

# Appendix A

# Definitions and methodologies used in the 22nd Annual Report – data to the end of 2018

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## 1. The UK Renal Registry (UKRR) Annual Report

The UKRR was established by the Renal Association in 1995 with the primary aim of collating data centrally from all adult UK renal centres to improve the care of patients with end-stage kidney disease (ESKD). Children on renal replacement therapy (RRT) were initially captured by a separate registry established by the British Association for Paediatric Nephrology, but this activity passed over to the UKRR from 2009. The Renal Association has an active and involved patient council.

Although originally limited to patients on RRT – dialysis treatments and kidney transplant (Tx) recipients – the UKRR has started to collect all cases of acute kidney injury (AKI) in primary and secondary care (in England only) and some cases of chronic kidney disease (CKD) in secondary care not on dialysis. Collecting and reporting AKI and CKD data will in time allow the UKRR to report the journeys of patients who go on to start RRT, as well as those who choose conservative management instead of RRT.

The UKRR collects data to benchmark each of the UK's 71 adult and 13 paediatric renal centres against the Renal Association audit standards and publishes an annual report comprising centre comparisons, attainment of audit standards, national averages and long term trends for measures of renal care and patient outcomes.

The UKRR Annual Report focuses predominantly on patients with ESKD who are on RRT. Each chapter of the report analyses a subset of these patients as detailed in section 7.

#### 1.1 Groups of patients with kidney disease

#### 1.1.1 Patients with ESKD on RRT

Throughout this report the term ESKD is used to describe individuals with kidney disease who have progressed to such a point in their disease trajectory that they require either RRT or conservative management. The term ESKD is synonymous with established renal failure, end-stage renal failure and end-stage renal disease. The start date of ESKD is defined as the date of the first dialysis session or receipt of a pre-emptive kidney Tx.

#### 1.1.2 Patients with ESKD on conservative management

Through the addition of CKD data, the UKRR will in future report on patients with ESKD who do not start RRT. However, identifying patients receiving conservative management will depend on patients being coded as such on the treatment timeline.

#### 1.1.3 Patients with CKD

A preliminary analysis of the CKD dataset suggests only around 15 renal centres currently return any CKD data. Before CKD data are incorporated into analyses, further work is required to understand the nature of the data and how patients are defined as having CKD.

#### 1.1.4 Patients with AKI

The term AKI applies to individuals who experience a sudden decline in kidney function, which can be graded from mild to severe. The UKRR reports on the subset of adult patients with severe AKI receiving acute dialysis whose data are submitted to the UKRR through quarterly renal centre data returns, but there may be gaps in the dataset because of coding issues, or because some patients start in an intensive therapy unit (renal filtration) rather than in a renal centre. Some of these people may recover their kidney function and therefore not require any further RRT, some may die during this period, while others may become established on RRT and be classified as ESKD patients. Patients with AKI are not included in the current UKRR Annual Report – instead, see the AKI report.

## 2. Data flows to the UKRR and data completeness

#### 2.1 Data flows

Patient data flows to the UKRR from different sources, in different ways and with varying frequency, but primarily via quarterly electronic returns from renal centres. Data are collected without patient consent using section 251 permissions of the NHS Act (2006) as detailed in the Renal Association patient privacy notice. The current dataset (version 4.2) – the data variables which the UKRR has permission to collect – is available here. In reality, many variables are currently not well reported to the UKRR and an exercise is being undertaken to improve transparency and completeness of data collection – see the data completeness portal.

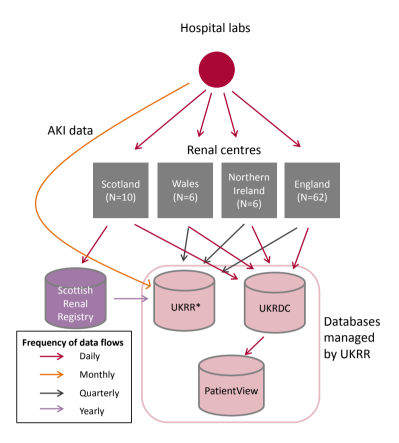
The UK Renal Data Collaboration (UKRDC) is an ongoing development which will allow data to flow daily from renal centres to the UKRR. Some renal centres have started sending their full data submission to the UKRDC and in future, quarterly returns for all renal centres will be replaced by near real time reporting across all centres. The UKRDC is also being used for those patients signed up to PatientView, a mobile-friendly platform allowing patients live access to much of the information in their health record.

#### 2.1.1 CKD and RRT data

The main source of data is the UK's 84 renal centres (adult and paediatric) which mostly send automatic downloads to the UKRR at the end of each quarter (figure A1), although not all centres routinely manage this. English, Welsh and Northern Irish renal centres send their data directly to the UKRR, where data are cleaned and validated prior to analysis. Scottish data are collected, validated and published by the Scottish Renal Registry before they are shared with the UKRR. The data items collected from renal centres are detailed below.

#### 2.1.2 AKI data

Hospital laboratories in England send AKI data to the UKRR on a monthly basis forming the AKI master patient index – an NHS England safety initiative (figure A1). The aim is to collect all episodes of AKI in England. The blood results of a patient with AKI continue to be reported to the UKRR for a further 15 months after the AKI episode to monitor potential renal recovery. The data items in the AKI data flow are limited to demographic and location data rather than the extensive data collected as specified in the CKD and ESKD 4.2 dataset. AKI data are also collected under section 251 permissions. The AKI data are not currently used in the UKRR Annual Report and are instead subject to separate analysis, reporting and publication.



**Figure A1** Frequencies and directions of patient data flows between hospital labs, renal centres and databases \*The UKRR database includes the British Association for Paediatric Nephrology database.

UKRDC – UK Renal Data Collaboration

#### 2.1.3 Other regular sources of data for the UKRR Annual Report

*NHS Blood and Transplant (NHSBT)* – the UKRR and NHSBT share a dataset on patients who are wait-listed for or who have received a kidney Tx.

*Public Health England (PHE)* – PHE send the UKRR a dataset on patients on dialysis who have had specific types of blood stream or gut infection in a 12 month period.

NHS Digital Hospital Episode Statistics (HES) for England and the Patient Episode Database for Wales (PEDW) – these datasets include information on patient comorbidities, hospital admissions and lengths of stay, surgical procedures and causes of death. These linkages have the potential to enhance UKRR data in a number of ways, by:

- enabling adjustment for case-mix in centre survival comparisons
- providing information about differences in rates of hospital admission between renal centres
- making it possible to study equity of access to other non-renal services, such as cardiology, stroke and orthopaedics.

#### 2.2 Data completeness

Unless otherwise stated, the data completeness threshold for a data item is  $\geq$ 70%, i.e. where a renal centre's data completeness for a data item falls below 70%, the individual centre will be excluded from an analysis, but the national total will include the centre's available data. Centres providing relevant data from <10 patients are also excluded from funnel and caterpillar plots for biochemistry and dialysis access analyses. While poor completeness may reflect a failure to accurately record patient data, other contributing factors include the incompatibility of local renal information technology (IT) systems and the loss of data during the transfer and validation processes on account of coding issues. Data completeness is likely to improve with the development of the UKRDC and increasing uptake of the UKRR dataset 4.2. The dataset has evolved and expanded over time in response to audit guidelines, with an understandable variable lag in the ability of local renal IT systems to respond to those changes.

#### 2.2.1 RRT data

Completeness of data items for patients receiving RRT varies between renal centres as detailed within each chapter.

#### 2.2.2 CKD data

So far only a small number of renal centres are returning CKD data as part of their quarterly extract.

#### 2.2.3 AKI data

Currently most hospital laboratories in England submit AKI data to the UKRR, although not all manage this every month. Until 100% of laboratories consistently report AKI data, some caution will be needed in their interpretation. The UKRR reports AKI data in quarterly Clinical Commissioning Group (CCG) reports (thinkkidneys.nhs.uk/aki/aki-data).

#### 2.2.4 Comorbidity data

Comorbidity data derived from diagnostic and procedure codes in HES and PEDW are used to augment comorbidity data for adults submitted by renal centres to the UKRR. A corresponding analysis of paediatric patients will be published later this year. Where UKRR comorbidities are absent (meaning the patient does not have the comorbid condition), but the comorbidity in HES/PEDW is present (meaning the patient has the comorbid condition), the UKRR 'absent' comorbidity is overwritten with the HES/PEDW 'present' comorbidity. Enriching the 2018 dataset with comorbidities from HES/PEDW increased comorbidity completeness from 66% to 99% and all renal centres in England and Wales had  $\geq$ 85% comorbidity completeness.

