All treatments for chronic kidney disease (CKD) need to be tailored to the individual patient. The following recommendations are useful starting points for the majority of cats at each stage. Serial monitoring of these patients is ideal and treatment should be adapted according to the response to treatment. Note that staging of disease is undertaken following diagnosis of CKD – an increased blood creatinine concentration alone is not diagnostic of CKD.

Treatment recommendations fall into two broad categories, namely:

1. Those that slow progression of CKD and so preserve remaining kidney function for longer
2. Those that address the quality of life of the cat, addressing the symptoms of CKD

In general, at the early stages of CKD (stages 1 and 2), there are few clinical extra-renal signs of the disease and the therapeutic emphasis is on slowing progression. From stage 3 onwards, extra-renal signs become more common and more severe. The importance of administering treatments which are symptomatic and improve the cat’s quality of life assumes greater importance and exceeds the importance of treatments designed to slow progression by stage 4.

Some of the treatment recommendations are not authorized for use in all geographical regions and some may not be authorized for use in cats. Such recommended dose rates are therefore empirical. It is the treating veterinarian’s duty to make a risk:benefit assessment for each patient prior to administering any treatment.
Treatment Recommendations for CKD in Cats (2017)

Treatment recommendations for Cats with Chronic Kidney Disease

Stage 1 Feline patients:
1. Discontinue all potentially nephrotoxic drugs if possible.
2. Identify and treat any pre-renal or post-renal abnormalities.
3. Rule out any treatable conditions like pyelonephritis (any urinary tract infection should be regarded as a potential pyelonephritis and treated appropriately) and renal urolithiasis with radiographs and/or ultrasonography.
4. Measure blood pressure and urine protein to creatinine ratio (UP/C).

Management of dehydration:
In these patients urine concentrating ability may be somewhat impaired and therefore
- correct clinical dehydration/hypovolemia with isotonic, polyionic replacement fluid solutions (e.g., lactated Ringer’s) IV or SQ as needed.
- have fresh water available at all times for drinking.

Systemic hypertension:
The blood pressure above which progressive renal injury may be induced is unknown. Our goal is to reduce systolic blood pressure to <160 mm Hg and to minimize the risk of extra-renal target organ damage (CNS, retinal, cardiac problems/damage). If there is no evidence of such damage but systolic blood pressure persists above 160 mm Hg, increasing the risk of this occurring, treatment should be instituted.

‘Persistence’ of increased systolic blood pressure should be judged on multiple measurements made over the following time-scales in these blood pressure substages:
- Hypertensive (moderate risk of future target organ damage) – systolic blood pressure 160 to 179 mm Hg measured over 1 to 2 months
- Severely hypertensive (high risk of future target organ damage) – systolic blood pressure ≥ 180 mm Hg over 1 to 2 weeks.

If evidence of target organ damage exists, cats should be treated without the need to demonstrate persistently increased systolic blood pressure. Reducing blood pressure is a long term aim in managing the patient with CKD and a gradual and sustained reduction should be the goal, avoiding any sudden or severe decreases leading to hypotension.

A logical stepwise approach to managing hypertension is as follows:
1. Dietary sodium (Na) reduction - there is no evidence that lowering dietary Na will reduce blood pressure. If dietary Na reduction is attempted, it should be accomplished gradually and in combination with pharmacological therapy.
2. Calcium channel blocker (CCB), such as amlodipine (0.125 to 0.25 mg/kg once daily).
3. Double the dose of amlodipine (0.25 to 0.5 mg/kg once daily).

4. Combine an inhibitor of the renin-angiotensin-aldosterone system (RAAS; either an angiotensin converting enzyme inhibitor [ACEI, such as benazepril] or an angiotensin receptor blocker [ARB, such as telmisartan]) and CCB treatment. Note: Take care not to introduce CCB/RAAS inhibitor treatment to unstable dehydrated cats as glomerular filtration rate may drop precipitously if these drugs are introduced before the patient is adequately hydrated.

**Monitoring response to antihypertensive treatment:**

Hypertensive cats normally require lifelong therapy and may require treatment adjustments. Serial monitoring is essential. After stabilization, monitoring should occur at least every 3 months.

