A Study into the Identity, Patterns of Infection and Potential Pathological Effects of Rumen Fluke and the Frequency of Co-Infections with Liver Fluke in Cattle and Sheep

Valentina Busin 1,* , Eilidh Geddes 1,2, Gordon Robertson 1, Gillian Mitchell 2, Philip Skuce 2, Katie Waine 3, Caroline Millins 1,4 and Andrew Forbes 1

1 School of Biodiversity, One Health & Veterinary Medicine, University of Glasgow, Glasgow G61 1QH, UK
2 Moredun Research Institute, Pentlands Science Park, Midlothian EH26 0PZ, UK
3 Finn Pathologists, One Eyed Lane, Norfolk IP21 5TT, UK
4 Department of Livestock and One Health, Institute of Infection, Veterinary and Ecological Sciences, University of Liverpool, Liverpool CH64 7TE, UK
* Correspondence: vale.busin@gmail.com

Abstract: Rumen fluke (RF) are trematode parasites that have increased in prevalence within European ruminant livestock since the 1990s. Morbidity and mortality can result from the development of juvenile flukes in the duodenum, however, evidence for significant impacts of adult fluke in the rumen of the final host is equivocal. The presence of rumen fluke in Scotland had not previously been quantified, so the purpose of this study was to use historical coprological data collected between 2008 and 2018 at the School of Veterinary Medicine in Glasgow to evaluate the frequency of rumen fluke infection and to compare this with the presence of liver fluke. This analysis showed evidence of adult rumen fluke in 6.6% of samples submitted, with a substantial and significant increase in positive diagnoses from 2016, following which positive rumen fluke diagnoses equaled or slightly outnumbered those of liver fluke. A prospective post-mortem examination study was also carried out to determine the presence of rumen and liver fluke in cattle and sheep, to quantify adult rumen fluke burdens and to assess any pathological changes in the reticulorumen and proximal duodenum associated with infection. The presence of rumen fluke in post-mortem cases was 26.9% (n = 18/67), the majority (66.7%, n = 12/18) with burdens of less than 100 adult rumen flukes. There was no significant difference in mean ruminal papillar density and length in animals with and without adult rumen fluke and no significant gross pathology was observed in the rumen or reticulum. Examination of animals with adult rumen fluke provided no evidence of any consistent associations with acute or chronic inflammatory changes in the duodenum. All rumen fluke collected at necropsy were identified by PCR and sequencing as Calicophoron daubneyi.

Keywords: rumen fluke; liver fluke; Calicophoron daubneyi; Fasciola hepatica; surveillance; pathology; diagnostics

1. Introduction

Trematodes are important livestock parasites with a worldwide distribution that are responsible for serious economic losses and compromised animal welfare. Two of the most significant trematodes in temperate regions are rumen fluke (RF) and liver fluke, Fasciola hepatica. RF are digenean trematodes belonging to the family Paramphistomidae, which includes several genera such as Calicophoron and Paramphistomum, which commonly infect ruminants worldwide [1]. Adult RF are conically shaped, pink or red in colour and measure between 5–12 mm in length and 2–4 mm in width, and are found primarily within the reticulum and rumen. Juveniles develop in the duodenum and measure 1–2 mm in length [2]. There are over 10 species of paramphistomes known to infect domestic ruminants, however, the species Calicophoron daubneyi has been confirmed as the most
paramphistomes in livestock is largely based upon post-mortem examination (PME) and faecal egg counts (FECs). FECs are useful in detecting infections with adult RF, but not during the pre-patent period when juvenile flukes only are present; that said, juvenile fluke can be detected in faecal samples during acute paramphistomosis [6]. PME of animals for evidence of haemorrhagic duodenitis with immature flukes confirmed on microscopic examination of intestinal scrapes [7] or recovery of adult RF from the rumen and reticulum are regarded as conclusive proof of RF infection [8]. RF have been present in the UK and Ireland for many years, however, their previously low prevalence and perceived low pathogenicity have meant they have received limited attention from veterinary parasitologists and livestock owners [8]. In recent years, the prevalence of RF in continental Europe, Ireland and the UK has increased [9,10], with reports of herd-level prevalence in the UK reaching 59% [11]. Furthermore, serious consequences have been associated with the presence of juveniles in the duodenum, where heavy burdens can result in clinical disease and mortality in both sheep and cattle [12,13]. In juvenile infections there is evidence of erosion, petechiation and necrosis, leading to haemorrhagic enteritis and duodenitis [7]. Little is known, however, about any long-term effects on the duodenal mucosa caused by juvenile RF infection. In endemic areas, it is quite possible that ruminants could play host to both RF and liver fluke simultaneously if they graze on pastures contaminated with metacercariae of both species. Co-infection with C. daubneyi and F. hepatica could also confound the interpretation of clinical, pathological and parasitological observations [14].