## 3. How the UKRR looks after patient data

#### 3.1 Information governance

The UKRR continues to receive support under section 251 of the NHS Act (2006) to collect data without individual patient consent. This ensures the robustness and validity of analyses. The fair processing of patient data remains a key principle of the General Data Protection Regulation (2016) and the Data Protection Act (2018). This requires organisations to be clear and open with individuals about how their information is used. The UKRR publishes this information on the UKRR website as well as in patient information leaflets and posters, which are distributed to all renal centres. Each year the UKRR completes NHS Digital's Data Security and Protection Toolkit. Further information on information governance is available on the Renal Association website.

#### 3.2 Small numbers

From time to time, due to the rarity of a condition or other factors, data for only a small number of patients (<5) will be available for analysis and inclusion in the UKRR's annual report. With so few patients the risk of reidentification is increased. To assess this risk, the UKRR conducts an assessment on each chapter of the annual report, identifying the level of risk of re-identification for each cell containing a small number and balancing this with the benefits of publication. Where the risk of re-identification is deemed too high, or the benefits of publication fail to outweigh that risk then the cell is supressed. Where small numbers are included in this report, it was deemed that the risk of re-identification was low, because no one cell can provide insight into an individual patient, unless that patient is already known to the reader.

# 4. How the UKRR codes and organises data prior to analysis or categorisation

The data collected by the UKRR are organised onto a chronological timeline of events and treatments for each patient. Some key dates are detailed below. For patients receiving haemodialysis (HD), the treatment element of the timeline can be validated against data supplied each time the patient has a dialysis treatment – this is termed 'session data'. UKRR data managers check timeline entries and liaise with renal centres to identify discrepancies within timelines, and between timeline and session data.

#### 4.1 Key dates – the renal 'treatment timeline'

#### 4.1.1 Date first seen by a nephrologist

For England, Wales and Northern Ireland, this is the date the patient first attended clinic or was an inpatient under the care of a nephrologist (whichever is the earlier). If a patient transfers into a renal centre from another renal centre then this date should be left blank by the new renal centre. For the purposes of this report, referral date is defined as the same as date first seen by a nephrologist.

The Scottish Renal Registry has provided date of referral to nephrology by general practitioner (GP) for people starting RRT in adult renal centres. This clearly differs from date first seen by a nephrologist because of the delay between a GP referral letter being issued and the actual appointment with the nephrologist.

#### 4.1.2 Late presentation

First seen date and date of RRT start (see below) are used to define late presentation, with a 90 day cut-off differentiating early versus late presentation. Scottish centres are included in some of the analyses on time of presentation, acknowledging the difference in definition described in 4.1.1 and the consequent underestimation in late presentation compared to the rest of the UK. Centre and national level data for Scotland are shown but UK results are not calculated. Two year cohorts may be used for analyses to make the late presentation percentages more reliably estimated and to allow these to be shown for subgroups of patients. Only data from those centres with  $\geq$ 70% completeness for the relevant year are used. This data item is investigated with centres, and possibly excluded, if an unexplained large proportion of patients are reported to have started RRT on the same date as the first presentation, because this is likely due to incorrect recording of data.

#### 4.1.3 Date of RRT start

A patient with ESKD starting RRT on 'chronic' HD (or PD or pre-emptive Tx) should be entered as such on the UKRR timeline on the date of the first HD (or PD) episode.

If a patient starts RRT with an episode of AKI in which it was felt that kidney function had potential to recover, then 'acute' HD (or acute HD or renal filtration) or acute PD (where appropriate) should be entered on the UKRR timeline. If subsequently it is felt that kidney function is no longer likely to recover, a timeline modality should be added of 'chronic dialysis' at the time when this becomes apparent (accepting that the timing of this change will vary by clinician practice and interpretation). The UKRR will interrogate the timeline of patients starting 'chronic' RRT and if there is evidence of recent 'acute' RRT, will backdate the date of start of RRT to the first episode of 'acute' RRT, provided there has been <90 days recovery of kidney function between acute and chronic episodes.

If a patient was started on dialysis and dialysis was temporarily stopped for <90 days for any reason (including access failure and awaiting the formation of further access), the date of start of RRT in UKRR analyses remains the date of first dialysis. If a patient recovers for  $\ge90$  days, subsequent RRT start dates are used.

The date of start of PD is defined as the date of first PD fluid exchange given with the intention of causing solute or fluid clearance. This is in contrast with a flush solely for confirming or maintaining PD catheter patency. In general, exchanges which are part of PD training should be considered as the start of PD (unless earlier exchanges have already been given). However, if it is not planned that the patient starts RRT until a later date, exchanges as part of PD training need not be considered the start of RRT.

#### 4.1.4 Change of modality date

Renal centres are requested to log in their timeline changes between PD and HD if the modality switch is for >30 days.

#### 4.1.5 Date of death

See section 8.1.

#### 4.2 Allocation of patients to a renal centre

The default method for allocating a patient to a renal centre is based on the centre sending their quarterly data.

Where applicable, pre-emptive Tx patients are allocated to their work-up centre rather than their Tx centre. This is not possible for all patients because some centres do not supply the 'transfer out for pre-emptive Tx' timeline code. Consequently, some patients remain allocated to their transplanting centre.

More generally, there are centre-specific variations in the repatriation of Tx recipients. Some Tx centres continue to follow-up and report on all patients they transplant, whereas others refer patients back to non-transplanting centres at some point post-Tx. Some Tx centres only refer back patients when their graft is failing. The time post-transplantation that a patient is referred back to their local centre varies between Tx centres, but the UKRR can detect patients being reported from both Tx and referring centres and in such situations care is usually attributed to the referring centre (see section 7.2).

# 5. Variables used to categorise patients in the UKRR Annual Report

#### 5.1 Demographics

#### 5.1.1 Location

This includes renal centre, region, country and CCG.

#### 5.1.2 Sex

Patients are defined as male or female as reported by the renal centre.

#### 5.1.3 Age

Age-adjusted analyses allow comparisons between centres with differing age distributions by adjusting the analysis as if all the patients were the same chosen age.

#### 5.1.4 Biometrics

Height, weight, body mass index (BMI) – these variables are only used for paediatric analyses. Data for height, weight, BMI and systolic blood pressure (SBP) vary with age, sex and size in children under 16 years and are therefore presented as z-scores as described in the relevant chapter. See section 7.7 for definitions.

#### 5.1.5 Ethnicity

Most centres electronically upload ethnicity coding to their renal IT system from the hospital patient administration system (PAS). Ethnicity coding in PAS is based on self-reported ethnicity. For the remaining centres, ethnicity coding is performed by clinical staff and recorded directly into the renal IT system (using a variety of coding systems). The details of regrouping the PAS codes into these ethnic categories are detailed below.

Tables A1 and A2 show the old and new groupings of ethnicity information used in this report as centres transition to the new codes. Ethnic categories are condensed into five groups (White, South Asian, Black, Chinese and Other). For all current analyses Chinese are grouped into Other.

**Table A1** Old ethnicity groupings

Code	Ethnic category	Assigned group
9S1	White	White
9SA9.	Irish (NMO)	White
SAA.	Greek Cypriot (NMO)	White
SAB.	Turkish Cypriot (NMO)	White
SAC.	Other European (NMO)	White
S6	Indian	South Asian
S7	Pakistani	South Asian
S8	Bangladeshi	South Asian
SA6.	East African Asian	South Asian
SA7.	Indian Subcontinent	South Asian
SA8.	Other Asian	South Asian
S2	Black Caribbean	Black
S3	Black African	Black
S4	Black/Other/NMO	Black
S41.	Black British	Black
S42.	Black Caribbean	Black

Table A1 Continued

Code	Ethnic category	Assigned group
9S43.	Black North African	Black
9S44.	Black other African country	Black
9S45.	Black East African Asian	Black
9S46.	Black Indian subcontinent	Black
9S47.	Black Other Asian	Black
9S48.	Black Black Other	Black
9S5	Black other/mixed	Black
9S51.	Other Black - Black/White origin	Black
9S52.	Other Black - Black/Asian origin	Black
9S9	Chinese	Chinese
9T1C.	Chinese	Chinese
9SA	Other ethnic non-mixed (NMO)	Other
9SA1.	British ethnic minority specified (NMO)	Other
9SA2.	British ethnic minority unspecified (NMO)	Other
9SA3.	Caribbean Island (NMO)	Other
9SA4.	North African Arab (NMO)	Other
9SA5.	Other African countries (NMO)	Other
9SAD.	Other ethnic NEC (NMO)	Other
9SB	Other ethnic/mixed origin	Other
9SB1.	Other ethnic/Black/White origin	Other
9SB2.	Other ethnic/Asian/White origin	Other
9SB3.	Other ethnic/mixed White origin	Other
9SB4.	Other ethnic/Other mixed origin	Other

NEC - not elsewhere contained; NMO - non-mixed origin

**Table A2** New ethnicity groupings

Code	Ethnic category	Assigned group
A	White – British	White
В	White - Irish	White
C	Other White background	White
D	Mixed - White and Black Caribbean	Other
E	Mixed - White and Black African	Other
F	Mixed - White and Asian	Other
G	Other Mixed background	Other
Н	Asian or Asian British – Indian	South Asian
J	Asian or Asian British – Pakistani	South Asian
K	Asian or Asian British – Bangladeshi	South Asian
L	Other Asian background	South Asian
M	Black Caribbean	Black
N	Black African	Black
P	Other Black background	Black
R	Chinese	Chinese
S	Other ethnic background	Other

#### 5.2 Health

#### 5.2.1 Primary renal disease (PRD)

Patients should be allocated a code for the PRD based on the histological or clinical picture, with codes available for where the cause is unknown. New PRD codes were produced by the European Renal Association – European Dialysis and Transplant Association (ERA-EDTA) in 2012. The data used for this report include a mixture of old and new ERA-EDTA codes. Old codes cannot be mapped to new codes, but the reverse mapping

is possible. Therefore, the old codes are used where available, and for those people without an old code, new codes (where available) are mapped back to old codes, using the mapping available on the ERA-EDTA website. As recommended in the notes for users in the ERA-EDTA's PRD code list document, the mapping of new to old codes is provided for guidance only and has not been validated. Therefore, care must be taken not to over interpret data from this mapping.