- Systolic blood pressure <120 mm Hg and/or clinical signs such as weakness or tachycardia indicate hypotension, which is to be avoided.
- Blood creatinine concentration – reducing blood pressure may lead to small and persistent increases in creatinine concentration (<45 µmol/l or 0.5 mg/dl increase), but a marked increase suggests an adverse drug effect. Progressively increasing concentrations indicate progressive kidney damage/disease.

**Proteinuria:**

Cats in Stage 1 with UP/C >0.4 should be investigated for disease processes leading to proteinuria (see 1 and 2 below) and treated with anti-proteinuric measures (see 3 and 4 below).

Those with borderline proteinuria (UP/C 0.2 to 0.4) require close monitoring (see 1 and 4 below).

1. Look for any concurrent associated disease process that may be treated/corrected.
2. Consider kidney biopsy as a means of identifying underlying disease (see Appendix and consult experts if unsure of indications for kidney biopsy).
3. Administer an RAAS inhibitor (ACEI or ARB) and feed a clinical renal diet.
4. Monitor response to treatment / progression of disease:
   - stable blood creatinine concentration and decreasing UP/C = good response.
   - serially increasing blood creatinine concentrations and/or increasing UP/C = disease is progressing.

Ordinarily therapy will be maintained lifelong unless the underlying disease has been resolved in which case dose reduction whilst monitoring UP/C might be considered.

Note:

a. The use of an RAAS inhibitor is contraindicated in any animal that is clinically dehydrated and/or is showing signs of hypovolemia. Correct dehydration before using these drugs otherwise glomerular filtration rate may drop precipitously.

b. Cats with proteinuria and hypoalbuminemia likely share the same thromboembolic
risk as dogs, but aspirin is difficult to use in cats to achieve a selective antiplatelet effect. A suggested dose rate if plasma albumin is below 20 g/l (2 g/dl) is 1 mg/kg every 72 hours.

c. Cats with serum phosphate within the IRIS target may be at increased risk of developing hypercalcemia when renal diets are introduced. Monitor serum calcium and if total calcium exceeds 12 mg/dl (3 mmol/l) switch the cat to a senior diet or mix renal diet (50:50 by volume) with standard grocery food.

d. If borderline proteinuria persists, antiproteinuric treatment could be offered, because the association between progression of CKD and proteinuria includes the borderline category. However, there is at present no evidence that intervention with anti-proteinuric drugs slows progression.

Stage 2 Feline patients:
All of the above listed for Stage 1 plus any additional steps indicated.

1. Discontinue all potentially nephrotoxic drugs if possible.
2. Identify and treat any pre-renal or post-renal abnormalities.
3. Rule out any treatable conditions like pyelonephritis (any urinary tract infection should be regarded as a potential pyelonephritis and treated appropriately) and renal urolithiasis with radiographs and/or ultrasonography.
4. Measure blood pressure and urine protein to creatinine ratio (UP/C).
5. Consider feeding a clinical renal diet: this may be accomplished more easily early in the course of CKD, before inappetence develops.

Management of dehydration:
These patients have decreased urine concentrating ability and therefore
- correct clinical dehydration/hypovolemia with isotonic, polyionic replacement fluid solutions (e.g., lactated Ringer’s) IV or SQ as needed.
- have fresh water available at all times for drinking.

Systemic hypertension:
The blood pressure above which progressive renal injury may be induced is unknown. Our goal is to reduce systolic blood pressure to <160 mm Hg and to minimize the risk of extra-renal target organ damage (CNS, retinal, cardiac problems/damage). If there is no evidence of this but systolic blood pressure persistently exceeds 160 mm Hg, increasing the risk of such damage, treatment should be instituted.

‘Persistence’ of increased systolic blood pressure should be judged on multiple measurements made over the following time-scales in these blood pressure substages:
- Hypertensive (moderate risk of future target organ damage) – systolic blood pressure 160 to 179 mm Hg measured over 1 to 2 months.
- Severely hypertensive (high risk of future target organ damage) – systolic blood pressure ≥180 mm Hg over 1 to 2 weeks.

If evidence of target organ damage exists, cats should be treated without the need to demonstrate persistently increased systolic blood pressure. Reducing blood pressure
is a long term aim in managing the patient with CKD and a gradual and sustained reduction should be the goal, avoiding any sudden or severe decreases leading to hypotension.

