

1 **A CASE OF RENAL AMYLOIDOSIS ASSOCIATED WITH UDDER CLEFT DERMATITIS**
2 **IN AN ADULT DAIRY COW**

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10 **SUMMARY**

11 Amyloidosis is a group of disorders characterised by the injurious deposition of abnormal proteins in
12 tissues. Most common in cattle and other animals is secondary or reactive systemic amyloidosis
13 associated with chronic inflammation, resulting in deposition of acute-phase serum amyloid A (SAA)
14 in organs like the kidneys, liver, and spleen. While chronic diseases like mastitis, metritis and
15 pododermatitis are usually the most common diseases identified as inflammatory sources, any
16 persistent inflammation can trigger this disorder. Cattle affected by amyloidosis often exhibit
17 symptoms like weight loss and kidney dysfunction. Here we present the case of a five-year-old
18 Holstein Friesian referred to the University of Glasgow for weight loss and chronic diarrhoea. Clinical
19 examination revealed low body condition score, watery diarrhoea, mild dehydration (5%), mild
20 submandibular oedema, left renomegaly, and udder cleft dermatitis. Biochemistry and urine analysis
21 indicated hypoalbuminemia with normal globulin levels and marked proteinuria. Given the poor
22 prognosis, the animal was euthanised on welfare grounds. Gross *post-mortem* findings suggested a
23 diagnosis of secondary amyloidosis, and histopathology confirmed SAA deposition in the glomeruli
24 and renal medullary interstitium. In the absence of another grossly appreciable chronic inflammatory
25 focus/ foci the udder cleft dermatitis was considered the likely contributing comorbidity.

26 **KEYWORDS**

27 Bovine; renal disease; amyloidosis; chronic inflammation; udder cleft dermatitis.

28 **INTRODUCTION**

29 Renal amyloidosis in cattle is uncommon, with abattoir reports ranging from 0.8 % to 5% (1,2).
30 Amyloidosis is associated with chronic inflammatory diseases, such as reticuloperitonitis or any other
31 long-term inflammation causing an elevated concentration of serum amyloid A (SAA), which can
32 form aggregates that are systemically deposited, predominantly in the kidney, liver, and spleen (2,3).
33 Even if the disease can affect different organs, the clinical signs draw attention when the kidneys are
34 involved (4,5). There are no specific preventative measures for amyloidosis and the prognosis is
35 usually poor (6).

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37 **CASE HISTORY**

38 A 5-year-old Holstein Friesian cow was referred on 24th February 2021 to the University of Glasgow
39 (UofG) School of Veterinary Medicine. The cow was 164 days in milk in her third lactation.
40 According to the farmer's history, the cow had a mild milk drop (from 44 to 40 litres) in December
41 2020. On 9th February 2021, a significant milk drop (from 38 to 26 litres) associated with diarrhoea
42 was noticed. The referring veterinary surgeon reported profuse diarrhoea, left renomegaly, and low
43 body condition score. The cow was treated on farm with a three-day course of 15 mg/Kg bodyweight
44 trimethoprim 40 mg – 200 mg sulfadiazine, IM, SID (Norodine®, Trimethoprim 4 mg/ml
45 Sulfadiazine 20 mg/ml, Norbrook, UK); with no clinical improvement.

46 The cow was referred from a dairy farm milking 200 pedigree Holstein Friesian cows and averaging
47 12.500 kg per cow/year. Udder cleft dermatitis was reported sporadically on the farm (prevalence <
48 3 %). The farm was closed with biosecurity measures in place for visitors and contractors. A control
49 plan was in place for paratuberculosis with all milking cows tested quarterly for antibodies in milk.
50 All tests were negative in the previous 2 years. The herd had Bovine Viral Diarrhoea (BVD) negative
51 status. Vaccinations against BVD, Leptospirosis, Infectious Bovine Rhinotracheitis and mastitis
52 (*Escherichia coli-Staphylococcus aureus*) were performed yearly. Adult cattle were kept indoors all

53 year round in cubicles, with some cows spending 2-3 weeks on pasture during the dry period. *Fasciola*
54 *hepatica* was absent based on bulk tank mild antibody testing and abattoir reports.

