Collagen type III alpha I is a gastro-oesophageal reflux disease susceptibility gene and a male risk factor for hiatus hernia

B Åslng,1 J Jirholt,1 P Hammond,2 M Knutsson,1 A Walentinsson,1 G Davidson,2 L Agreus,3 A Lehmann,1 M Lagerström-Fermer1

ABSTRACT

Background and objectives: Gastro-oesophageal reflux disease (GORD) is a common gastrointestinal disorder with a genetic component. Our aim was to identify genetic factors associated with GORD.

Results: A region on chromosome 2, containing collagen type III alpha 1 (COL3A1), was identified (LOD = 3.3) in families with dominant transmission of GORD, stratified for hiatus hernia (HH). COL3A1 showed significant association with GORD in an independent paediatric trio cohort (p corr = 0.003). The association was male specific (p corr = 0.018). The COL3A1 association was replicated in an independent adult case-control cohort (p corr = 0.022). Moreover, male specific association to HH (p corr = 0.019) was found for a SNP not associated to GORD. Collagen type III protein was more abundant in oesophageal biopsies from male patients with GORD (p = 0.03).

Conclusion: COL3A1 is a disease-associated gene in both paediatric and adult GORD. Furthermore, we show that COL3A1 is genetically associated with HH in adult males. The GORD- and HH-associated alleles are different, indicating two separate mechanisms leading to disease. Our data provides new insight into GORD aetiology, identifying a connective tissue component and indicating a tissue remodelling mechanism in GORD. Our results implicate gender differences in the genetic risk for both GORD and HH.

Gastro-oesophageal reflux disease (GORD) is characterised by a retrograde movement of stomach contents into the oesophagus, leading to symptoms such as heartburn and regurgitation.1 In severe cases, the disease causes tissue erosion and inflammation in the oesophageal mucosal lining. GORD is increasingly common in the Western world, with a prevalence of 25–40% in population-based studies.2 3 Patients suffering from GORD have a severely impaired quality of life,4 and the cost to the society is substantial.5

The establishment of diagnostic criteria for GORD is made complicated by the fact that patients with GORD represent a heterogeneous patient group. Apart from heartburn and acid regurgitation, diverse additional symptoms, including extra-oesophageal manifestations, have been recognised as important disease components.6 Today, diagnosis is generally based on symptomatic presentation complemented by endoscopic and pH probe findings together with data from validated multidimensional questionnaires.6 Less than half of the patients suffer from erosive reflux disease, characterised by mucosal damage in the oesophagus. However, the majority of patients have non-erosive reflux disease, and experience typical GORD symptoms without visible oesophageal injury.7

Epidemiological studies have pinpointed a number of lifestyle-related factors affecting the disease.2 One such risk factor is hiatus hernia (HH), a condition characterised by a protrusion of the upper part of the stomach into the thorax through a tear or weakness in the diaphragm.9 10 HH leads to reflux episodes through an attenuation of the pressure barrier, constituted by the lower oesophageal sphincter in conjunction with the diaphragm.10 Interestingly, there are data indicating a genetic contribution to the development of HH.11

The age of onset of GORD is variable and many individuals develop the disease during childhood. GORD is the most common oesophageal disorder of children, affecting about 11% of all infants during their first year of life.12 It has been suggested that adult GORD may sometimes originate in childhood.13–16 The disease aetiology is further complicated by a substantial genetic contribution as shown by: familial clustering,17 autosomal dominant familial transmission of disease,18 19 as well as twin studies.20–22 Hu and colleagues addressed this and identified a linked region on chromosome 13q14 in families with severe paediatric GORD.18 Orenstein and colleagues, however, failed to replicate this linkage finding in a different GORD family material.19 Although this region is relatively well defined, subsequent work has, so far, not led to the identification of a disease susceptibility gene on chromosome 13q14.23