A logical stepwise approach to managing hypertension is as follows:

1. Dietary sodium (Na) reduction - there is no evidence that lowering dietary Na will reduce blood pressure. If dietary Na reduction is attempted, it should be accomplished gradually and in combination with pharmacological therapy.
2. Calcium channel blocker (CCB) such as amlodipine (0.125 to 0.25 mg/kg once daily).
3. Double the dose of amlodipine (0.25 to 0.5 mg/kg once daily).
4. Combine an inhibitor of the renin-angiotensin-aldosterone system (RAAS; either an angiotensin converting enzyme inhibitor [ACEI, such as benazepril] or an angiotensin receptor blocker [ARB, such as telmisartan]) with the CCB

Take care not to introduce CCB/RAAS inhibitor treatment to unstable dehydrated cats as glomerular filtration rate may drop precipitously if these drugs are introduced before the patient is adequately hydrated.

Monitoring response to antihypertensive treatment:

Hypertensive cats normally require lifelong therapy and may require treatment adjustments. Serial monitoring is essential. After stabilization, monitoring should occur at least every 3 months.

Systolic blood pressure <120 mm Hg and/or clinical signs such as weakness or tachycardia indicate hypotension, which is to be avoided.

Blood creatinine concentration – reducing blood pressure may lead to small and persistent increases in creatinine (<45 µmol/l or 0.5 mg/dl increase), but a marked increase suggests an adverse drug effect. Progressively increasing creatinine concentrations indicate progressive kidney damage/disease.

Proteinuria:

Cats in Stage 2 with UP/C >0.4 should be investigated for disease processes leading to proteinuria (see 1 and 2 below) and treated with anti-proteinuric measures (see 3 and 4 below).

Those with borderline proteinuria (0.2 to 0.4) require close monitoring (see 1 and 4 below).

1. Look for any concurrent associated disease process that may be treated/corrected.
2. Consider kidney biopsy as a means of identifying underlying disease (see Appendix and/or consult experts if unsure of indications for kidney biopsy).
3. Administer an RAAS inhibitor (ACEI or ARB) and feed a clinical renal diet.
4. Monitor response to treatment/progression of disease:
   – stable blood creatinine concentration and decreasing UP/C = good response.
   – serially increasing creatinine concentrations and/or increasing UP/C = disease is progressing.
Ordinarily therapy will be maintained lifelong unless the underlying disease has been resolved in which case dose reduction while monitoring UP/C might be considered.

Note:

a. Use of an RAAS inhibitor is contraindicated in any animal that is clinically dehydrated and/or showing signs of hypovolemia. Correct dehydration before using these drugs otherwise glomerular filtration rate may drop precipitously.

b. Cats with proteinuria and hypoalbuminemia likely share the same thromboembolic risk as dogs, but aspirin is difficult to use in cats to achieve a selective antiplatelet effect. A suggested dose rate if plasma albumin is below 20 g/l (2 g/dl) is 1 mg/kg every 72 hours.

c. Cats with serum phosphate within the IRIS target may be at increased risk of developing hypercalcemia when renal diets are introduced. Monitor serum calcium and if total calcium exceeds 12 mg/dl (3 mmol/l) switch the cat to a senior diet or mix renal diet (50:50 by volume) with standard grocery food.

d. If borderline proteinuria is persistent, antiproteinuric treatment could be offered. The rationale for doing so is that there is an association between progression of CKD and proteinuria which extends into the borderline category. There is no evidence that intervention with antiproteinuric drugs slows progression, however.

Reduction of phosphate intake:

Many cats in Stage 2 will have normal plasma phosphate concentrations but will have increased plasma PTH concentration. Evidence suggests that chronic reduction of phosphate intake to maintain a plasma phosphate concentration below 1.5 mmol/l (but not less than 0.9 mmol/l; <4.6 mg/dl but >2.7 mg/dl) is beneficial to patients with CKD.