While adult RF are still considered largely non-pathogenic, it is known that infections can result in focal chronic ulcerative ruminitis, ruminal papillar atrophy and ulceration at the point of attachment of the adults [15]. The rumen is a complex fermentative organ and infection with C. daubneyi has been shown to change an animal’s volatile fatty acid profile, potentially affecting nutrient absorption and utilisation [16]. Additionally, alterations in papillar shape and length have been observed in clinical cases associated with adult RF [15].

Based on the increase in RF diagnoses in the UK and the relative lack of evidence of the subclinical effects of RF, the aims of this study were the following: (1) to analyse available historical parasitological data and establish the frequency and pattern of RF infection in Central and South West Scotland; (2) to investigate animals submitted for PME for the presence and burden of RF and liver fluke infection; (3) to determine the species of RF most prevalent in samples submitted to the laboratory; (4) to assess anatomical changes in the rumen and duodenum of cattle and sheep affected by adult RF; and (5) to provide a preliminary assessment of the frequency of rumen and liver fluke co-infections.

2. Materials and Methods
2.1. Parasitological Data
At the University of Glasgow Veterinary Diagnostic Service, faecal samples from ruminants, collected for routine analysis of University associated farms (n = 6) and from farm animal clinical cases referred from first opinion veterinary practitioners [17], are routinely tested for evidence of trematode eggs. Faecal samples are tested for both RF and liver fluke eggs using a modified Boray sedimentation technique as follows. A total of 3 grams of faeces were homogenised with 42 mL of tap water and the fluid suspension was passed through a coarse sieve (250 microns) into a 1 L measuring cylinder and allowed to sediment for two minutes. The supernatant was removed and two drops of 5% methylene blue was added to the sediment material. The resultant solution was placed in a lined petri
dish and examined using a stereo microscope at 12× magnification. The total number of eggs in the sample for both RF and liver fluke was recorded from 0 to >25 eggs per gram (epg). After counting 25 eggs, the samples were discarded and a value of >25 epg was recorded. Differentiation of the eggs of rumen and liver fluke was through morphological features and the colour of the eggs.

2.2. Post-mortem Examination (PME)
2.2.1. Presence and Burden of Adult Rumen Fluke

An 8-week study was conducted for two consecutive years (January and February 2018 and 2019). All ovine and bovine cases submitted to the University of Glasgow Veterinary Diagnostic Service for PME during these time periods were inspected for the presence of RF and liver fluke as follows: the forestomachs were thoroughly inspected for the presence of adult RF by gross examination of the contents and the mucosa. If RF were present, the infection burden was classified between 0–4 [18]: 0—no fluke, 1—between 1 and 10, 2—between 11 and 100, 3—between 100 and 200, 4—more than 200 adult rumen fluke. The liver and gall bladder of each case were also examined by longitudinal section for the presence of adult liver fluke. The presence of liver fluke was recorded as either present or absent. All the animals submitted for PME were over 8 months old.