The old codes (both those received from centres and those mapped back from new codes) are then grouped into the same eight categories as in previous reports as shown in table A3.

**Table A3** Old primary renal disease (PRD) groupings

	Old PRD grouping	Assigned group
)	Chronic renal failure; aetiology uncertain unknown/unavailable	Uncertain aetiology
0	Glomerulonephritis; histologically NOT examined	Glomerulonephritis*
1	Focal segmental glomerulosclerosis with nephrotic syndrome in children	Glomerulonephritis
12	IgA nephropathy (proven by immunofluorescence, not code 76 and not 85)	Glomerulonephritis
13	Dense deposit disease; membrano-proliferative glomerulonephritis; type II (proven by immunofluorescence and/or electron microscopy)	Glomerulonephritis
.4	Membranous nephropathy	Glomerulonephritis
15	Membrano-proliferative glomerulonephritis; type I (proven by immunofluorescence and/or electron microscopy – not code 84 or 89)	Glomerulonephritis
16	Crescentic (extracapillary) glomerulonephritis (type I, II, III)	Glomerulonephritis
17	Focal segmental glomerulosclerosis with nephrotic syndrome in adults	Glomerulonephritis
9	Glomerulonephritis; histologically examined, not given above	Glomerulonephritis
20	Pyelonephritis – cause not specified	Pyelonephritis
21	Pyelonephritis associated with neurogenic bladder	Pyelonephritis
22	Pyelonephritis due to congenital obstructive uropathy with/without vesico-ureteric reflux	Pyelonephritis
23	Pyelonephritis due to acquired obstructive uropathy	Pyelonephritis
24	Pyelonephritis due to vesico-ureteric reflux without obstruction	Pyelonephritis
25	Pyelonephritis due to urolithiasis	Pyelonephritis
29	Pyelonephritis due to other cause	Pyelonephritis
30	Interstitial nephritis (not pyelonephritis) due to other cause, or unspecified (not mentioned above)	Other
1	Nephropathy (interstitial) due to analgesic drugs	Other
32	Nephropathy (interstitial) due to cis-platinum	Other
3	Nephropathy (interstitial) due to cyclosporin A	Other
34	Lead induced nephropathy (interstitial)	Other
39	Drug induced nephropathy (interstitial) not mentioned above	Other
10	Cystic kidney disease – type unspecified	Polycystic kidney
1	Polycystic kidneys; adult type (dominant)	Polycystic kidney
2	Polycystic kidneys; infantile (recessive)	Polycystic kidney
13	Medullary cystic disease; including nephronophthisis	Other
19	Cystic kidney disease – other specified type	Other
50	Hereditary/Familial nephropathy – type unspecified	Other
1	Hereditary nephritis with nerve deafness (Alport's syndrome)	Other
2	Cystinosis	Other
3	Primary oxalosis	Other
4	Fabry's disease	Other
9	Hereditary nephropathy – other specified type	Other
60	Renal hypoplasia (congenital) – type unspecified	Other
1	Oligomeganephronic hypoplasia	Other
3	Congenital renal dysplasia with or without urinary tract malformation	Other
6	Syndrome of agenesis of abdominal muscles (Prune Belly)	Other
70	Renal vascular disease – type unspecified	Renal vascular disease
71	Renal vascular disease – type unspectificular Renal vascular disease due to malignant hypertension	Hypertension
72	Renal vascular disease due to hypertension	Hypertension

**Table A3** Continued

Code	Old PRD grouping	Assigned group	
73	Renal vascular disease due to polyarteritis	Renal vascular disease	
74	Wegener's granulomatosis	Other	
75	Ischaemic renal disease/cholesterol embolism	Renal vascular disease	
76	Glomerulonephritis related to liver cirrhosis	Other	
78	Cryoglobulinemic glomerulonephritis	Other	
79	Renal vascular disease – due to other cause (not given above and not code 84-88)	Renal vascular disease	
80	Type 1 diabetes with diabetic nephropathy	Diabetes	
81	Type 2 diabetes with diabetic nephropathy	Diabetes	
82	Myelomatosis/light chain deposit disease	Other	
83	Amyloid	Other	
84	Lupus erythematosus	Other	
85	Henoch-Schoenlein purpura	Other	
86	Goodpasture's syndrome	Other	
87	Systemic sclerosis (scleroderma)	Other	
88	Haemolytic ureaemic syndrome (including Moschcowitz syndrome)	Other	
89	Multi-system disease – other (not mentioned above)	Other	
90	Tubular necrosis (irreversible) or cortical necrosis (different from 88)	Other	
91	Tuberculosis	Other	
92	Gout nephropathy (urate)	Other	
93	Nephrocalcinosis and hypercalcaemic nephropathy	Other	
94	Balkan nephropathy	Other	
95	Kidney tumour	Other	
96	Traumatic or surgical loss of kidney	Other	
98	Not known	Missing	
99	Other identified renal disorders	Other	
199	Code not sent	Missing	

<sup>\*</sup>Prior to the 15th UKRR Annual Report categorised as 'uncertain aetiology'.

#### 5.2.2 Cause of death

ERA-EDTA codes for cause of death are grouped as shown. Patients with a cause of death code 107 (advanced CKD not on dialysis) with no other information to determine the group were assigned to missing cause of death.

**Table A4** Cause of death groupings

Code	Cause of death grouping	Assigned group	
0	Cause of death uncertain/not determined	Uncertain aetiology	
11	Myocardial ischaemia and infarction	Cardiac disease	
12	Hyperkalaemia	Other	
13	Haemorrhagic pericarditis	Other	
14	Other causes of cardiac failure	Cardiac disease	
15	Cardiac arrest/sudden death; other cause or unknown	Cardiac disease	
16	Hypertensive cardiac failure	Cardiac disease	
17	Hypokalaemia	Other	
18	Fluid overload/pulmonary oedema	Cardiac disease	
21	Pulmonary embolus	Other	
22	Cerebro-vascular accident, other cause or unspecified	Cerebrovascular disease	
23	Gastro-intestinal haemorrhage (digestive)	Other	
24	Haemorrhage from graft site	Other	
25	Haemorrhage from vascular access or dialysis circuit	Other	
26	Haemorrhage from ruptured vascular aneurysm (not codes 22, 23)	Other	
27	Haemorrhage from surgery (not codes 23, 24, 26)	Other	
28	Other haemorrhage, (not codes 23–27)	Other	

IgA – immunoglobulin A

**Table A4** Continued

Code	Cause of death grouping	Assigned group	
29	Mesenteric infarction	Other	
31	Pulmonary infection bacterial (not code 73)	Infection	
32	Pulmonary infection (viral)	Infection	
33	Pulmonary infection (fungal or protozoal; parasitic)	Infection	
34	Infections elsewhere except viral hepatitis	Infection	
35	Septicaemia	Infection	
36	Tuberculosis (lung)	Infection	
37	Tuberculosis (elsewhere)	Infection	
38	Generalised viral infection	Infection	
39	Peritonitis (all causes except for PD)	Infection	
41	Liver disease due to hepatitis B virus	Other	
42	Liver disease due to other viral hepatitis	Other	
43	Liver disease due to drug toxicity	Other	
44	Cirrhosis – not viral (alcoholic or other cause)	Other	
45	Cystic liver disease	Other	
46	Liver failure – cause unknown	Other	
47	Patient refused further treatment for ESKD	Treatment withdrawal	
51	Patient refused further treatment for ESKD	Treatment withdrawal	
52	Suicide	Other	
53	ESKD treatment ceased for any other reason	Treatment withdrawal	
54	ESKD treatment withdrawn for medical reasons	Treatment withdrawal	
61	Uraemia caused by graft failure	Treatment withdrawal	
62	Pancreatitis	Other	
63	Bone marrow depression (aplasia)	Other	
64	Cachexia	Other	
66	Malignant disease in patient treated by immunosuppressive therapy	Malignancy	
67	Malignant disease: solid tumours (except code 66)	Malignancy	
68	Malignant disease: lymphoproliferative disorders (except code 66)	Malignancy	
69	Dementia	Other	
70	Peritonitis (sclerosing, with PD)	Other	
71	Perforation of peptic ulcer	Other	
72	Perforation of colon	Other	
73	Chronic obstructive pulmonary disease	Other	
81	Accident related to ESKD treatment (not code 25)	Other	
82	Accident unrelated to ESKD treatment	Other	
90	Uraemia caused by graft failure	Treatment withdrawal	
99	Other identified cause of death	Other	
100	Peritonitis (bacterial, with PD)	Infection	
101	Peritonitis (fungal, with PD)	Infection	
102	Peritonitis (due to other cause, with PD)	Infection	