The following measures can be introduced sequentially in an attempt to achieve this:

1. Dietary phosphate restriction (i.e., clinical renal diet therapy).

2. If plasma phosphate concentration remains above 1.5 mmol/l (4.6 mg/dl) after dietary restriction, give enteric phosphate binders (such as aluminum hydroxide, calcium carbonate, calcium acetate, lanthanum carbonate) to effect, starting at 30-60 mg/kg/day in divided doses to be mixed with each meal (mixed with the food). The dose required will vary according to the amount of phosphate being fed and the stage of kidney disease. Treatment with phosphate binders should be to effect (as outlined above), with signs of toxicity limiting the upper dose rate possible in a given patient. Monitor serum calcium and phosphate concentrations every 4-6 weeks until stable and then every 12 weeks. Microcytosis and/or generalized muscle weakness suggests aluminum toxicity if using an aluminum containing binder – switch to another form of phosphate binder should this occur. Hypercalcemia should be avoided – combinations of aluminum and calcium containing phosphate binders may be necessary in some cases.
Metabolic acidosis:
If metabolic acidosis exists (blood bicarbonate or total CO₂ is stabilized on the diet of choice, supplement with oral sodium bicarbonate (or potassium citrate if hypokalemic) to effect to maintain blood bicarbonate / total CO₂ in the range of 16-24 mmol/L.

Additional recommendations for Stage 2 patients:
If the patient is hypokalemic, then potassium gluconate or potassium citrate should be supplemented to effect (typically 1-2 mmol/kg/day).
If serum or plasma SDMA is >25 µg/dl in a Stage 2 patient, consider the suitability of treatments recommended for Stage 3 patients.

Stage 3 Feline patients:
The range of presentations for cats in Stage 3 is likely to be wide, from no clinical signs to quite marked extra-renal clinical signs. The main treatments mentioned so far for stages 1 and 2 are aimed at slowing progression of CKD also apply in stage 3 and may be the only therapies needed for cats with no or mild extra-renal signs. However, treatments designed to improve the quality of life of the cat become more important, the more extra-renal signs are present. These include treatments to address dehydration, nausea and vomiting, anemia and acidosis.
Treatments include all of the steps listed for Stage 1 and 2 plus any additional steps indicated.
1. Discontinue all potentially nephrotoxic drugs if possible.
2. Identify and treat any pre-renal or post-renal abnormalities.
3. Rule out any treatable conditions like pyelonephritis (any urinary tract infection should be regarded as a potential pyelonephritis and treated appropriately) and renal urolithiasis with radiographs and/or ultrasonography.
4. Measure blood pressure and urine protein to creatinine ratio (UP/C).
5. Feed a clinical renal diet.

Management of dehydration:
These patients have decreased urine concentrating ability and therefore
- correct clinical dehydration/hypovolemia with isotonic, polyionic replacement fluid solutions (e.g., lactated Ringer’s) IV or SQ as needed.
- have fresh water available at all times for drinking.

Systemic hypertension:
The blood pressure above which progressive renal injury may be induced is unknown. Our goal is to reduce systolic blood pressure to <160 mm Hg and to minimize the risk of extra-renal target organ damage (CNS, retinal, cardiac problems/damage). If there is no evidence of this but systolic blood pressure persistently exceeds 160 mm Hg,
increasing the risk of extra-renal target organ damage, treatment should be instituted. ‘Persistence’ of increase in systolic blood pressure should be judged on multiple measurements made over the following time-scales in these blood pressure substages:

- Hypertensive (moderate risk of future target organ damage) – systolic blood pressure 160 to 179 mm Hg measured over 1 to 2 months.

- Severely hypertensive (high risk of future target organ damage) – systolic blood pressure ≥180 mm Hg over 1 to 2 weeks.

If evidence of target organ damage exists, cats should be treated without the need to demonstrate persistently increased systolic blood pressure. Reducing blood pressure is a long term aim in managing the patient with CKD and a gradual and sustained reduction should be the goal, avoiding any sudden or severe decreases leading to hypotension.

A logical stepwise approach to managing hypertension is as follows:

1. Dietary sodium (Na) reduction – there is no evidence that lowering dietary Na will reduce blood pressure. If dietary Na reduction is attempted, it should be accomplished gradually and in combination with pharmacological therapy.
2. Calcium channel blocker (CCB) such as amlodipine (0.125 to 0.25 mg/kg once daily).
3. Double the dose of amlodipine (0.25 to 0.5 mg/kg once daily).
4. Combine an inhibitor of the renin-angiotensin-aldosterone system (RAAS; either an angiotensin converting enzyme inhibitor [ACEI, such as benazepril] or an angiotensin receptor blocker [ARB, such as telmisartan]) with the CCB.