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56 **CASE PRESENTATION**

57 On the day of the admission, the cow appeared bright, alert and responsive. Her Body Condition
58 Score (BCS) was 2.00 (range 0-5; bodyweight 622 Kg (7)). A greenish and watery diarrhoea without
59 visible blood was observed. Normal spontaneous micturition was observed, resulting in the voiding
60 of pale yellow transparent urine. Mild oedema was observed in the submandibular area but not in the
61 brisket. No jugular venous distension was observed. Mild dehydration was observed (5%). The
62 mucous membranes were pink and moist and capillary refill time was less than 2 seconds.
63 Examination of the oral cavity revealed no abnormalities. Palpable lymph nodes were normal in size
64 and shape. The heart rate was 60 bpm (reference range: 40-80 (8)); there were no anomalies in
65 frequency and rhythm. The respiratory rate was 30 bpm (reference range: 12-36 bpm (8)); no
66 adventitious sounds were auscultated. Auscultation and percussion of both right and left abdomen
67 sides revealed no abnormal sounds and rumination was regular (1 contraction every 40 seconds). No
68 evidence of thoracic or abdominal pain was present on the “withers test”. Rectal temperature was
69 38.0 °C (reference range: 38-39 °C (8)). On the trans-rectal examination, the left kidney was
70 moderately enlarged but not painful on palpation. Palpation of other organs was unremarkable
71 (rumen, intestines, sub-aortic lymph nodes, urinary bladder, uterus). A focal 10 cm diameter light red
72 ulcerative lesion typical of a healing udder cleft dermatitis lesion on the skin of the ventral midline
73 between the front udder quarters. Palpation of the rest of the udder was unremarkable and the milk
74 was normal in colour and consistency. The California Mastitis Test (CMT) was negative for all
75 quarters (9).

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77 **PROBLEM LIST**

- 78 • Milk drop during mid-lactation
- 79 • Low BCS (2.00 out of 5)
- 80 • Chronic marked watery diarrhoea
- 81 • Enlarged and non-painful left kidney per rectum
- 82 • Mild submandibular oedema
- 83 • Mild dehydration (5%)
- 84 • Udder cleft dermatitis

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86 **DIFFERENTIAL DIAGNOSES**

87 From most to least likely:

- 88 • Urinary tract diseases: Renal amyloidosis, Glomerulonephritis (GN), Pyelonephritis,
89 Hydronephrosis, Cystitis.
- 90 • Protein-losing enteropathies: Paratuberculosis (*Mycobacterium avium* sub *paratuberculosis*-
91 MAP), Gastrointestinal nematodes, Chronic salmonellosis.
- 92 • Hepatic failure: Fasciolosis.
- 93 • Heart failure: Congestive heart failure.

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95 **ANCILLARY TESTING**

96 A blood sample for complete haematology and biochemistry was taken on 25th February 2021, and
97 the results are reported in Table 1. The haematology was unremarkable. Biochemical features
98 included hypoalbuminemia (19 g/L (31-38 g/L)) and mild hypoproteinaemia (53 g/L (63-89 g/L)).
99 Globulins were within the normal range (34 g/L (30-48 g/L)).

100 A urine sample was collected by catheterisation of the urethra after careful disinfection of the vulva
101 to evaluate renal functionality (Table 2). The urine sample was analysed with a dipstick (Multistix
102 SG (Bayer)); urine pH was 8.5, and marked proteinuria (+4) was observed. Proteinuria was confirmed

103 with sulfosalicylic acid precipitation; some white blood cells (WBCs) were observed on microscopy
104 (Table 2).

105 Both kidneys were examined by ultrasonography: the left one by trans-rectal ultrasonography (7.5
106 MHz linear probe) and the right one trans-abdominally (2.5-5 MHz convex probe) in the dorsal part
107 of the right paralumbar fossa (10). The ultrasonography of the left kidney showed renomegaly (15 x
108 35 cm). Similarly, the right kidney was evaluated trans-cutaneously, and renomegaly (15 x 35 cm)
109 was observed; no other abnormalities, such as flocculent fluid or deformed sinuses, were observed
110 (11,12). Ultrasonography of the bladder (7.5 MHz linear probe) was unremarkable. Ultrasonography
111 of the heart (2.5-5 MHz convex probe) was unremarkable. Other ancillary testing, such as faecal
112 culture, *Mycobacterium avium* sub *paratuberculosis* - PCR, and parasitology, did not highlight any
113 relevant findings (Table 2).