#### 5.2.3 Infections

Patients on dialysis are susceptible to infections because of an impaired immune system and the need to regularly access the vascular system in HD or use of a catheter in PD. PHE carries out mandatory enhanced surveillance of methicillin-resistant *Staphylococcus aureus* (MRSA) bacteraemia, methicillin-sensitive *Staphylococcus aureus* (MSSA) bacteraemia, *Escherichia coli* bacteraemia and *Clostridium difficile* reporting for NHS acute trusts. A data sharing agreement exists between the UKRR and PHE to identify infections in dialysis patients in England in a given year through data linkage. In the 21st UKRR Annual Report, Wales provided data for the first time, which were extracted locally from the renal and hospital IT systems.

Until the 21st Annual Report, infection data were validated by securely emailing individual renal centres to confirm that infections were related to dialysis patients. Historically, this has resulted in only a small number of alterations in cases and was discontinued from the 21st report onwards.

PHE reports individual blood culture results. However, the annual report details individual infection episodes – repeated positive blood cultures within a two week timeframe are treated as a single infection episode for MSSA/MRSA/E. coli bacteraemia; beyond two weeks they are treated as a new episode or re-infection. Four weeks, rather than two weeks, is used as the cut-off for repeated *C. difficile* infections. Centre-specific rates for each infection are presented per 100 dialysis patient years. The denominator for this rate is calculated for each centre by summing the number of days that each dialysis patient contributes between 1 January and 31 December of the year in question. When calculating the modality specific rates, the number of days that every dialysis patient spends on each modality during the collection period is totalled.

To illustrate the variation in precision of the estimated infection rate, the rate of bacteraemia (MRSA and MSSA) per 100 dialysis patient years is plotted against the centre size in a funnel plot. This is plotted for each infection type.

#### 5.2.4 Comorbidity

The comorbidity data items collected in the UKRR dataset are listed below.

At the time of each patient starting RRT, clinical staff in each centre are responsible for recording, in yes/no format on their renal IT system, the presence or absence of the following comorbid conditions and information on current smoking status. Patients are classified as having complete comorbidity data if there is at least one entry (yes/no) for any one or more of the comorbid conditions, excluding smoking.

'Ischaemic heart disease' is defined as the presence of one or more of the following conditions: angina, myocardial infarction (MI) in the three months prior to starting RRT, MI more than three months prior to starting RRT or coronary artery bypass graft (CABG)/angioplasty.

'Peripheral vascular disease' is defined as the presence of one or more of the following conditions: claudication, ischaemic or neuropathic ulcers, non-coronary angioplasty, vascular graft, aneurysm or amputation for peripheral vascular disease.

'Non-coronary vascular disease' is defined as the presence of cerebrovascular disease or any of the data items that comprise 'peripheral vascular disease'.

Specific consideration needs to be made regarding diabetes coding. The UKRR also collects data on PRD and uses these data alongside the comorbidity data to determine which patients have diabetes mellitus. The comorbidity screen is intended to capture those patients who have diabetes only when it is not the PRD, but some clinicians enter 'yes' in the comorbidity field in such cases. Prior to statistical analyses, these fields are examined together to identify these cases and to ensure diabetes is only counted as either the PRD or a comorbid condition for a certain individual.

Several renal centres submit an expanded list of comorbidities (non-ST segment elevation MI; atrial fibrillation; transient ischemic attack; cerebrovascular event/stroke; peripheral vascular disease; and dementia) with associated dates as specified in the current dataset (version 4.2). Comorbidities at start of RRT are subsequently derived from the date of the comorbidity and the date of starting RRT.

Angina – history of chest pain on exercise with or without electrocardiogram (ECG) changes, exercise tolerance test, radionucleotide imaging or angiography.

*Previous MI within last three months* – detection of rise and/or fall of a biomarker (creatinine kinase [CK], CK-MB or troponin) with at least one value above the 99th percentile, together with evidence of myocardial ischaemia with at least one of either:

- ischaemic symptoms
- ECG changes indicative of new ischaemia (new ST-T changes or new left bundle branch block)
- development of pathological Q waves
- imaging evidence of new loss of viable myocardium or new regional wall motion abnormality.

Previous MI more than three months ago – any previous MI at least three months prior to start of RRT.

Previous CABG or coronary angioplasty.

Previous episode of heart failure – whether or not due to fluid overload.

Cerebrovascular disease – any history of strokes (whatever cause) and including transient ischaemic attacks caused by carotid disease.

Diabetes (not causing ESKD, i.e. not as the PRD) – this includes diet controlled diabetics.

*Chronic obstructive pulmonary disease (COPD)* – this is characterised by airflow obstruction. The airflow obstruction is usually progressive, not fully reversible and does not change markedly over several months:.

- airflow obstruction is defined as a reduced forced expiratory volume in one second (FEV1) and a reduced FEV1/FVC ratio (where FVC is forced vital capacity), such that FEV1 is <80% predicted and FEV1/FVC is <0.7
- the airflow obstruction is due to a combination of airway and parenchymal damage
- the damage is the result of chronic inflammation that differs from that seen in asthma and which is usually the result of tobacco smoke.

There is no single diagnostic test for COPD. Making a diagnosis relies on clinical judgement based on a combination of history (exertional breathlessness, chronic cough, regular sputum production, frequent winter 'bronchitis', wheeze), physical examination and confirmation of the presence of airflow obstruction using spirometry (source: British Thoracic Society guidelines).

*Liver disease* – persistent enzyme evidence of hepatic dysfunction or biopsy evidence or hepatitis B antigen or hepatitis C antigen (polymerase chain reaction) positive serology.

*Malignancy* – defined as any history of malignancy (even if curative) e.g. removal of melanoma, excludes basal cell carcinoma.

Claudication - current claudication based on a history, with or without Doppler or angiographic evidence.

*Ischaemic/neuropathic ulcers* – current presence of these ulcers.

Angioplasty, stenting, vascular graft (all non-coronary) – this category now includes vascular grafts (e.g. aortic bifurcation graft) and renal artery stents.

Amputation for peripheral vascular disease.

Smoking – current smoker or history of smoking within the last year.

#### 5.2.5 Hypo/hypertension

Hypertension is analysed for Tx and paediatric patients using the relevant targets described in the chapters. Hypotension during dialysis is not currently routinely analysed.

#### 5.2.6 Diabetic/non-diabetic

In general, where the UKRR report refers to diabetics it refers to patients with diabetes as a PRD, but excludes patients with diabetes as a comorbidity. Non-diabetics, by contrast, includes patients with diabetes as a comorbidity.

#### 5.3 Treatment

#### 5.3.1 Referral time and surgical assessment

Time of presentation, the time a patient first sees a nephrology specialist and referral time are interchangeable for the purposes of this report and late presentation is defined above. Surgical assessment is the time at which a patient is seen by a surgeon for assessment for dialysis access – either an arteriovenous fistula (AVF), arteriovenous graft (AVG) or PD catheter. As with late presentation, three months prior to RRT start is used as a measure of care.

#### 5.3.2 RRT modality

The RRT treatment modalities available are a Tx, home haemodialysis (HHD), in-centre haemodialysis (ICHD) and PD – these are defined in the relevant chapters of the report. Paediatric patients on ICHD or HHD are reported in a combined HD group.

#### 5.3.3 Dialysis access

AVF, AVG, central venous catheter (CVC) – non-tunnelled line (NTL) and tunnelled line (TL) – and catheter insertion technique for PD are defined in chapter 1.

#### 5.3.4 HD session frequency and length

For patients on ICHD, the length and frequency of HD sessions are described in chapter 4. Patients on HHD are reported in a separate chapter in this year's annual report.

#### 5.3.5 Tx type

Donor after brain death (DBD), donor after circulatory death (DCD) and living kidney donor (LKD) Tx are defined in chapter 3.

