	5837(35.0)	13928
Control (N=7282)	3742 (51.4)	60.8 (13.3)	9.6 (2.4)	1.1 (0.4-2.3)	1257 (17.3)	2281
Type 2 diabetes (N=9417)	4146 (44.0)	66.9 (11.3)	8.2 (3.5)	3.8 (2.4-7.4)	4580 (48.6)	11647
No CKD (N=11090)	5089 (45.9)	59.9 (11.8)	9.2 (2.9)*	1.8 (0.8-3.2)	2263 (20.4)	3952
Control (N=6032)	3135 (52.0)	57.9 (12.1)	9.8 (2.3)	0.9 (0.4-1.7)	651(10.8)	951
Type 2 diabetes (N=5058)	1954 (38.6)	62.4 (11.0)	8.4 (3.4)	2.9 (2.0-4.7)	1602 (31.7)	3001
CKD (N=5610)	2799 (49.9)	72.8 (8.9)	8.1 (3.4)*	5.1 (3.0-9.4)	3584 (63.9)	9976
Control (N=1251)	607 (48.5)	75.3 (8.1)	8.7 (3.0)	3.3 (2.0-6.4)	606 (48.4)	1330
Type 2 diabetes (N=4359)	2192 (50.3)	72.1 (9.0)	8.0 (3.5)	5.7 (3.4-10.3)	2978 (68.3)	8646
Prior to CKD diagnosis (N=2442)	1114 (45.6)	69.2 (8.9)	4.42 (3.1) ⁺	2.9 (1.8-4.6)	571 (23.4)	942
Control (N=587)	273 (46.5)	72.7 (8.2)	4.7 (2.9)	1.8 (1.1-3.3)	58 (9.9)	120
Type 2 diabetes (N=1855)	841 (45.3)	68.2 (8.9)	4.3 (3.1)	3.2 (2.2-5.0)	493 (26.6)	822
Post CKD diagnosis (N=5610)	2799 (49.9)	74.9 (8.3) [§]	6.2 (3.5) [‡]	5.7 (3.2-11.1)	3352 (59.8)	9034
Control (N=1251)	607 (48.5)	77.6 (7.7)	6.5 (3.5)	4.0 (2.3-7.5)	569 (45.5)	1210
Type 2 diabetes (N=4359)	2192 (50.3)	74.1 (8.3)	6.1 (3.5)	6.3 (3.6-12.0)	2883 (66.1)	7824

*from recruitment until end of follow-up (RRT/death/out with HB/May 2017 whichever happened first).

⁺from recruitment until development of CKD.

[‡]from development of CKD/recruitment, whichever happened last, until end of follow-up.

[§]Age at recruitment or development of CKD, whichever happened last.

Table 5. Descriptive statistics showing sex, age, follow-up time and number of SCr tests as well as number of patients experiencing AKI and

number of AKI episodes in the diabetic vs control groups depending on CKD status.

	AKI episodes per 1000 person-years					
Patients' groups	Un-adjust	ed	Adjusted for age and sex			
	Mean rate (SE)	Rate ratio (95%CI)	Mean rate (SE)	Rate ratio (95% CI)		
All patients (N=16700)	131.6 (126.8-136.6)	-	114.8 (110.5-119.5)	-		
Control (N=7282)	38.2 (36.0-40.5)	1.0	38.4 (36.2-40.8)	1.0		
Type 2 diabetes (N=9417)	204.8 (196.4-213.6)	5.4 (5.0-5.8)	179.0 (171.5-186.9)	4.7 (4.3-5.0)		
No CKD (N=11090)	54.6 (51.4-58.0)	-	66.3 (61.1-72.1)	-		
Control (N=6032)	18.0 (16.6-19.6)	1.0	24.6 (22.3-27.2)	1.0		
Type 2 diabetes (N=5058)	101.1 (93.9-108.8)	5.6 (5.0-6.3)	121.5 (111.0-133.0)	4.9 (4.4-5.5)		
CKD (N=5610)	276.0 (265.1-187.3)	-	267.0 (252.1-282.8)	-		
Control (N=1251)	148.5 (135.8-162.3)	1.0	130.1 (117.7-143.8)	1.0		
Type 2 diabetes (N=4359)	312.6 (299.2-326.6)	2.1 (1.9-2.3)	299.3 (282.4-317.2)	2.3 (2.1-2.5)		
Prior to CKD diagnosis (N=2442)	93.8 (85.4-108.0)	-	92.9(81.0-106.1)	-		
Control (N=587)	45.8 (36.7-57.2)	1.0	47.4 (37.2-60.5)	1.0		
Type 2 diabetes (N=1855)	109.9 (99.3-121.6)	2.4 (1.9-3.1)	109.3 (94.8-126.1)	2.3 (1.8-3.0)		
Post CKD diagnosis (N=5610)	337.2 (323.3-351.7)	-	350.8 (321.8-382.5)	-		
Control (N=1251)	187.3 (170.5-205.8)	1.0	180.0 (159.1-203.8)	1.0		
Type 2 diabetes (N=4359)	379.2 (362.1-397.1)	2.0 (1.8-2.2)	384.8 (353.1-419.3)	2.1 (1.9-2.4)		

Table 6. AKI episode rates and rate ratios in the diabetic and non-diabetic groups depending on the CKD status