114 During the hospitalisation, which lasted three weeks, weekly haematology and blood biochemistry
115 was carried out. Hypoalbuminaemia (in the range of 18-20 g/L (31-38 g/L)) was stable, and no further
116 abnormalities were detected.

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118 **DIAGNOSIS**

119 Based on the clinical examination and ancillary testing results, a non-infectious protein-losing kidney
120 disease was diagnosed. The gold standard to differentiate between renal amyloidosis and
121 glomerulonephritis *in vivo* would have been renal biopsy (5). Considering the poor prognosis for the
122 mentioned diseases, it was decided to euthanise the animal using 140 mg/kg, Pentobarbital Sodium,
123 IV (Euthasol® Vet. 400 mg/ml, Dechra, UK). A *post-mortem* examination was performed.

124 On the gross pathology, both kidneys were enlarged (Figure 1.1). The ventral subcutis was moderately
125 expanded by clear to pale yellow gelatinous material (oedema). In the large intestines, the wall was
126 markedly expanded by clear to pale yellow gelatinous material (oedema). On the skin of the udder, a
127 lesion of approximately 10 cm diameter was observed between the two front quarters (Figure 1.2).

128 On histopathology of the kidney, the glomeruli and interstitium of the corticomedullary and
129 medullary regions were expanded by moderate to large amounts of pale eosinophilic homogenous
130 material that on thick sections displays moderate to marked congophilia (Figure 2.1) with apple-green
131 birefringence in polarised light (Figure 2.2) confirming that this was SAA substance. The
132 histopathology confirmed the SAA deposition in the glomeruli and renal interstitium of the medulla
133 (4). A final diagnosis of renal amyloidosis was made.

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135 **DISCUSSION**

136 Typical clinical signs of renal amyloidosis, such as chronic diarrhoea, weight loss, inappetence,
137 peripheral oedema and reduced milk production, were potentially indicative of a wide range of
138 disorders and, therefore, of low specificity. Renomegaly is more indicative of kidney disease, but it
139 is a subjective evaluation, and when dilatation is subtle, it can be easily missed at trans-rectal
140 palpation. Blood chemistry indicated hypoalbuminemia, suggesting a protein-losing disease rather
141 than congestive heart failure as the cause of submandibular oedema. Heart diseases were then ruled
142 out by ultrasonographic examination. Protein loss can be caused by kidney diseases, enteropathy or
143 hepatic failure (5). Interestingly, the albumin level did not diminish below 10 g/L, which is considered
144 the cut-off to create a change in the oncotic pressure to produce significant generalised oedema (5).
145 Protein-losing enteropathies (such as paratuberculosis, gastrointestinal nematodes, and chronic
146 salmonellosis) and fasciolosis can cause a clinical picture similar to the present case (5,13). These
147 diseases were ruled out by ancillary testing. Faecal culture isolated *Campylobacter jejuni*, which was
148 considered to be a secondary finding resulting from an opportunistic infection. In a study conducted
149 by Wesley and collaborators, faecal samples were collected from 2,085 dairy cattle across 31 farm
150 operations in the USA. *Campylobacter jejuni* was identified in healthy cows without any clinical
151 signs (14). Having ruled out protein-losing enteropathy and liver failure, protein-losing nephropathy
152 was considered the primary cause of the clinical signs.

153 Urinalysis was shown to be a useful tool in narrowing the diagnostic suspicion. The urinary dipstick
154 showed marked proteinuria, which was indicative of kidney damage. It is worth remembering the