2.2.2. Rumen Fluke Species Identification

During the first year of the project (2018), for each positive case, up to 10 individual adult RF were collected from the rumen and reticulum and stored in 100% ethanol and subsequently processed for species identification [19]. A Qiagen DNEasy Blood and Tissue kit (QIAGEN, Hilden, Germany) was used as detailed in the manufacturing instructions to extract DNA from each individual adult RF [20,21]. The PCR programme was performed using a GeneAMP PCR system 2720 thermal cylinder (Applied Biosystems, Foster City, CA, USA) and the gel was viewed on an Alphamagel Imaging System (Alpha Innotech, San Leandro, CA, USA). The purification of the resultant PCR products achieved using the QIAquick PCR Purification Kit as described by the manufacturer (QIAGEN, Hilden, Germany) and the purified products were sent for Direct Nucleotide Sequencing (Eurofins MWG Operon, Ebersberg, Germany). The resultant sequences were assessed with a Lasergene 10 core suite Software SeqMan Pro (DNASTAR, Madison, WI, USA) and compared to reference sequences from the GenBank database using BLASTn at the European Bioinformatics Institute website (http://www.ebi.ac.uk/, accessed on 22 February 2018).

2.2.3. Anatomical Changes

For the second year of the project, the rumen and duodenum of cattle and sheep submitted for PME were assessed macroscopically (gross examination of ruminal papillae) and microscopically (histopathology of duodenum samples). A 10 cm² square sample of the atrium ruminis (Figure 1A) was obtained from all cases. If the sample contained adult RF, the flukes were carefully removed to permit a blinded analysis. All samples were stapled to a numbered cardboard during fixation in a 10% buffered formalin solution for a minimum of 48 hours. Ruminal papillae measurements carried out were: density (average number of papillae within a plastic template of three squares of 3 × 1 cm size—Figure 1B) and length (average of 10 randomly selected papillae throughout the sample measured in millimetres using a ruler as adapted from [22]).

To examine the duodenum for microscopic changes, a 2.5 cm sample from the proximal duodenum (approximately 5 cm distal to the pylorus) was collected and stored for a minimum of 24 hours in 10% buffered formalin, embedded in paraffin and sectioned at 2 μm, and stained with haematoxylin and eosin for histopathological analysis. The stained slides were then examined at 200× and 400× magnification using an Olympus BX 43 microscope by two veterinary pathologists. The pathologists examining the sections did not have access to the PME findings.
Following initial histopathological examination of the sections, a scoring system was developed, based on standards used in small animal medicine [23] to assess several non-specific microscopic features present: inflammation of the mucosa and submucosa, and evidence of chronic changes such as fibrosis or granulomas (Appendix A).

2.3. Statistical Analysis

All data analysis in this study was performed using R version 4.1.1 and RStudio version 2021.09.0 (R Core Team, 2022; RStudio Team, 2020, Vienna, Austria). Statistical significance was defined at $p$-value $\leq 0.05$.

The relationship between the number of positive RF infections and year was analysed using a logistic regression model. This was then repeated for liver fluke. A Pearson’s Chi-squared test for independence was used to establish whether a difference existed in the overall number of RF infections between bovine and ovine samples. The Pearson’s Chi-squared test was also used to establish whether there was a significant seasonal pattern in RF and liver fluke diagnoses, cumulative across all years.

For the PME data, a generalised linear regression was used to test for a relationship between bovine/ovine and papillar density, then papillar length. Furthermore, the same method was also utilised to test for a relationship between adult RF presence (adult rumen fluke burden score 1–4 combined) and papillar length, then adult RF presence and papillar density.

A binomial generalised linear model with rumen fluke presence (adult rumen fluke burden score 1–4 combined) and absence (score 0) as the explanatory variable was used to assess the relationship between adult RF presence and each of the seven categories of histopathological changes (Appendix A). For category ‘a’ (inflammation of the lamina propria with lymphocytes and plasma cells), classification 1 (0–5 cells between crypts) was compared to classification 2–4 combined (5 or more cells). For all statistical analysis, co-infected animals were included in all analyses where RF presence/absence was tested.