The aim of this study was to identify genes associated with GORD and to investigate if these genes are shared between paediatric and adult forms of the disease. To address this, four separate patient cohorts were examined in a step-wise manner. First, genome-wide linkage analysis was carried out in families displaying an apparently dominant inheritance of the disease. Next, gene association analyses were performed in a paediatric trio cohort, followed by replication of results in an adult case-control cohort. Lastly, protein levels...
The family patient collection includes an additional 99 individuals with unknown gastro-oesophageal reflux disease (GORD) status. Affected children of the trios are listed as cases and their parents as controls. In the extended Kalixanda cohort, 98 individuals were diagnosed with both GORD and hiatus hernia (HH). The remaining 131 HH individuals were added from the original Kalixanda cohort, producing a separate HH case control material, consisting of 229 cases and the 485 controls.

METHODS

Patient collections

Informed consent was obtained before enrolment. All data and DNA/tissue samples were coded. Ethical approval was obtained for all patient collections.

Families

Enrolment of patients was done at the Gastroenterology Unit at Women’s and Children’s Hospital (WCH, Adelaide, Australia), from 2001 to 2005 by identifying children (age up to 18) diagnosed with GORD having at least one of the following criteria within the last 5 years: an abnormal finding from endoscopic examination or 24 h oesophageal pH test or having been subject to fundoplication. Proband were identified by examination of patient records at WCH and by contacting the local GORD support association. Parents of the proband were contacted and the family was enrolled if a positive family history of GORD could be shown. Disease status was assessed via physician diagnosis often with results of previously performed investigations.24–26 Patients with a medical condition known to predispose to GORD were excluded. HH was identified through previously performed endoscopic examination and/or radiological contrast studies.

Trios

The trio cohort consists of paediatric patients diagnosed with GORD between 3 months and 17 years of age without conditions predisposing to GORD, and their parents. The diagnosis of GORD relied on the following: a paediatric gastroenterologist’s evaluation of symptoms to be consistent with GORD, in addition to either endoscopic, histological or pH probe determined acid exposure consistent with GORD and/or a definite and significant improvement in symptoms of GORD shortly after commencing anti-reflux treatment. Previous anti-reflux surgery was also considered as evidence of GORD. Patients were identified from databases of endoscopy, pH probe and outpatient diagnoses from the Gastroenterology Unit at WCH.

Adult GORD case–control cohort (Extended Kalixanda)

The Kalixanda cohort is a collection of patients from northern Sweden where epidemiological factors related to GORD have been investigated.27–28 We extended this cohort with 100 patients with GORD from local clinics, resulting in a new cohort, referred to as the extended Kalixanda cohort. All individuals went through an endoscopic examination together with a written questionnaire to diagnose GORD and HH.29

Adult GORD case–control cohort (EsoNerd)

EsoNerd is a collection of 80 adult GORD cases (18 males) and eight healthy controls (four males) from western Sweden. Oesophageal biopsies were obtained and used for gene chip expression studies and immunohistochemistry (manuscript in preparation, Pierrou et al).

Healthy volunteers and GORD patients went through pH metric and endoscopic examinations together with an evaluation of symptoms by a gastroenterologist. Symptom-free individuals with negative pH metric and endoscopic examination were chosen as control individuals. Individuals with GORD symptoms with positive acid exposure measurements were defined as GORD patients.

Techniques

DNA extraction

DNA was obtained from blood or buccal swabs using Qiagen Blood DNA extraction kit (Qiagen, Valencia, California, USA) or was whole genome amplified using GenomPhi (Amersham Biosciences, Uppsala, Sweden) according to the manufacturer’s protocol. In cases where the DNA amount was limited, whole genome amplification was performed by Molecular Staging (Denver, Colorado, USA) through their Repli-g service.