155 importance of specifically testing proteinuria, like this case, with sulfosalicylic acid precipitation. In
156 fact, alkaline urine pH can give a false positive result and lead to a misinterpretation (5). Among
157 cattle diseases, pyelonephritis is the most common (5). Urinalysis revealed a lack of other evidence
158 consistent with urinary tract inflammation and/ or infection. The small numbers of white blood cells
159 observed in the present case was considered physiological based on the method of urine collection
160 and the mixed culture of bacteria isolated was consistent with contamination. On urine culture, there
161 was no evidence of *Corynebacterium renale* or *Escherichia coli*, which are commonly isolated in
162 cases of pyelonephritis; the other isolated ones were considered non-pathogenic. These findings were
163 supported by clinical examination, ancillary testing, and post-mortem findings. At clinical
164 examination, no signs of pain such as stranguria or renal pain at palpation were detected, and
165 proteinuria in the absence of significant leukocyturia or renal pain made significant pyelonephritis
166 unlikely, like a bladder infection. Ultrasonographically, no signs of infection were detected in the
167 kidney and bladder. Finally, post-mortem and histopathology did not reveal any evidence of
168 pyelonephritis or cystitis, completely ruling out urinary tract infections. These findings were
169 consistent with non-infectious protein-losing nephropathy, such as amyloidosis and
170 glomerulonephritis. Clinically significant glomerulonephritis, such as renal amyloidosis, is rarely
171 reported in cattle (5). Further pathological findings confirmed the presence of serum amyloid A in
172 both glomeruli and renal interstitial spaces, confirming the renal amyloidosis diagnosis (4).
173 Considering no other sources of chronic inflammation were identified, udder cleft dermatitis was
174 thought to be the inciting cause of the renal amyloidosis in this case (15).

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176 **CONCLUSION**

177 For the first time, we report a case of renal amyloidosis in which udder cleft dermatitis was considered
178 the causative factor in the absence of another identifiable inflammatory foci (15). There are no
179 specific preventative measures for amyloidosis, and it was not considered relevant at the herd level.
180 The present case highlights the importance of a comprehensive diagnostic approach, including
181 history, clinical examination, ancillary testing, and post-mortem, to achieve the correct diagnosis of
182 renal amyloidosis. Additionally, the causative inflammatory focus was identified by careful exclusion
183 criteria.

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185 **Acknowledgements**

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187 at the University of Glasgow without whom investigation of this case would not have been possible.

188

189 **Author Contributions**

190 Giovanni Capuzzello: Resources, Conceptualization; Investigation; Visualization; Data Curation;
191 Writing - original draft.

192 Isabella Nicola: Conceptualization; Writing - review & editing.

193 Alexander Gray: Conceptualization; Investigation; Methodology; Data Curation; Writing - review &
194 editing.

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196 **Conflict of Interest**

197 The authors declare no conflict of interest.

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TABLES AND FIGURES

Table 1 - Haematological and biochemical parameters of a 5 year old cow referred for diarrhoea, wasting condition and renomegaly to theUofG

Parameter	Result	Reference Interval
RBC	6.35	5-10 x 10 ¹²
Hb	10.7	8.5-12.2g/dL
HCT	30.4	22-33 %
MCV	47.9	38-50 fl
MCHC	35.2	30-36/dL
WBC	9.72	4.6-12 x 10 ⁹ /L
Neutrophils	3.56	0.6-4.0 x 10 ⁹ /L
Lymphocytes	3.72	1.5-7.5 x 10 ⁹ /L
Monocytes	0.5	0.025-0.84 x 10 ⁹ /L
Eosinophils	0	0-9 x 10 ⁹ /L
Basophils	0.051	0-0 x 10 ⁹ /L
PLT	456	100-800 x 10 ⁹
Toxic neutrophils	Absent	-
Phosphate	1.81	1.13-2.84 mmol/L
Calcium	2.24	2.2-3.3mmol/L
Magnesium	0.54	0.65-1.39 mmol/L
Sodium	137.3	135-157 mmol/L
Potassium	4.8	3.2-5.8 mmol/L
Total protein	53	63-89 g/L
Albumin	19	31-38 g/L
Globulin	34	30-48 g/L
Albumin/globulin ratio	0.56	0.88-1.31
GGT	22	0-27 U/L
GLDH	66.9	0-30 U/L
AST	62	0-140 U/L
ALK phosphatase	37	20-280 U/L
Creatinine	36	113-212 U/L
Urea	2.8	1.6-5.9 mmol/L

238 RBC: Red blood cells. Hb: Haemoglobin. HCT: Haematocrit, MCV: Mean corpuscular volume. MCHC: Mean
 239 corpuscular haemoglobin concentration. WBC: White blood cells. PLT: Platelet. GGT: Gamma-glutamyl transferase.
 240 GLDH: Glutamate dehydrogenase. AST: Aspartate aminotransferase. ALK: Alkaline.
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Table 2 - Other ancillary tests of a 5 years old cow referred for diarrhoea, wasting condition and renomegaly to the UofG