#### 5.3.6 Tx wait-listing

Pre-emptive Tx wait-listing is presented in chapter 1, while Tx wait-listing in the dialysis population is presented in chapter 2. Listing status before start of RRT for incident patients (analysis in chapter 1) or at end of year for the prevalent dialysis cohort (analysis in chapter 2) are obtained using NHSBT data regularly matched to the UKRR database.

#### 5.3.7 Laboratory data items

The UKRR does not currently collect data regarding different assay methods, mainly because a single dialysis centre may process samples in several different laboratories.

The UKRR dataset contains a number of laboratory variables, many of which are not currently returned by renal

centres. It is planned to expand this work as part of an ongoing data completeness exercise.

The collection methods and statistical analyses undertaken on the core laboratory data items of the annual report are as follows.

#### 5.3.7.1 Incident biochemical and haematology variables

For the analyses of biochemical variables for incident patients (with the exception of start estimated glomerular filtration rate [eGFR] – see below), those patients commencing RRT (HD/PD/Tx) are included. Measurements for variables taken from after starting dialysis, but still within the same quarter of RRT start are used. Therefore, depending on when in the quarter a patient starts RRT, the data could be from zero to 90 days later. Due to possible deficiencies with extract routines it is possible that a small number of the values extracted electronically may actually be from before the person started dialysis. This problem will not occur for Scottish data. Results are also shown with the cohort subdivided into early and late presenters (date first seen by a nephrologist ≥90 days and <90 days before starting dialysis, respectively). For these analyses only centres with at least 70% completeness of presentation time data are included.

*eGFR at RRT start* – eGFR is calculated from serum creatinine. The start eGFR is studied amongst patients with eGFR data within 14 days before the start of RRT. In line with the National Institute for Clinical Excellence advice and for consistency across the UKRR Annual Report, the Chronic Kidney Disease Epidemiology Collaboration (CKD-EPI) creatinine equation is used to calculate eGFR. In previous years, up to the 19th Annual Report, the Modification of Diet in Renal Disease equation was used. In light of this change, the UKRR advises caution in comparing eGFR results with previous UKRR annual reports.

A wide variety of creatinine assays are in use in clinical biochemistry laboratories in the UK and it is not possible to ensure that all measurements of creatinine concentration collected by the UKRR are harmonised.

For the purpose of the eGFR calculation, patients who have missing ethnicity data but a valid serum creatinine measurement are classed as White. The eGFR values are log transformed due to their skewed distribution and geometric means are calculated.

In children, eGFR is calculated using the updated 'bedside' Schwartz formula, using centre-specific individual correction factors submitted to the UKRR. For young adults (16–18 years old), the Full Age Spectrum (FAS) creatinine equation is used because of low completeness of height in young adults managed in adult centres.

#### 5.3.7.2 Prevalent biochemical and haematology variables

Haemoglobin (Hb) and ferritin – for the analyses of prevalent dialysis patients, those patients receiving dialysis on 31 December at the end of the analysis year are included if they have been on the same dialysis modality in the same centre for at least three months. To improve completeness, the last available measurement for each patient from the last two quarters of the year are used for Hb and from the last three quarters for ferritin.

Erythropoiesis stimulating agents (ESA) – ESA data from the last quarter of the year of analysis are used to define which patients are receiving ESAs, with the exception of Scottish data, for which the second quarter is used because the Scottish Renal Registry submit ESA data only for the second quarter of each year. Scotland is included in the ESA analysis for ICHD patients, but not PD patients, because Scotland does not submit ESA data for PD patients. Each individual is defined as being on an ESA if a drug type and/or a dose is present in the data. Centres reporting <70% of HD or PD patients being treated with ESAs, respectively, are considered to have incomplete data and are excluded from further analysis. The percentage of patients on ESAs is calculated from these data and incomplete data returns risk seriously impacting on any conclusions drawn.

For analyses of ESA dose, values are presented as weekly ESA dose. Doses of <150 IU/week (assumed to be

darbepoietin or methoxy polyethylene glycol–epoetin beta) are harmonised with ESA data by calculating a weekly dose and multiplying by 200. No adjustments are made with respect to route of administration. Patients who are not receiving ESAs are not included in analyses of dose (rather than being included with a dose of zero). Many centres provide data on ESA dose but not on ESA frequency. The ESA dose field is defined as the weekly dose and the dose is presumed to have been converted accordingly on submission to the UKRR, unless otherwise indicated from the centre. This may be an incorrect assumption for a number of patients, and this needs to be considered when interpreting the ESA information.

The ESA data are collected electronically from renal IT systems, but in contrast to laboratory linked variables the ESA data require manual data entry. The reliability depends upon the data source – whether the entry is linked to the prescription or whether the prescriptions are provided by the primary care physician. In the latter case, doses may not be as reliably updated because the link between data entry and prescription is indirect. It is worth noting that ESA data are the only medication that is reported by the UKRR, because of data completeness (iron replacement is also not included).

Quarterly values are extracted from the database for the last two quarters for calcium, phosphate, bicarbonate and potassium and the last three quarters for parathyroid hormone (PTH). Patients who do not have these data are excluded from the analyses.

*Calcium* – the adjusted calcium is calculated by adjusting for the binding of albumin to a proportion of the calcium in the blood depending on albumin levels. Not all centres return adjusted calcium. For centres providing adjusted calcium values, these data are analysed directly because it is these values on which clinical decisions within centres are based. For centres providing unadjusted calcium values, the formula provided by each centre (or, if this is not available, the standard formula in widespread use) is used to calculate adjusted calcium.

*PTH and phosphate* – these variables no longer have target ranges in the most recent adult Renal Association audit guidelines and are therefore not currently reported in the UKRR Annual Report for the adult dialysis population. However, they are reported in paediatric patients and at the national level for the adult transplant population.

*Bicarbonate* – the audit measures used for serum bicarbonate in the HD cohort and PD cohort differ as per the most recent guidelines. For children and young people aged <18 years, the paediatric reference range has been used (see section 7.7)

Potassium – centres are requested to send pre-dialysis potassium levels for HD patients, which like all biochemical samples should be collected from a short-gap session (i.e. a gap of one day between HD sessions rather than the longer two day gap). Outlying centres are contacted and if it is identified that post-dialysis potassium data have inadvertently been submitted, these centres are excluded from the analysis. However, post-dialysis samples may remain within the analysis for some centres. Future data extracts will aim to ensure that only pre-dialysis results are submitted.

*Urea reduction ratio* (*URR*), *session duration and frequency* – the prevalent adult ICHD patient population for a given year is analysed using URR data taken from the third quarter of the year, unless that data point is missing, in which case data from the second quarter are taken. The use of URR data from the third quarter is preferred over the fourth quarter due to better data completeness.

Since 2015, centres have been submitting quarterly HD sessional data as specified in version 4.2 of the UKRR dataset. These data are used to augment the quarterly data on the frequency and duration of dialysis sessions across all centres, for those centres with poor completeness on those two items.

Data from patients known to be receiving more than or less than thrice weekly HD are omitted from the analysis. Patients who have missing data for the number of dialysis sessions per week are assumed to be dialysing thrice weekly. However, because not all centres report frequency of HD, it is possible that data from a small number of patients receiving HD at a different frequency are included in the analyses. HHD patients are excluded from the analysis.

The URR is calculated as the percentage fall in urea during a dialysis session by taking a urea sample before and after the dialysis session. Post-dialysis blood samples should be collected either by the slow-flow method, the simplified stop-flow method, or the stop dialysate flow method. The method used should remain consistent within renal centres and should be reported to the UKRR.

# 6. Statistical methods and analyses used in the UKRR Annual Report

SAS software (sas.com) is used for all analyses.

#### 6.1 Estimation of renal centre dialysis catchment populations

Estimates of each renal centre's catchment area are needed to calculate estimates of the incidence and prevalence rates of RRT at renal centre level. The UKRR database of the incident dialysis population between 1 January 2008 and 31 December 2012 was used to estimate the size of each UK renal centre's catchment area. This used the postcode and renal centre for each patient at the time of starting RRT on dialysis.

Polygons were constructed to define an area around the geographical location of each dialysis patient. The lines of the polygons, representing the boundaries between areas, were drawn such that they were equidistant between adjacent patients, creating a map of non-overlapping polygons covering the entire area of England, Northern Ireland, Scotland and Wales (the process was done separately for each country). This method produces Thiessen polygons which have the property that all locations within each polygon share the same nearest dialysis patient (Boots BN: Voronoi (Thiessen) Polygons (Concepts and Techniques in Modern Geography); Norwich: Geo Books, 1986).

The polygons of all patients starting at the same renal centre were combined to create the catchment area for that centre. The catchment area for one centre might comprise multiple unconnected polygons as a result of adjacent patients attending different renal centres. The Office for National Statistics (ONS) map of 2011 census merged wards contains population estimates for England and Wales divided into 8,546 wards.