Genotyping

Microsatellite genotyping was performed in 419 familial DNA samples using ABI Prism Linkage Mapping set v2.5 HD5 for DNA fragment analysis (Applied Biosystems, Foster City, California, USA). Additional microsatellites were amplified using public primer sequences. The forward primer was ordered 5’–fluorophore labelled (Sigma–Genosys, Cambridge, UK) while the reverse primer was optimised.27 Size fractioning of DNA fragments was done in ABI S700 or 5700 DNA analysers (Applied Biosystems). Alleles were called using the Genotyper v 3.0 software (Applied Biosystems). The mean distance between adjacent markers was 4.32 centimorgan (cM) (SD 2.65); range, 0–14.78. SNP genotyping was performed using the TaqMan assay (Applied Biosystems) and detection in ABI 7900HT (Applied Biosystems) according to the manufacturer’s recommendations, apart from reducing the total reaction volume to 2.5 μl. All data was analysed with Sequence Detection Systems software v2.1 (Applied Biosystems).

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Table 1 The four different GORD patient collections used in this study to identify collagen type III alpha I (COL3A1) as a susceptibility gene for GORD

<table>
<thead>
<tr>
<th>Patient collection</th>
<th>Origin</th>
<th>Usage</th>
<th>Total no of individuals</th>
<th>No of patients with GORD</th>
<th>No of healthy individuals</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Families</td>
<td>Women’s &amp; Children's Hospital Adelaide, Australia</td>
<td>Whole-genome linkage analysis</td>
<td>504</td>
<td>237</td>
<td>168</td>
</tr>
<tr>
<td>2 Trios</td>
<td>Women’s &amp; Children’s Hospital Adelaide, Australia</td>
<td>Genetic association of candidate genes</td>
<td>1092</td>
<td>364</td>
<td>728</td>
</tr>
<tr>
<td>3 Case–control extended Kalixanda</td>
<td>Karolinska Institute, Sweden</td>
<td>Genetic association, replication of findings</td>
<td>741</td>
<td>256</td>
<td>485</td>
</tr>
<tr>
<td>4 Case–control EsoNerd</td>
<td>Sahlgrenska Hospital, Sweden</td>
<td>Immunohistochemistry</td>
<td>38</td>
<td>30</td>
<td>8</td>
</tr>
</tbody>
</table>

Summary

|                  |                  |                  | 2375 | 887 | 1389 |

Patient collections Origin Usage Total no of individuals No of patients with GORD No of healthy individuals

1 Families Women’s & Children’s Hospital Adelaide, Australia Whole-genome linkage analysis 504 237 168
2 Trios Women’s & Children’s Hospital Adelaide, Australia Genetic association of candidate genes 1092 364 728
3 Case–control extended Kalixanda Karolinska Institute, Sweden Genetic association, replication of findings 741 256 485
4 Case–control EsoNerd Sahlgrenska Hospital, Sweden Immunohistochemistry 38 30 8

The four different GORD patient collections used in this study to identify collagen type III alpha I (COL3A1) as a susceptibility gene for GORD.
Sequencing
Sequencing was performed using the Applied Biosystems 3700 or 3730 automated DNA sequencers (Applied Biosystems) according to the manufacturer’s recommendation. Sequencing was performed from both directions and analysis was performed using Sequencher 4.6 (Genecodes, Ann Arbor, Michigan, USA) and compared with the public genomic sequence (ENST00000304636, Ensembl release 49). Sequence differences were manually checked and remaining inconsistencies were resequenced.

Immunohistochemistry
Paraffin-embedded oesophageal tissue sections were treated according to standard protocols for rehydration and antigen retrieval. Immunohistochemistry was performed using the collagen type III specific antibody CL50311AP (Cederlane Laboratories, Burlington, Ontario, Canada) diluted 1:1500. A biotinylated secondary antibody was used (DakoCytomation, Glostrup, Denmark) and the immunoreaction was detected using the chromogen DAB kit: peroxidase/DAB +, Rabbit/Mouse (DAKOytement Code K5001). Digital images from the tissue sections were captured and computerised planimetry was performed using the program Picsara (Euromed Networks, Stockholm, Sweden). The area of interest (total area) was set to the epithelial layer, defined to be between the lumen and the muscularis mucosae. The strong positive immunoreactivation seen in the submucosa of the biopsies was used as an internal positive control for the experiment. The staining in the submucosa was used to confine the positive reaction using the upper and lower threshold values (Picsara). The identified setting was then used in all images.