3. Results

3.1. Parasitological Data

The first positive RF egg count recorded was in August 2009 (data recording for trematodes started in 1979), while only F. hepatica eggs were found in previous faecal samples. FECs data were therefore analysed from January 2008 to December 2018. In this period,
3068 ruminant faecal samples were submitted for FECs analysis. A total of 595 samples (19.4%) contained trematode eggs: 124 had only RF eggs (4%) and 394 (12.8%) contained only liver fluke eggs, with 77 (2.5%) positive for both trematodes, giving an overall presence of adult rumen fluke infections of 6.6% (n = 201/3068). Of the 595 trematode egg positive samples, 437 (73.5%) were of known bovine or ovine origin, the remainder were not specified (but were either bovine or ovine). Over the 10-year study, a total of 192 (n = 192/437; 43.9%) known bovine samples and 245 (n = 245/437; 56.1%) known ovine samples submitted were positive for RF.

Considering the total number of samples submitted (n = 3068), diagnoses of rumen fluke increased yearly, with the highest percentage of positive RF diagnoses being recorded in 2016 (n = 104/586; 17.8%). In 2017, the occurrence of RF infection fell to 10.9% (n = 42/386), rising back in 2018 to 17.2% (n = 36/209). There were two notable peaks in liver fluke diagnoses, one in 2013, when the proportion of liver fluke positive samples reached 24.4% (n = 83/340), and another in 2016, reaching 22.7% (n = 133/586). From 2017, the percentage of positive samples for RF exceeded that of liver fluke (Figure 2). There was no statistical difference in RF infection between bovine and ovine samples (p = 0.20; x² = 1.66).

Figure 2. The percentage of faecal samples testing positive for rumen fluke eggs and liver fluke eggs during the study period (2008–2018).

RF seasonal variation in positive samples (Figure 3) displayed the highest proportion of positive samples in winter (33%), and lowest in summer (15%); however, there was no significant statistical difference between number of positive diagnoses made each season (p = 0.14; x² = 5.5). The highest proportion of samples with liver fluke present occurred in autumn (37%). This decreased in winter (28%) and the lowest occurrence was observed in spring (17%). A significant difference in the number of positive liver fluke diagnoses made per season was observed (p = 0.001; x² = 15.8).
significant statistical difference between number of positive diagnoses made each season \((p = 0.14; \chi^2 = 5.5)\). The highest proportion of samples with liver fluke present occurred in autumn (37%). This decreased in winter (28%) and the lowest occurrence was observed in spring (17%). A significant difference in the number of positive liver fluke diagnoses made per season was observed \((p = 0.001; \chi^2 = 15.8)\).

Figure 3. Seasonal distribution of positive faecal samples for rumen fluke \((n = 201)\) and liver fluke \((n = 471)\).

### 3.2. Post-Mortem Examination (PME)

#### 3.2.1. Presence and Burden of Adult Rumen Fluke

A total of 67 animals (38 in year 1 and 29 in year 2) were submitted for PME. Of these animals, 18 (26.9%: 11 in year 1 and seven in year 2) had adult RF present within their forestomach: thirteen bovine (13/30; 43.3%) and five ovine (5/37; 13.5%). Eleven animals (11/67; 16.4%) were positive for liver fluke across both years. Five of these had co-infection with RF (5/67; 7.5%) (four in year 1 and one in year 2). Different burdens of adult RF were found in the RF positive animals \((n = 18)\) during the PME study, with most \((12/18; 66.7\%)\) showing scores below 100 adult RF (Table 1).

#### Table 1. Number of animals (total, bovines and ovines) in relation to adult RF burden.

<table>
<thead>
<tr>
<th>Adult Rumen Fluke Burden Score</th>
<th>Number of Adult Rumen Fluke</th>
<th>Total Number of Infected Animals (Percentage %)</th>
<th>Total Number of Infected Bovines (Percentage %)</th>
<th>Total Number of Infected Ovines (Percentage %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1–10</td>
<td>5 (27.8%)</td>
<td>1 (7.7%)</td>
<td>4 (80%)</td>
</tr>
<tr>
<td>2</td>
<td>11–100</td>
<td>7 (38.9%)</td>
<td>6 (46.2%)</td>
<td>1 (20%)</td>
</tr>
<tr>
<td>3</td>
<td>101–200</td>
<td>3 (16.6%)</td>
<td>3 (23.1%)</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>201+</td>
<td>3 (16.6%)</td>
<td>3 (23.1%)</td>
<td>0</td>
</tr>
</tbody>
</table>