Note: Take care not to introduce CCB/RAAS inhibitor treatment to unstable dehydrated cats as glomerular filtration rate may drop precipitously if these drugs are introduced before the patient is adequately hydrated

**Monitoring response to antihypertensive treatment:**

Hypertensive cats normally require lifelong therapy and may require treatment adjustments. Serial monitoring is essential. After stabilization, monitoring should occur at least every 3 months.

Systolic blood pressure <120 mm Hg and/or clinical signs such as weakness or tachycardia indicate hypotension, which is to be avoided.

Blood creatinine concentration – reducing blood pressure may lead to small and persistent increases in creatinine concentration (<45 µmol/l or 0.5 mg/dl increase), but a marked increase suggests an adverse drug effect. Progressively increasing concentrations indicate progressive kidney damage/disease.

**Proteinuria:**

Cats in Stage 3 with urine protein to creatinine ratio (UP/C) >0.4 should be investigated for disease processes leading to proteinuria (see 1 and 2 below) and treated with anti-proteinuric measures (see 3 and 4 below).
Those with borderline proteinuria (0.2 to 0.4) require close monitoring (see 1 and 4 below).

1. Look for any concurrent associated disease process that may be treated/corrected.

2. Consider kidney biopsy as a means of identifying underlying disease (see Appendix and/or consult experts if unsure of indications for kidney biopsy).

3. Administer an RAAS inhibitor (ACEI or ARB) and feed a clinical renal diet.

4. Monitor response to treatment / progression of disease:
   - stable blood creatinine concentration and decreasing UP/C = good response.
   - serially increasing creatinine concentrations and/or increasing UP/C = disease is progressing.

Ordinarily therapy will be maintained lifelong unless the underlying disease has been resolved in which case dose reduction whilst monitoring UP/C might be considered.

Note:

a. Use of an RAAS inhibitor is contraindicated in any animal that is clinically dehydrated and/or is showing signs of hypovolemia. Correct dehydration before using these drugs otherwise glomerular filtration rate may drop precipitously.

b. Cats with proteinuria and hypoalbuminemia likely share the same thromboembolic risk as dogs, however, aspirin is difficult to use in cats to achieve a selective antiplatelet effect. A suggested dose rate if plasma albumin is below 20 g/l (2 g/dl) is 1 mg/kg every 72 hours.

c. Cats with serum phosphate within the IRIS target may be at increased risk of developing hypercalcemia when renal diets are introduced. Monitor serum calcium and if total calcium exceeds 12 mg/dl (3 mmol/l) switch to a senior diet or mix renal diet (50:50 by volume) with standard grocery food.

d. If borderline proteinuria is persistent, antiproteinuric treatment could be offered. The rationale for doing so is that there is an association between progression of CKD and proteinuria which extends into the borderline category. However, there is at present no evidence that intervention with anti-proteinuric drugs slows progression.

**Reduction of phosphate intake:**

Evidence suggests that chronic reduction of phosphate intake to maintain a plasma phosphate concentration below 1.5 mmol/l (but not less than 0.9 mmol/l; <4.6 mg/dl but >2.7 mg/dl) is beneficial to patients with CKD.

A more realistic post-treatment target plasma phosphate concentration for cats at Stage 3 is <1.6 mmol/l (5.0 mg/dl).

The following measures can be introduced sequentially in an attempt to achieve this:

1. Dietary phosphate restriction (i.e., clinical renal diet therapy).

2. If plasma phosphate concentration remains above 1.6 mmol/l (4.6 mg/dl) after dietary restriction, give enteric phosphate binders (such as aluminum hydroxide, aluminum carbonate, calcium carbonate, calcium acetate, lanthanum carbonate) to effect, starting at 30-60 mg/kg/day in divided doses to be mixed with each meal (mixed with the food). The dose required will vary according to the amount of phosphate being fed and the IRIS stage. Treatment with phosphate binders should be to effect (as outlined above), with signs of toxicity limiting the upper dose rate possible in a given patient. Monitor serum calcium and phosphate concentrations
every 4-6 weeks until stable and then every 12 weeks. Microcytosis and/or generalized muscle weakness suggests aluminum toxicity if using an aluminum containing binder – switch to another form of phosphate binder should this occur. Hypercalcemia should be avoided – combinations of aluminum and calcium containing phosphate binders may be necessary in some cases.