Statistics
Quality control of familial genotype data
Mendelian inconsistencies were detected using PedCheck (v.1.1). Markers with a large number of errors were either rechecked or discarded. Hardy–Weinberg testing was performed in Pedstats (v.0.6.4) and, in the case of the largest family, SimWalk2 (v.2.91) were used to find genotypes due to unlikely recombinations. Although such genotypes were not automatically discarded, it was ascertained that they did not influence the linkage peaks.

Quality control of trio and extended Kalixanda genotype data
Mendelian inconsistencies in trio genotypes were detected using PedCheck (v.1.1). Hardy–Weinberg testing as well as calculation of descriptive statistics were performed in both Pedstats (v.0.6.4) and Haploview v.3.2 for both the Trios and the extended Kalixanda cohort.

Linkage analysis
Genetic map positions used in the linkage analysis were obtained from the Decode map. Markers that did not have an assigned position were extrapolated into the map based on physical distance to the flanking markers. Family-wise multipoint logarithm of the odds ratio (LOD) score curves were calculated for all autosomal chromosomes and the X chromosome using Genehunter v.2.1 and, in the case of the largest family, SimWalk2 (v.2.91), assuming a dominant mode of inheritance. Single point LOD scores were calculated and compared with the multipoint LOD scores for data consistency. Linkages are reported according to established guidelines. Sensitivity analysis was performed by (1) varying the parameters in the dominant model, (2) assessing an additive mode of inheritance, and (3) performing non parametric analysis. These analyses provided no further information and the results are thus omitted.

Association analysis trios
Single SNPs TDT analysis was performed in Genehunter (v.2.1) and Haploview (v.3.2). TDT analysis based on haplotypes was performed in Genehunter (v.2.1). p values adjusted for multiple
transmit/untransmit-files provided by Genehunter. Written to obtain adjusted p values for haplotypes, using the Haploview (v.3.2) was used, whereas a SAS (v.8.2) macro was committed and untransmitted haplotypes. For single SNPs, et al by Orenstein represented in our material, a finding in line with the results families showed evidence for linkage to chromosome 13 (data region was estimated to be 35 megabase pairs. Chromosome 2q24–q33 with a LOD of 3.3 (fig 1). The size of the families, we were able to identify a common linked region on was absent were selected for further analysis. In this group of that families where HH was present and Barrett’s oesophagus different subsets of the families, the material was stratified so the assumption that different genetic factors are segregating in the EsoNerd study, Pierrou genes within the linkage region (Affymetrix chip U133A & B., Genetic association analysis in the trio cohort RESULTS Linkage analysis The four patient collections used in this study are summarised in table 1. In the initial analysis we investigated the existence of disease-linked chromosomal regions using the entire family collection. We collected 36 families, encompassing a total of 504 individuals. Of these, 237 were affected with GORD and 99 were assigned with unknown disease status. The families were composed of three generations for 22 families, four generations for 12 families and five generations for two families. Based on the assumption that different genetic factors are segregating in different subsets of the families, the material was stratified so that families where HH was present and Barrett’s oesophagus was absent were selected for further analysis. In this group of families, we were able to identify a common linked region on chromosome 2q4–q33 with a LOD of 3.3 (fig 1). The size of the region was estimated to be 35 megabase pairs.

Hu and colleagues previously identified linkage at chromosome 13q14 to GORD in a family material. None of our 36 families showed evidence for linkage to chromosome 13 (data not shown). We conclude that the 13q14 locus is not represented in our material, a finding in line with the results by Orenstein et al. Genetic association analysis in the trio cohort Oesophageal gene expression was used to identify candidate genes within the linkage region (Affymetrix chip U133A & B., EsoNerd study, Pierrou et al, manuscript in preparation). Three genes, CFLAR (CASP8 and FADD-like apoptosis regulator), COL5A1 (collagen type III alpha I) and ALS2CR2 (amyotrophic lateral sclerosis 2 (juvenile) chromosome region, candidate 2, also called STRADB), showed differential expression in oesophageal biopsies from patients with GORD compared with healthy controls. Genotyping of single nucleotide polymorphisms (SNPs) in the three genes in the 364 Trios, followed by genetic association analysis using transmission disequilibrium test (TDT), revealed significant association for SNPs only in COL3A1.