Urine dipstick	Proteinuria (+4): No other abnormalities were detected.		
Urine analysis	Parameter	Results	Reference Interval
	Quantitative Protein	351.7 mg/100ml	0- 25.0
	Qualitative Protein	+++	Considered physiological in ruminants
	Quantitative Creatinine	59.06 mg/100mL	0-142.3
	Protein creatinine ratio (UPC)	5.95	0.04.-0.25
	pH	8.5	-
	WBC (microscopy)	+	-
Urine culture and sensitivity	<i>Corynebacterium</i> Group G (formerly <i>Arcanobacterium haemolyticum</i>), <i>Aerococcus viridans</i> and <i>Bacillus</i> were isolated.		
Faecal culture and sensitivity	<i>Campylobacter jejuni</i> ssp <i>doylei</i> was isolated.		
Paratuberculosis	Serology (Ab ELISA): Negative (1%). Cut-off (P%): Negative <= 50%.		
	PCR: negative.		
Parasitology	McMaster: no nematode eggs detected. Boray (faecal sedimentation): - No liver egg flukes were detected.		

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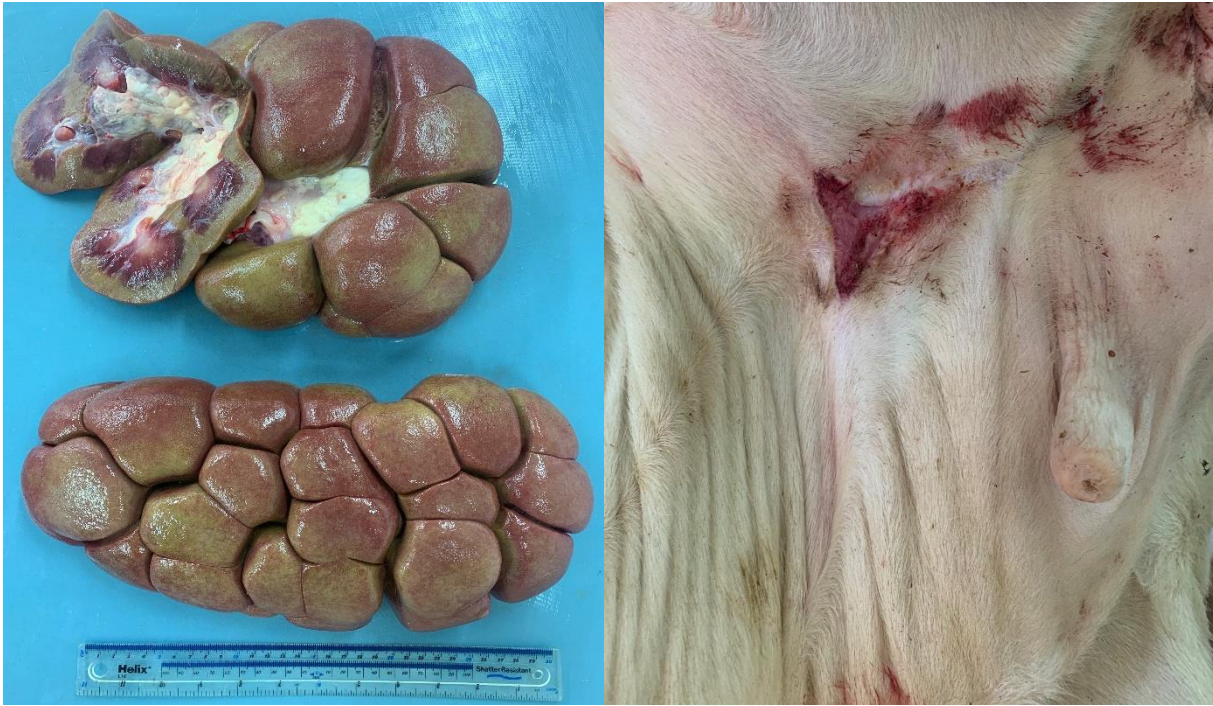


Figure 1.1 Kidneys bilaterally enlarged, measuring 15 x 20 x 35cm (normal size: 14 x 18 x 24), with pale yellow discolouration of the subcapsular surface and cortex.

Figure 1.2 Udder cleft dermatitis. The lesion was irregularly shaped, depressed, well-delineated, and dark red, indicating a chronic ulcerative dermatitis.