Upon inspection of the animals at PME, adult RF were identified morphologically. The highest densities of RF were generally located in the passage between the rumen and reticulum (Figure 4) and in the atrium ruminis. Adult flukes were attached via their ventral sucker (acetabulum) to ruminal and reticular mucosa, with animals exhibiting higher burdens (score 3 and 4) also presenting with RF within the ruminal and reticular contents. No gross pathological changes were visible.
3.2.2. Rumen Fluke Species Identification

Thirty-three adult RF collected from seven animals (twenty-five samples from five cattle and eight samples from two sheep) during the first year of the study were used for species identification. All 33 RF samples were confirmed to be *C. daubneyi* by PCR and sequencing (Genbank OQ102004–OQ102036). A total of 31 out of 33 samples matched as 99–100% identity to *C. daubneyi* from Ireland (Genbank KP201674), two bovine RF samples had an identity match of 98% to *C. daubneyi*.

3.2.3. Anatomical Changes

During the second year, 28 rumen and 27 duodenal samples were collected and evaluated. The average papillar density was 21.8 papillae/cm$^2$ and 19.9 papillae/cm$^2$ for ovine and bovine samples, respectively. There was no significant difference between species ($p = 0.321; \text{OR} = 6.9; \text{CI} = 0.16–288.2$). Additionally, no significant relationship was found between adult RF presence and papillar density ($p = 0.38; \text{OR} = 7.0; \text{CI} = 0.1–511.1$). The average papillar length was 3.3 mm for ovine samples and 8.6 mm for bovine samples, with a significant difference ($p < 0.0001; \text{OR} = 0.005; \text{CI} = 0.002–0.02)$ between species. However, no significant difference was found between adult RF presence and papillar length when the bovine/ovine difference was taken into account ($p = 0.06; \text{OR} = 0.27; \text{CI} = 0.08–0.98$).

All but one of the twenty-seven duodenal samples had at least one non-specific microscopic change noted (see Appendix A for details of these changes). Immature rumen flukes were not observed in any of the sections examined. There was no significant relationship between any of the seven categories of microscopic duodenal lesions and the presence/absence of adult RF (Table 2).
Table 2. Estimate and $p$-value output from the generalised linear models examining the relationship between adult rumen fluke presence and the histopathological changes of the microscopic duodenal lesions ($n = 27$).

<table>
<thead>
<tr>
<th>Histopathological Changes</th>
<th>Estimate</th>
<th>Standard Error</th>
<th>$p$-Value</th>
<th>Odds Ratio (OR)</th>
<th>95% Confidence Interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inflammation of lamina propria (Lymphocytes and plasma cells)</td>
<td>1.18</td>
<td>1.18</td>
<td>0.32</td>
<td>3.23</td>
<td>0.32–32.48</td>
</tr>
<tr>
<td>Inflammation of lamina propria (Eosinophils)</td>
<td>−1.14</td>
<td>0.91</td>
<td>0.21</td>
<td>0.32</td>
<td>0.05–1.90</td>
</tr>
<tr>
<td>Inflammation of submucosa/Brunner’s glands (Lymphocytes and plasma cells/eosinophils)</td>
<td>−0.12</td>
<td>0.89</td>
<td>0.90</td>
<td>0.89</td>
<td>0.16–5.08</td>
</tr>
<tr>
<td>Inflammation of submucosa/Brunner’s glands (Granuloma)</td>
<td>0.82</td>
<td>1.05</td>
<td>0.43</td>
<td>2.27</td>
<td>0.29–17.58</td>
</tr>
<tr>
<td>Crypt hyperplasia</td>
<td>−2.03</td>
<td>1.32</td>
<td>0.13</td>
<td>0.13</td>
<td>0.01–1.76</td>
</tr>
<tr>
<td>Dilation or hyperplasia of Brunner’s glands</td>
<td>−1.39</td>
<td>1.00</td>
<td>0.17</td>
<td>0.25</td>
<td>0.04–1.77</td>
</tr>
<tr>
<td>Fibrosis submucosa/Brunner’s glands</td>
<td>−0.22</td>
<td>0.94</td>
<td>0.81</td>
<td>0.80</td>
<td>0.13–5.09</td>
</tr>
</tbody>
</table>