The Northern Ireland Statistics and Research Agency (NISRA) published population estimates based on the 2011 census for 4,537 geographical regions referred to as small areas. The General Register Office for Scotland published 2011 population estimates at 6,505 data zone level areas. Wards, small areas and data zones will collectively be referred to as wards in the following paragraphs.

The wards were overlaid on the map of renal centre catchment areas, enabling the proportion of each ward's area covered by each of the renal centre catchment areas to be calculated. Each ward's population was then allocated to the renal centres in proportions equal to the proportions of the overlaid areas. Summing these proportions of populations across all of the wards for each renal centre produced the estimates of the total catchment population for each centre.

There are some limitations to these estimates. The main one is that the ward allocated to each renal centre was based upon dialysis patients only. Therefore it is possible that non-dialysis patients may come from a different

catchment population. This is more likely where a renal centre provides specialist services and especially likely for patients undergoing kidney transplantation. The catchment population for kidney Tx patients will depend largely upon the distribution of workload between the referral centre and the transplanting centre for pre-Tx work-up, donor nephrectomy work-up and post-Tx care (including if and when care is returned to the referring centre).

Despite the limitations, this is the most valid methodology to date to estimate the size of the catchment populations for renal centres in the UK. Thanks are expressed to Professor Andrew Judge for calculating these catchment populations for the UKRR.

There is a need for centre catchment populations to be re-estimated for future analyses because populations will have changed. For the 2018 analyses, the catchment population of each centre was updated by upscaling the previous estimate so that for each UK country, the sum of centres' populations covered was equal to the 2018 national adult ( $\geq$ 18 years) population estimates from the ONS.

#### 6.2 Adjusted analyses

Most analyses presented in this report are unadjusted. However, a few analyses are adjusted to take into account the difference in baseline characteristics between groups that may influence the outcome, thereby allowing better comparisons between renal centres. See each chapter for more details.

#### 6.3 Graphs

Percentages achieving The Renal Association guidelines and other standards are displayed in several ways in the UKRR Annual Report.

#### 6.3.1 Caterpillar plots

Caterpillar plots show the percentage meeting the targets along with 95% confidence intervals (CIs) for each centre, country and overall.

#### 6.3.2 Funnel plots

Funnel plots show the percentage meeting the target plotted against the size of the centre (the number of people with a measurement, or the number of patient years at risk). A 'funnel' is plotted either around the average percentage meeting the target or the target itself as specified in the plot title. There is evidence that any centres which fall outside the funnel are significantly different from the average or the target. The funnel shape of the limits reflects the fact that for smaller centres, for which the percentage meeting the target is less reliably estimated, a greater observed difference from the average/target is required for it to be statistically significantly different.

In each funnel plot, the lines (see legends) indicate the national mean and the 95% and 99.7% CIs as stated, corresponding to two and three standard deviations from the mean, respectively. Each point on the plot represents one renal centre. For each outcome measure, if no significant inter-centre variation was present, three of 71 adult renal centres would be expected to fall between the 95% and 99.7% CIs and no centre should fall outside the 99.7% CI. In survival analysis the funnel plot methodology is similar except that the funnel plots show the percentage survival plotted against the size of the centre (the number of patients in the cohort) and a 'funnel' is plotted around the average survival in the UK. Survival for any centres falling outside the 95% CIs is therefore significantly different from the average survival in the UK.

#### 6.3.3 Box and whisker plots

These are only used to report MSSA and MRSA infection rates. The box shows the median in the middle and

the upper and lower quartiles, i.e. 25th and 75th centiles. The whiskers show the full measured range of that variable.

#### 6.3.4 Kaplan-Meier (KM) method/plots

In the KM method, the probability of surviving more than a given time period can be estimated for all members of a cohort of patients overall (or by subgroup such as age group). Its estimator is a series of declining horizontal steps that approaches the true survival function for the given population with a large enough sample size. The declining step function (i.e. the KM curve) takes the censoring of data into account (right-censoring in the UKRR analysis), which occurs if the patient is lost to follow-up or is alive without the event occurrence at last follow-up. The KM method can also estimate median time to event in conjunction with right-censoring information; median time is when 50% of patients within the population experienced the event (see section 8.1 for further discussion of the KM methods used in the survival analysis).

#### 6.4 How to interpret centre-specific analyses and outlying centres

The UKRR continues to advise caution in the interpretation of the comparisons of centre-specific attainment of clinical audit measures provided in this report. As in previous reports, the UKRR does not test for 'significant difference' between centres and arbitrary 95% and 99.7% CIs are created from the data to show compliance with an audit standard.

For a number of years de-anonymised centre-specific reports on survival of RRT patients have been published in the annual report. Centres are contacted if survival is lower than expected in patients starting dialysis and for prevalent RRT patients

The UKRR has no statutory powers. However, because the UKRR provides centre-specific de-anonymised analyses of important clinical outcomes, including survival, it is important to define how the UKRR responds to apparent under-performance. The UKRR senior management team communicates survival outlier status with the renal centres prior to publication. Centres are asked to report their outlying status internally at trust level and to follow-up with robust mortality and morbidity meetings. They are also asked to provide evidence that the clinical governance department and chief executive of the trust housing the service have been informed. In the event that no such evidence is provided, the chief executive officer or medical director of the UKRR inform the president of the Renal Association, who then takes action to ensure that the findings are properly investigated.

# 7. Populations and analyses by annual report chapter

Analyses in the report are presented on cohorts of patients who share either the time at which they initiated RRT e.g. incident population, or share a treatment modality e.g. PD patients.

#### 7.1 Incident adult renal replacement therapy (RRT) population (chapter 1)

The incident adult RRT population is all patients aged 18 years and over with ESKD who started RRT (dialysis or pre-emptive Tx) at a UK renal centre for the first time in the calendar year applicable to the analyses. It excludes patients who recover their renal function for >90 days within 90 days of starting dialysis. Furthermore, patients restarting dialysis after a failed Tx are not counted as incident patients. A patient can therefore appear only once in the incident cohort.

The treatment timeline is used to define incident patients. If a patient has timeline entries from more than one centre these are combined and sorted by date. The first RRT treatment entry from any centre is used to determine the first date they commenced RRT. This is defined as a 'start date'. However, in the following

situations there is evidence that the patient was already receiving RRT before this 'start date' and consequently these people are not classed as incident patients:

- those with an initial entry on the timeline of transferred in (modality codes 39 to 69)
- those with an initial entry of transferred out (modality code 38)
- those with an initial treatment of lost to follow-up (modality code 95)
- those who had an initial graft acute rejection (modality code 31) and did not have a Tx on the same day
- those with an initial entry of transfer to adult nephrology (modality code 37)
- those with an initial entry of graft functioning (modality code 72)
- those with an initial entry of nephrectomy Tx (modality code 76).

Where none of the above apply, the patient is defined as an incident patient, providing there is no recovery of >90 days starting within 90 days of the start date. If there is a recovery lasting >90 days, modality codes after this date would indicate that the patient restarted RRT. If they did, this second (or third etc.) starting point is defined as their start date, providing that they did not have a recovery lasting >90 days starting within 90 days of start.

Provided the UKRR received a modality code 36 (pre-emptive Tx) from the work-up centre, pre-emptive Tx are allocated as incident patients of the work-up centre and not of the centre where the Tx took place.

NHS England mandates the collection of data regarding acute HD sessions. However, sessional HD data carry no information about whether the HD was for AKI or ESKD. Distinguishing between these two indications depends entirely upon the accuracy of timeline data provided by centres.

Patients who receive acute HD are only reported if their dialysis is subsequently recoded as being for ESKD, when they fail to recover native renal function. Recoding to RRT is automatically applied at 90 days for individuals still on RRT, unless the centre confirms a patient was on an unusually long period of dialysis for acute renal failure, but can also be applied at any point between zero and 90 days by the reporting centre. Individuals who commence HD for AKI (i.e. acute HD by definition) and subsequently recover renal function or die within the first 90 days of treatment without receiving an ESKD code are the focus of a separate piece of work.

Differences in RRT incidence can be seen in the most recent years when compared with previous publications because of retrospective updating of data in collaboration with renal centres. In addition, patients with AKI requiring dialysis may be coded in the subsequent year as having developed ESKD, allowing the UKRR to backdate the start date of RRT.

#### 7.2 Prevalent adult RRT population (chapter 2)

The prevalent adult RRT population is all patients on RRT for ESKD aged 18 years and over at a UK renal centre who were alive on 31 December of the year applicable to the analyses. It includes both incident patients for that year (who remained on RRT until the end of the year) and patients who have been on RRT for longer. Excluded are patients who had transferred out, recovered renal function, stopped treatment without recovery of function, died or were lost to follow-up before the end of the year. Patients who had transferred out, then transferred in to another centre before the end of the year would be included at the incoming centre. Also excluded are any patients aged 18 years and over still being treated at a paediatric renal centre.