3. Although there is evidence that judicious use of calcitriol prolongs survival in dogs at IRIS Stage 3, beneficial effects of ultra-low dose calcitriol have not yet been established in cats.

**Metabolic acidosis:**

If metabolic acidosis exists (blood bicarbonate or total CO₂ <16 mmol/l) once the patient is stabilized on the diet of choice, supplement with oral sodium bicarbonate, (or potassium citrate if hypokalemic) to effect to maintain blood bicarbonate / total CO₂ in the range of 16-24 mmol/l.

**Additional recommendations for Stage 3 patients:**

1. If the patient is hypokalemic, then potassium gluconate or potassium citrate should be supplemented to effect (typically 1-2 mmol/kg/day).

2. Treat vomiting / decreased appetite / nausea / weight loss with an antiemetic / appetite stimulant / antinausea agent (such as maropitant, ondansetron or mirtazapine). Evidence suggests mirtazapine (1.88 mg/cat every 48 h for 3 weeks) reduces vomiting, improves appetite and leads to weight gain in cats showing signs of inappetence and vomiting in this stage. Maropitant (1 mg/kg daily for 2 weeks) reduced vomiting but did not lead to weight gain/ improved appetite. Further investigations are needed on the use of these and other drugs to determine whether they are useful for managing gastrointestinal disturbances in cats with CKD and uremia when administered longer term.

3. Give appropriate maintenance fluids parenterally as necessary to maintain hydration (see Footnote).

4. If serum or plasma SDMA is >45 µg/dl in a Stage 3 patient, consider the suitability of treatments recommended for Stage 4 patients.

Drugs that rely predominantly on renal function for their clearance from the body should be used with caution in patients in Stage 3 CKD. It may be necessary to adjust the dose of these drugs (depending on their therapeutic indices) to avoid accumulation.
Stage 4 Feline patients:

Most cats with stage 4 CKD have many extra-renal signs present. Although it is still important to administer treatments which slow progression of CKD, the importance of improving quality of life for these dogs is greater at this stage. Symptomatic therapy to improve quality of life includes management of dehydration, acidosis, vomiting and nausea and anemia.

Treatments include all of the steps listed for listed for Stages 1, 2 and 3 plus any additional steps indicated.

1. Discontinue all potentially nephrotoxic drugs if possible.
2. Identify and treat any pre-renal or post-renal abnormalities.
3. Rule out any treatable conditions like pyelonephritis (any urinary tract infection should be regarded as a potential pyelonephritis and treated appropriately) and renal urolithiasis with radiographs and/or ultrasonography.
4. Measure blood pressure and urine protein to creatinine ratio (UP/C).
5. Feed a clinical renal diet.

Management of dehydration:

These patients have decreased urine concentrating ability and therefore
- correct clinical dehydration/hypovolemia with isotonic, polyionic replacement fluid solutions (e.g., lactated Ringer’s) IV or SQ as needed.
- have fresh water available at all times for drinking.

Systemic hypertension:

The blood pressure above which progressive renal injury may be induced is unknown. Our goal is to reduce systolic blood pressure to <160 mm Hg and to minimize the risk of extra-renal target organ damage (CNS, retinal, cardiac problems/damage). If there is no evidence of this but systolic blood pressure persistently exceeds 160 mm Hg, increasing the risk of such damage, treatment should be instituted.

‘Persistence’ of increase in systolic blood pressure should be judged on multiple measurements made over the following time-scales in these blood pressure substages:

- Hypertensive (moderate risk of future target organ damage) – systolic blood pressure 160 to 179 mm Hg measured over 1 to 2 months.
- Severely hypertensive (high risk of future target organ damage) – systolic blood pressure ≥180 mm Hg over 1 to 2 weeks

If evidence of target organ damage exists, cats should be treated without the need to demonstrate persistently increased systolic blood pressure. Reducing blood pressure is a long term aim in managing the patient with CKD and a gradual and sustained reduction should be the goal, avoiding any sudden or severe decreases leading to hypotension.