4. Discussion

In this study, the trends seen in the presence of RF eggs in routine faecal samples are similar to those seen in other surveys in the UK [1] and Ireland [9,24], where the first significant detection of eggs occurred in 2007–2008, followed by a rapid increase in positive diagnoses. A similar trend was observed in France in the 1990s [25], but it is not known where the original introduction of *C. daubneyi* occurred, nor how it spread so rapidly in Europe. The original isolation and identification of *C. daubneyi* was in the Kenyan highlands [26] and it seems likely that spread to other countries was caused by movement of livestock; however, long distance movement of snails can also occur [27]. This study also suggests *C. daubneyi* is the predominant, if not the only, species of RF present in cattle and sheep within Central and Southwest Scotland.

The number of PMEs carried out over the two-year period was relatively small and therefore interpretation of the results should be guarded. The observed percentage of positive RF cases in bovine animals (43.3%) is somewhat higher than previous studies on finished cattle examined at abattoirs (25% in England and Wales [18] and 29% in Scotland [1]). This could be explained by the different source of samples in these studies. Abattoir samples are collected from a healthy, mainly young population of animals, while cases submitted to the University of Glasgow for PME are primarily animals with undiagnosed or chronic diseases. In contrast, the percentage of sheep that had fluke in the rumen was relatively low (13.5%) compared with other recent surveys in Ireland and Wales [5,9,11], though the values were higher than a study conducted in the Netherlands [28]. Whether these are true differences or the consequence of our small sample size cannot be determined from these data.

Over the 10 years of the FECs study, co-infections with RF and liver fluke in sheep and cattle were quite low (2.5%), despite the fact that all infected animals presumably grazed in habitats containing their intermediate host, *G. truncatula*. Co-infection of snails is considered a rare occurrence (<1%), both experimentally and naturally [2,29]; this points to competition of RF and liver fluke within the intermediate hosts. It has also been speculated that duodenal pathology resulting from immature RF could impede the migration of juvenile liver fluke through the intestinal wall into the abdominal cavity en route to the liver [27]. A potential confounder of these observations is that animals sampled or submitted may have been previously treated with flukicides (other than oxyclozanide) and their liver fluke burdens reduced or eliminated.

Adult RF found at PME were observed at highest densities within the atrium ruminis, surrounding the reticular groove and within the reticulum and, in heavy burdens, flukes were also often observed within the ruminal contents. These observations are in line with current literature citing the location of infection [1,30]. Parasite burdens were predomi-
nantly below 100 adult RF, which agrees with findings from a previous abattoir study in the UK, where most parasitized cattle were found to have ≤100 adult RF at necropsy [18], lower than those reported in Spain [31], Belgium [8] and the Netherlands [28]. No gross pathology was observed in the rumen and reticulum of parasitized animals, nor was there any significant association between the density or length of papillae and RF infection status. Previous studies have found shortening, ulceration and necrosis of parasitized papillae [15] and papillar atrophy [15,32], which could suggest either a different response between healthy cattle and this study cohort, or might reflect a different spectrum of host responses to adult rumen fluke, ranging from apparently healthy tissue through to local inflammatory changes around the attachment sites.

No juvenile RF were observed in the duodenum of any parasitized animals and there was no association between duodenal histopathology and the presence of adult RF. This suggests that previous colonisation of the duodenum by juvenile C. daubneyi did not have a long-term impact on duodenal morphology or function. Collectively, these data indicate that infections with adult rumen fluke are not consistently associated with pathological changes in the rumen or duodenum and therefore lend support to the observed relative lack of pathogenicity of adult paramphistomosis in Europe [33,34].