When quarterly data are received from more than one centre (often when there is joint care of kidney Tx recipients between the referring centre and the Tx centre) the patient is only included under one of these. The

allocated centre is defined by the steps below (as many steps as necessary are followed in this order until data are only left from one centre):

- the treatment timeline is used to eliminate any centre(s) which the patient was not still attending, at the end of the quarter
- a centre with biochemistry data (at least one of the six fields: creatinine, Hb, albumin, aluminium, serum potassium, urea) is favoured over one without
- a centre with quarterly modality of Tx is favoured over one without
- non-Tx centres are favoured over Tx centres
- the centre with the highest number of the six biochemistry fields (listed above) populated is favoured
- if the above steps do not decide between centres (unusual) then the choice is made based on the order in which the centres appear in the data.

In some situations (generally where timeline data are seen to be inaccurate/incomplete) the centre used is set manually on an ad hoc basis.

There are exclusions for analyses of quarterly biochemistry or blood pressure data:

- patients who had 'transferred in' to the centre in that particular quarter are excluded
- patients who had changed treatment modality in that particular quarter are excluded
- patients who had been on RRT for <90 days are excluded.

Note the length of time on RRT is calculated from the most recent start date (i.e. the point at which they are defined as an incident patient using the definition detailed above). So if a patient starts, then recovers and then starts again, this second start date is used. Also, for patients who are not defined as incident patients because their start date is unknown (for example, if their first timeline entry is a transfer in code) it is assumed that they have been on RRT for  $\geq 90$  days and they are included for every quarter.

#### 7.3 Prevalent adult transplant (Tx) population (chapter 3)

There are 23 UK adult renal Tx centres – 19 in England, two in Scotland and one each in Northern Ireland and Wales.

Annual organ-specific updates with comprehensive data concerning the number of patients on the Tx waiting-list, percentage of pre-emptive listing, the number of transplants performed, the number of deceased kidney donors (DBD and DCD), LKDs, patient survival and graft survival are available on the NHSBT website.

Where joint care of kidney Tx recipients between the referring centre and the Tx centre occurs, the patient is usually allocated to the referring centre (see section 7.2). Thus, the number of patients allocated to a Tx centre is often lower than that recorded by the centre itself and conversely, pre-emptively transplanted patients are sometimes allocated to the transplanting centre rather than the referring centre if no transfer out code is submitted to the UKRR. Queries and updated information are welcomed by the UKRR at any point during the year if this has occurred.

The median PTH by CKD stages is reported nationally, despite poor PTH completeness across all centres – therefore this has to be interpreted with caution. PTH is submitted to the UKRR in two different units from different centres (pmol/L or pg/mL). We assume each centre submits PTH using the same unit for all patients within their centre. During our data cleaning process, we convert the data to pmol/L if the overall median PTH of the centre suggested they had used pg/mL.

In the eGFR slope analysis, a minimum duration of 18 months graft function is required and three or more creatinine measurements from the second year of graft function onwards are used to plot the eGFR slope. If a Tx failed but there are at least three creatinine measurements between one year post-Tx and graft failure, the patient is included, but no creatinine measurements after the quarter preceding the recorded date of Tx failure are analysed. Slopes are calculated using linear regression, assuming linear change in eGFR over time and the effect of age, ethnicity, sex, diabetes, donor type, year of Tx and current Tx status are analysed.

#### 7.4 Prevalent adult in-centre haemodialysis (ICHD) population (chapter 4)

This chapter describes the population of adult patients with ESKD who were receiving ICHD in the UK at the end of the year applicable to the analyses. Throughout this chapter, ICHD refers to all modes of ICHD treatment, including haemodiafiltration (HDF). Several centres reported significant numbers of patients on HDF, but other centres did not differentiate this treatment type in their UKRR returns. Analyses in this chapter exclude patients on HHD unless stated – HHD patients are analysed in a separate chapter.

#### 7.5 Prevalent adult peritoneal dialysis (PD) population (chapter 5)

The PD chapter includes analyses of prevalent patients on continuous ambulatory PD (CAPD) and automated PD (APD).

#### 7.6 Prevalent adult home haemodialysis (HHD) population (chapter 6)

The HHD chapter includes analyses of prevalent patients on home haemodialysis. Due to small numbers, haematological and biochemical results are not shown for many of the UK renal centres. Renal centres are not required to submit changes in dialysis modality that last <30 days, so it is difficult to correctly attribute an infection to HHD or ICHD. Therefore analysis of infections is presented in the ICHD chapter for ICHD and HHD combined.

#### 7.7 Paediatric RRT population (chapter 7)

This chapter describes the population of children (aged <18 years) with ESKD who received RRT in the year applicable to the analyses. Definitions of 'incident' and 'prevalent' cohorts are equivalent to those used in the analysis of adult RRT patients. However, by contrast to adult chapters, paediatric patients treated in paediatric renal centres and coded as ESKD who died within the first 90 days of RRT are excluded from the paediatric analyses.

In the UK, RRT for children is managed by 13 paediatric renal centres, all of which are equipped to provide both HD and PD. Ten of these centres also perform kidney transplantation. Young people aged 16–18 years may be managed in either paediatric or adult renal centres. This is variable across the UK and dependent on local practices, social factors and patient/family wishes.

In this chapter, data are reported separately for patients aged <16 years who are managed within UK paediatric renal centres and for young people aged 16–18 years (including both young adults managed by paediatric renal centres and those who received nephrology care from adult renal centres).

The populations used to calculate incidence and prevalence are obtained from the ONS. The mid-current-year population estimate produced by the ONS, based on the 2011 census, is used to calculate the current year incidence and prevalence rates. For analyses performed using historic years, an incident 15 year cohort is divided into three five year periods – with the mid-year estimate for each five year period being used as the population estimate. Incidence and prevalence for 16–18 year olds are also reported, however these are possibly

under-estimated because adult centres are not currently required to send data on young people aged <18 years.

PRD is coded according to 2012 diagnostic groupings used by the ERA-EDTA: these include tubulointerstitial disease, glomerular disease, familial and hereditary nephropathies, systemic disease affecting the kidney and miscellaneous. Further details on how PRDs are coded and grouped can be found on the ERA-EDTA website.

Data for height, weight, BMI and blood pressure vary with age, sex and size and are therefore presented as z-scores as described in the chapter.

Analysis of cardiovascular risk factors is shown in children <16 years old. Risk factors considered are hypertension (SBP and/or DBP over the 90th percentile), BMI (overweight or obese, defined as an height-age z-score  $\geq$ 1.3 in male and  $\geq$ 1.19 in female) and hypercholesterolaemia (cholesterol >5.2 mmol/L, and/or high triglycerides, defined as triglycerides >1.13 mmol/L for those aged under 9 years and >1.46 mmol/L for those aged 9 years and over).

Table A5 Summary of age-specific biochemical clinical audit measures for children

	Age (years)			
Parameter	<1	1–5	6–12	>12
Hb (g/L)	Maintain 95–115 if aged <2 years	Maintain 100–120 if aged ≥2 years	100-120	100-120
Adjusted calcium (mmol/L)	2.24-2.74	2.19-2.69	2.19-2.69	2.15-2.55
Phosphate (mmol/L)	1.10-1.95	1.05-1.75	1.05-1.75	1.05-1.75
PTH (individual centre)	Within twice the normal range			
	Levels may be maintained within normal range if growing appropriately			
Bicarbonate* (mmol/L)	Reported as either within or outside centre reference range			

<sup>\*</sup>In young adults, the range of 20–26 mmol/L was used.

# 8. Specific analyses for adults

### 8.1 Survival and cause of death analyses

The unadjusted survival probabilities (with 95% CIs) are calculated using the KM method, in which the probability of surviving more than a given time can be estimated for all members of a cohort of patients overall or by subgroup such as age group, but without any adjustment for confounding factors such as age that affect the chances of survival. Where centres are small, or the survival probabilities are >90%, the CIs are only approximate.

To estimate the difference in survival of different subgroups of patients within the cohort, a stratified proportional hazards model (Cox) is used where appropriate. The results from the Cox model are interpreted using a hazard ratio. When comparing two groups, the hazard ratio is the ratio of the estimated hazard for group A relative to group B, where the hazard is the risk of dying at time *t* given that the individual has survived until this time. The underlying assumption of a proportional hazards model is that the hazard ratio remains constant throughout the period under consideration. Whenever used, the assumptions of the proportional hazards model are tested.

To allow for comparisons between centres with differing age distributions, survival analyses are adjusted for age and reported as survival adjusted to age 60 years. This gives an estimate of what the survival would have been if all patients in that centre had been aged 60 years at the start of RRT. This age was chosen because it was

Hb – haemoglobin; PTH – parathyroid hormone

approximately the average age of patients starting RRT 17 years ago at the start of the UKRR's data collection. The average age of patients commencing RRT in the UK has recently stabilised around 64 years, but the UKRR has maintained age adjustment to 60 years for comparability with all previous years' analyses.