A logical stepwise approach to managing hypertension is as follows:

1. Dietary sodium (Na) reduction - there is no evidence that lowering dietary Na
will reduce blood pressure. If dietary Na reduction is attempted, it should be accomplished gradually and in combination with pharmacological therapy.

2. Calcium channel blocker (CCB) such as amlodipine (0.125 to 0.25 mg/kg once daily).

3. Double the dose of amlodipine (0.25 to 0.5 mg/kg once daily).

4. Combine an inhibitor of the renin-angiotensin-aldosterone system (RAAS; either an angiotensin converting enzyme inhibitor [ACEI, such as benazepril] or an angiotensin receptor blocker [ARB, such as telmisartan]) with the CCB.

Note: Take care not to introduce CCB/RAAS inhibitor treatment to unstable dehydrated cats as glomerular filtration rate may drop precipitously if these drugs are introduced before the patient is adequately hydrated.

**Monitoring response to antihypertensive treatment:**
Hypertensive cats normally require lifelong therapy and may require treatment adjustments. Serial monitoring is essential. After stabilization, monitoring should occur at least every 3 months.

Systolic blood pressure <120 mm Hg and/or clinical signs such as weakness or tachycardia indicate hypotension, which is to be avoided.

Blood creatinine concentration – reducing blood pressure may lead to small and persistent increases in creatinine (<45 µmol/l or 0.5 mg/dl increase), but a marked increase suggests an adverse drug effect. Progressively increasing concentrations indicate progressive kidney damage/disease.

**Proteinuria:**
Cats in Stage 4 with urine protein to creatinine ratio (UP/C) >0.4 should be investigated for disease processes leading to proteinuria (see 1 and 2 below) and treated with anti-proteinuric measures (see 3 and 4 below).

Those with borderline proteinuria (0.2 to 0.4) require close monitoring (see 1 and 4 below).

1. Look for any concurrent associated disease process that may be treated/corrected.

2. Consider kidney biopsy as a means of identifying underlying disease (see Appendix and/or consult experts if unsure of indications for kidney biopsy).

3. Administer an RAAS inhibitor (ACEI or ARB) and feed a clinical renal diet.

4. Monitor response to treatment / progression of disease:
   - stable blood creatinine concentration and decreasing UP/C = good response.
   - serially increasing creatinine concentrations and/or increasing UP/C = disease is progressing.

Ordinarily therapy will be maintained lifelong unless the underlying disease has been resolved, in which case dose reduction whilst monitoring UP/C might be considered.

Note:

a. Use of an RAAS inhibitor is contraindicated in any animal that is clinically dehydrated and/or is showing signs of hypovolemia. Correct dehydration before using these drugs otherwise glomerular filtration rate may
drop precipitously.

b. Cats with proteinuria and hypoalbuminemia likely share the same thromboembolic risk as dogs, however, aspirin is difficult to use in cats to achieve a selective antiplatelet effect. A suggested dose rate if plasma albumin is below 20 g/l (2 g/dl) is 1 mg/kg every 72 hours.

c. Cats with serum phosphate within the IRIS target may be at increased risk of developing hypercalcemia when renal diets are introduced. Monitor serum calcium and if total calcium exceeds 12 mg/dl (3 mmol/l) switch to a senior diet or mix renal diet (50:50 by volume) with standard grocery food.

d. If borderline proteinuria is persistent, antiproteinuric treatment could be offered. The rationale for doing so is that there is an association between progression of CKD and proteinuria which extends into the borderline category. However, there is at present no evidence that intervention with anti-proteinuric drugs slows progression.

Reduction of phosphate intake:
Evidence suggests that chronic reduction of phosphate intake to maintain a plasmaphosphate concentration below 1.5 mmol/l (but not less than 0.9 mmol/l; <4.6 mg/dl but >2.7 mg/dl) is beneficial to patients with CKD.

A more realistic post-treatment target plasma phosphate concentration for cats at Stage 4 is <1.9 mmol/l (6.0 mg/dl).