5. Conclusions

This study has demonstrated that, based on coprological methods, the appearance and increasing prevalence of rumen fluke since the late 2000s, which has been observed in other parts of the UK and in Ireland, has also occurred within Central and Southwest Scotland and, furthermore, that the species responsible is predominantly, if not exclusively, C. daubneyi. The frequency of rumen fluke at post-mortem within the source population (diseased animals) was different compared with those reported elsewhere in abattoir surveys and epidemiological studies. Co-infection with liver fluke in sheep and cattle appeared to be uncommon and this may have implications for the epidemiology of these trematodes in regions where they share the same molluscan intermediate hosts, G. truncatula. No evidence could be found for current or historic pathological changes as a direct result of rumen fluke in the rumen or duodenum, supporting evidence for a lack of severe pathological impact in the final host of moderate infections with adult C. daubneyi.

Author Contributions: Conceptualization, V.B., E.G. and A.F.; methodology, E.G., G.R., G.M., P.S., K.W. and C.M., software, E.G.; formal analysis, E.G.; writing—original draft preparation, V.B.; writing—review and editing, V.B., E.G., G.M., P.S., K.W., C.M. and A.F.; visualization, E.G.; supervision, V.B.; project administration, V.B. All authors have read and agreed to the published version of the manuscript.

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Institutional Review Board Statement: The study was conducted in accordance with the University of Glasgow Ethics Committee regulations (Ref 49a/18).

Data Availability Statement: Data are available upon request to the corresponding author.

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Conflicts of Interest: The authors declare no conflict of interest.
Appendix A

Table A1. Classification of microscopic duodenal lesions.

<table>
<thead>
<tr>
<th>Microscopic Feature</th>
<th>Description</th>
<th>Features Assessed</th>
<th>Classification Used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lymphocytes and plasma cells</td>
<td>Inflammation of the lamina propria</td>
<td>The number of lymphocytes and plasma cells between crypts</td>
<td>1 (0–5 cells between crypts) 2 (5–10 cells between crypts) 3 (10–20 cells between crypts) 4 (&gt;20 cells between crypts)</td>
</tr>
<tr>
<td>Eosinophils</td>
<td>Inflammation of lamina propria</td>
<td>The number of eosinophils in the lamina propria per high-power field (hpf)</td>
<td>1 (&lt;5 eosinophils per hpf) 2 (&gt;5 eosinophils per hpf)</td>
</tr>
<tr>
<td>Lymphocytes and plasmacells/eosinophils</td>
<td>Inflammation of the submucosa/Brunner’s glands</td>
<td>The presence of lymphocytes, plasma cells and/or eosinophils within the submucosa or between Brunner’s glands</td>
<td>Present (clusters of lymphocytes, plasma cells and/or eosinophils) Absent (Rare scattered leucocytes, or complete absence of leucocytes)</td>
</tr>
<tr>
<td>Granuloma</td>
<td>Submucosal granulomas</td>
<td>Presence of granulomas within the submucosa</td>
<td>Present Absent</td>
</tr>
<tr>
<td>Crypt hyperplasia</td>
<td>Hyperplasia of the crypt epithelium</td>
<td>The length and arrangement of crypts, level of epithelial proliferation and the number of mitoses</td>
<td>Present Absent</td>
</tr>
<tr>
<td>Dilation or hyperplasia of Brunner’s glands</td>
<td>Dilation or hyperplasia of Brunner’s glands</td>
<td>Gland size and width, and gland epithelial proliferation</td>
<td>Present Absent</td>
</tr>
<tr>
<td>Fibrosis</td>
<td>Fibrosis of the submucosa/between Brunner’s glands</td>
<td>The presence of greater than two fibrocyte layers separating Brunner’s glands within lobules, or nodules of collagen and fibrocytes surrounding Brunner’s glands.</td>
<td>Present Absent</td>
</tr>
</tbody>
</table>

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