For some analyses, further adjustment was carried out for not only age, but also sex and comorbidities. Comorbidity data derived from diagnostic and procedure codes in HES and PEDW were used to augment comorbidity data submitted by renal centres to the UKRR. A comorbidity score was derived from a multivariable Cox proportional hazards model including all the comorbidities. A score was allocated to each comorbidity according to the size of the hazard ratios estimated from the model. A score for each patient was calculated by summing the scores of the individual comorbidities present for the patient.

Defining when a patient starts RRT (day zero) is reliant on centres following consistently the methodology described in section 4.1.3. Previous work suggests that is not always the case. As well as variability in defining start date within the UK, there is international variability when patient data are collected by national registries (often for financial reimbursement or administrative reasons). Some countries define the 90th day after starting RRT as day zero, whilst others collect data only on those who have survived 90 days and report as zero the number of patients dying within the first 90 days.

Therefore, as many other national registries do not include reports on patients who do not survive the first 90 days, survival from 90 days onwards is also reported to allow international comparisons. This distinction is important, as there is a much higher death rate in the first 90 days, which would distort comparisons.

#### 8.1.1 Methodology for incident patient survival

Patients incident to RRT are analysed over a number of years as stated in each analysis to help more readily identify differences between the survival of the populations being compared. Two years' incidence data is used to identify differences between the four UK countries. One year after 90 day survival using a rolling four year combined incident RRT cohort is used to compare survival between centres. A 10 year rolling cohort is used when analysing trends over time and for long term survival.

The incident survival cohort is not censored at the time of transplantation and therefore includes the survival of the subset of patients who start RRT with a pre-emptive Tx. An additional reason for not censoring is to facilitate comparison between centres. Centres with a high proportion of patients of South Asian and Black origin are likely to have a healthier dialysis population, because South Asian and Black patients are less likely to undergo early transplantation and centres with a high pre-emptive Tx rate are likely to have a less healthy dialysis population because transplantation selectively removes fitter patients.

The one year incident survival is for patients who started RRT from 1 October or two years earlier until the 30 September of the previous year and followed-up for one full year (e.g. patients starting RRT on 1 December 2016 are followed through to 30 November 2017). Using the same example, for analysis of one year after 90 day survival, patients who started RRT from 1 October 2016 until 30 September 2017 are included in the cohort and are followed-up for a full year after the first 90 days of RRT.

The death rate per 1,000 patient years is calculated by dividing the number of deaths by the person years exposed. Person years exposed are the total years at risk for each patient (until death, recovery or lost to follow-up). The death rate is presented by age group.

Case-mix adjustment of one year after 90 day survival for the effect of age, sex and comorbidity is undertaken using a rolling four year combined incident RRT cohort. Data on age and sex are 100% complete. Only those centres returning ≥85% of comorbidity data (after augmentation from HES and PEDW) for patients in the combined cohort are included. A Cox proportional hazards model with statistical frailty was fitted to account for

heterogeneity and random effects between renal centres.

#### 8.1.2 Methodology for prevalent dialysis patient survival

The prevalent dialysis patient group is defined as all adults, alive and receiving dialysis at the start of the given year who had been on dialysis for at least 90 days at one of the UK adult renal centres. It does not include patients coded as being on chronic dialysis but yet to reach 90 days, unlike other definitions of the prevalent population. Prevalent dialysis patients on 31 December of the previous year are followed-up in the current year and are censored at transplantation. When a patient is censored at transplantation, this means that the patient is considered as alive up to the point of transplantation, but the patient's status post-Tx is not considered.

Case-mix adjusted 1 year survival for prevalent dialysis patients at the end of 2017 is reported. The methodology followed is the same as described in section 8.1.1.

As discussed in previous reports, comparison of survival of prevalent dialysis patients between centres is complex. Survival of prevalent dialysis patients can be studied with or without censoring at transplantation and it is common practice in some registries to censor at transplantation. Censoring could cause apparent differences in survival between those renal centres with a high Tx rate and those with a low Tx rate, especially in younger patients where the Tx rate is highest. Censoring at transplantation systematically removes younger, fitter patients from the survival data. The differences are likely to be small due to the relatively small proportion of patients being transplanted in a given year compared to the whole dialysis population (about 9% of the dialysis population aged <65 years and about 2% of the population aged  $\geq$ 65 years). To allow comparisons with other registries, the survival results for prevalent dialysis patients censored for transplantation are quoted. To understand survival of patients, including survival following transplantation, the incident patient analyses should be viewed.

# 8.1.3 Methodology for comparing mortality in prevalent RRT patients with mortality in the general population

Data on the UK population in mid-2018 and the number of deaths in each age group in 2018 were obtained from the ONS. The age-specific UK death rate was calculated as the number of deaths in the UK per 1,000 people in the population. The age-specific expected number of deaths in the RRT population was calculated by applying the UK age-specific death rate to the total of years exposed for RRT patients in that age group. This is expressed as deaths per 1,000 patient years. The age-specific number of RRT deaths is the actual number of deaths observed in 2018 in RRT patients. The RRT observed death rate was calculated as number of deaths observed in 2018 per 1,000 patient years exposed. Relative risk of death was calculated as the ratio of the observed and expected death rates for RRT patients. The death rate was calculated for the UK general population by age group and compared with the same age group for prevalent patients on RRT on 31 December 2018.

#### 8.1.4 Methodology of cause of death

Completeness of cause of death data is calculated for all prevalent patients on RRT who died in a specific year with cause of death data completed for that year. Patients who were lost to follow-up or who recovered are not included in the cause of death completeness calculation.

Adult patients from England, Wales, Scotland and Northern Ireland are included in the analyses of cause of death. The incident patient analysis included all patients starting RRT in the years 2014–2017. Analysis of prevalent patients included all those aged  $\geq$ 18 years and receiving RRT on 31 December 2017 and followed-up for one year in 2018.

#### 8.2 Dialysis access

Each year, all adult renal centres in England, Wales and Northern Ireland are asked to provide vascular and peritoneal access data for incident and prevalent dialysis patients. The Scottish Renal Registry provides a separate dataset including access at start for all incident patients. We do not include Scottish patients in any subgroup analysis by early/late referral because of differences in definition (see section 4.1.1 and 4.1.2), nor in analysis of surgical assessment time, PD catheter insertion technique and access failure because data are not provided. Access data for incident patients are collected at patient level, whereas centre level data are submitted for prevalent patients. Records are validated against the UKRR database to confirm that the population collected at each centre for the audit was representative of the incident/prevalent population at that centre collected via the routine quarterly return.

For the purposes of this audit, patients categorised as having AKI are excluded from the analyses as well as those with missing information for access at start, age and date of starting RRT. From this year onwards, patients who did not start dialysis for the first time in 2018 based on UKRR quarterly data submissions were excluded, as were those aged <18 years. If a centre returns audit data for less than 70% of the incident or prevalent patients, it is excluded from analyses of that centre. Cross-referencing with the UKRR database also enables ascertainment of mortality within three months of commencing dialysis.

Patients starting HD are grouped by type of first vascular access: AVF, AVG, TL and NTL. Patients starting PD are categorised by the insertion technique: open surgery, laparoscopic, peritoneoscopic or percutaneous. Access at three months is defined as the type of access in use at three months after starting dialysis. If a patient is no longer receiving dialysis at three months (but had not recovered renal function), the reason is recorded instead, for example, 'death' or 'transplantation'. Referral time is defined as the number of days between the date of first being seen by a renal physician (as an inpatient or outpatient) and the date of commencing dialysis. A patient is classified as presenting 'late' if they have a referral time of <90 days.

Access failure is defined as when it is no longer usable for dialysis, with the date and cause of access failure reported. PD technique failure is grouped into six causes: infection, catheter related, solute/water clearance, leaks/hernia, other and unknown. There were no patients with a failure caused by solute/water clearance this year. Access failure is censored for death, transplantation, withdrawal from RRT and elective switching of access type. It is the intention to only capture access failures relating to the first access that is performed. If the reason recorded for access failure is not related to the first type of access recorded, then the data are not included in this analysis.

Centres that report data on PD patients in the previous vascular and peritoneal access audit are asked to complete a one year follow-up of their PD patients. Additional information is requested on the date of PD catheter failure, the reason for catheter failure, the number of catheters used during the year and the modality in use at one year after starting PD. Analyses that use these data are titled 'PD follow-up audit'.

Dialysis access is best interpreted in the context of all patients starting RRT, thus data for pre-emptive Tx recipients are included and sourced from the UKRR database to augment the dialysis access audit data. This reflects the amended (2014) Renal Association guidelines for planned RRT initiation, which include Tx in the audit standard. Tx and non-Tx centres work together to prepare patients for Tx, but for the purpose of these analyses, patients are allocated to their most likely treatment centre (Tx or non-Tx).