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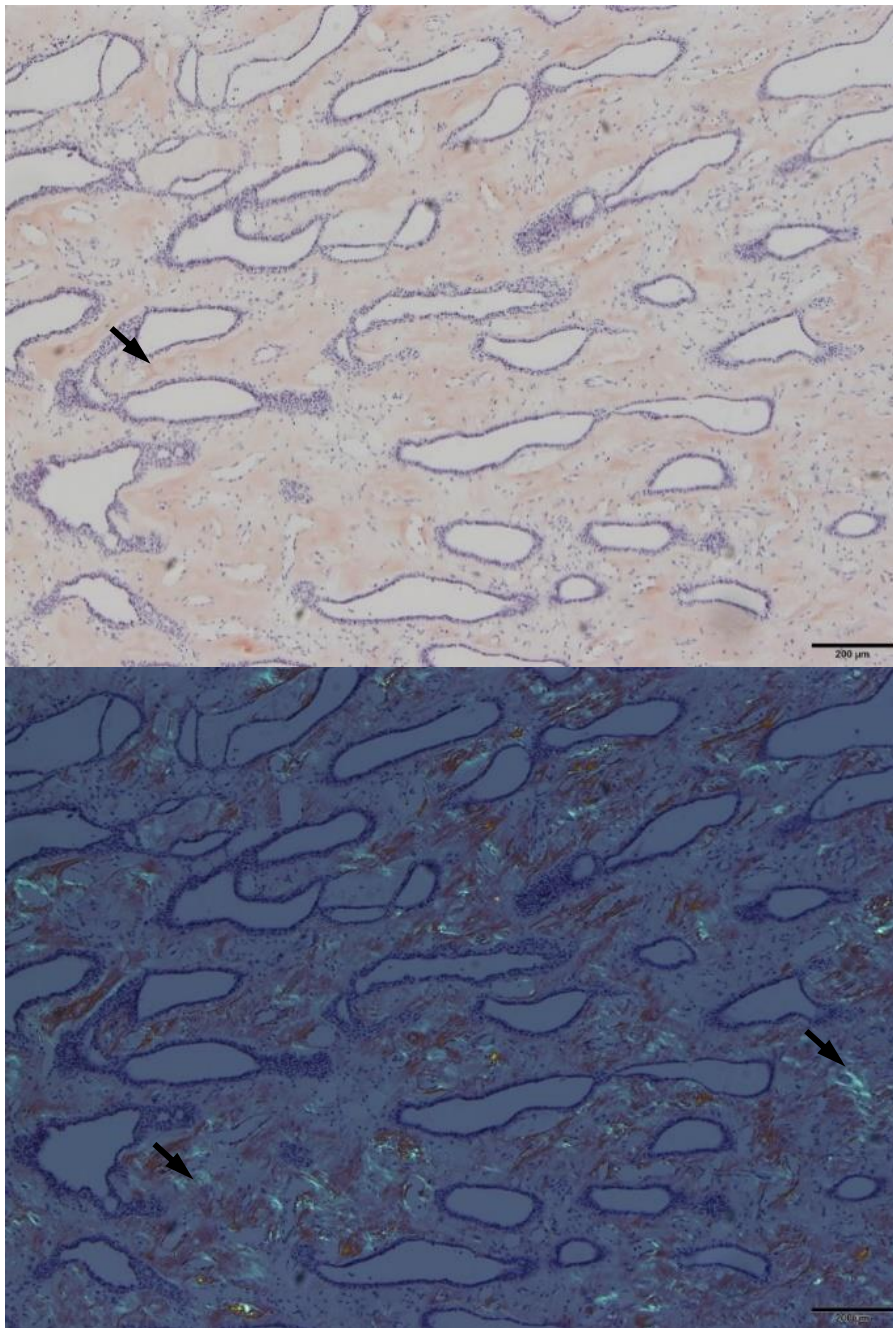


Figure 2.1 Histopathology section of the medulla, kidney. Moderate expansion of the interstitium of the renal medulla by homogeneous amorphous congophilic material suspected to be amyloid (black arrow). Congo red, 6 µm section, standard illumination, x 400, scale bar 200 µm.

Figure 2.2 Histopathology section of the medulla, kidney. Apple green birefringence under polarised light (black arrows) confirms the congophilic material as amyloid. Congo red, 6 µm section, polarised light, x 400, scale bar 200 µm.