The following measures can be introduced sequentially in an attempt to achieve this:

1. Dietary phosphate restriction (i.e., clinical renal diet therapy).

2. If plasma phosphate concentration remains above 1.9 mmol/l (6.0 mg/dl) after dietary restriction, give enteric phosphate binders (such as aluminum hydroxide, aluminum carbonate, calcium carbonate, calcium acetate, lanthanum carbonate) to effect, starting at 30-60 mg/kg/day in divided doses to be mixed with each meal (mixed with the food). The dose required will vary according to the amount of phosphate being fed and the stage of CKD. Treatment with phosphate binders should be to effect (as outlined above), with signs of toxicity limiting the upper dose rate possible in a given patient. Monitor serum calcium and phosphate concentrations every 4-6 weeks until stable and then every 12 weeks. Microcytosis and/or generalized muscle weakness suggests aluminum toxicity if using an aluminum containing binder – switch to another form of phosphate binder should this occur. Hypercalcemia should be avoided – combinations of aluminum and calcium containing phosphate binders may be necessary in some cases.

3. Although there is evidence that judicious use of calcitriol prolongs survival in dogs at IRIS Stage 4, beneficial effects of ultra-low dose calcitriol have not yet been established in cats.
Metabolic acidosis:
If metabolic acidosis exists (blood bicarbonate or total CO₂ <16 mmol/l) once the patient is stabilized on the diet of choice, supplement with oral sodium bicarbonate (or potassium citrate if hypokalemic) to effect to maintain blood bicarbonate / total CO₂ in the range of 16-24 mmol/l.

Additional recommendations for Stage 4 patients:
1. If the patient is hypokalemic, then potassium gluconate or potassium citrate should be supplemented to effect (typically 1-2 mmol/kg/day).
2. Consider treatment for anemia if it is affecting the patient’s quality of life: typically this occurs when the PCV is <0.20 l/l (20%). Human recombinant erythropoietin is the most effective treatment but is not approved for veterinary use: darbepoetin is preferable as it is less antigenic than epoetin alfa. Anabolic steroids are of no proven benefit and may be detrimental.
3. Treat vomiting / decreased appetite / nausea / weight loss with an antiemetic/appetite stimulant / antinausea agent (such as maropitant, ondansetron or mirtazapine). Evidence suggests mirtazapine (1.88 mg/cat every 48 h for 3 weeks) reduces vomiting, improves appetite and leads to weight gain in cats showing signs of inappetence and vomiting in this stage. Maropitant (1 mg/kg daily for 2 weeks) reduced vomiting but did not lead to weight gain/ improved appetite. Further investigations are needed on the use of these and other drugs to determine whether they are useful for managing gastrointestinal disturbances in cats with CKD and uremia when administered longer term.
4. Give appropriate maintenance fluids parenterally as necessary to maintain hydration (see Footnote).
5. Intensify efforts to prevent protein / calorie malnutrition. Consider feeding tube intervention (e.g., percutaneous gastrostomy tube).
6. Intensify efforts to prevent dehydration. Feeding tubes can be used to administer fluids as well as food.
7. Consider dialysis and/or renal transplantation.
Drugs that rely predominantly on renal function for their clearance from the body should be used with caution in patients in Stage 4 CKD. It may be necessary to adjust the dose of these drugs (depending on their therapeutic indices) to avoid accumulation.
Appendix

Reasons for Undertaking Renal Biopsy

1. Renomegaly
2. CKD in a young patient
3. Persistent and severe proteinuria (UP/C > 2.0) in a non-azotemic patient
4. Worsening proteinuria in a CKD patient
5. Acute kidney injury, where renal biopsy may provide a prognostic indicator

Footnote

Maintenance fluids to maintain hydration status are low in sodium (30-40 mmol/l) and ideally have added potassium (about 13 mmol/l) to ensure daily requirements for fluid and electrolytes are met (e.g. Normosol-M® or 5% Dextrose plus 0.18% NaCl with added KCl).

Disclaimer

Although every effort has been made to ensure the completeness and accuracy of the information provided herein, neither the IRIS Board nor Elanco Animal Health assumes any responsibility for the completeness or accuracy of the information. All information is provided “as is” without any warranties, either expressed or implied.