To establish that the association was confined to COL3A1, we analysed the linkage disequilibrium (LD) structure in the region by assaying multiple SNPs in COL3A1 as well as in the flanking genes DIRC1 (disrupted in renal carcinoma 1) and COL5A2 (collagen, type V, alpha 2). The LD block containing the disease-associated SNPs only harbouring COL3A1. In males, we identified an associated haplotype in COL3A1 (p corr of 0.0009; table 2) and an associated SNP (rs6434304) with a p corr (p value adjusted for multiple testing) of 0.02 (table 3). No single markers or haplotypes showed evidence for association in females in the Trio cohort.

Genetic association analysis in the case–control cohort Association analysis was repeated in the adult case–control cohort consisting of 256 GORD cases, 229 hiatus hernia cases, and 485 healthy controls. GORD and HH co-existed in 98 individuals. We used the same set of COL3A1 SNPs as previously, and were able to replicate the GORD association at a p corr = 0.022 (table 5). No male specific association was observed.

The linkage to chromosome 2 originated from families selected for GORD and HH. To investigate if COL3A1 also was associated with HH, we stratified the case–control cohort for this phenotype and repeated the analysis. Significant association was identified in COL3A1. Interestingly, this was also male specific, with the most significant SNP (rs5134646) showing a p corr = 0.019 (table 5). It was not possible to validate our finding in the paediatric trio cohort since HH typically develops after adolescence.

Mutation analysis of COL3A1 Mutation detection was performed by polymerase chain reaction (PCR) amplification and DNA sequencing of all 51 exons of COL3A1 as well as 2 kb upstream of the transcription start site in 48 males and females from the trio cohort. These individuals represented both contributors and non-contributors with respect to the observed association. Intron–exon boundaries were included in the analysis. No amino acid changing mutation segregating with the disease was identified.

Collagen type III protein expression analysis in oesophageal biopsies The abundance of collagen type III was investigated in the epithelial lining of the distal oesophagus. Biopsies from GORD patients and healthy controls were obtained and stained with an antibody directed against human collagen type III. Patients
had a significantly larger surface area staining than controls (p = 0.038 from 30 cases and eight controls. Thus, the data shows that collagen type III is upregulated in the epithelial lining of the oesophagus as a consequence of GORD. After gender stratification, the up regulation remained significant in males only (p = 0.033, 18 cases and four controls).

Discussion

In the present study, based on four independent GORD patient collections (table 1), we show that COL5A1 is associated with GORD and implicated as being a risk factor for hiatus hernia (table 5). We identified linkage to the COL5A1 region in a set of families in which GORD and hiatus hernia is transmitted (fig 1). We show genetic association for COL5A1 to GORD in a paediatric Trio cohort and replicate the genetic association in an adult GORD case–control cohort (tables 2 and 3). We also show genetic association between COL5A1 and hiatus hernia in the same adult case–control cohort (table 3). Our genetic findings from these three independent patient materials are further strengthened by the protein studies from a fourth patient material, where the COL5A1 protein is up regulated in oesophageal tissue biopsies taken from GORD patients in comparison to healthy controls. We show that COL5A1 is associated to both GORD and HH but with different alleles. This becomes even clearer when comparing gender distribution where only males contribute to the HH association (table 3).

The gene COL5A1 encodes type III collagen, a fibrillar monoclonal extra cellular matrix protein that is present in extensible connective tissues such as skin, lung, and the vascular system, frequently together with type I collagen. Collagen type III has an important role in adjusting the strength and flexibility of tissues where it is expressed.43 Furthermore, COL5A1 is modulated in the wound response process. It is acutely upregulated in the early phases of wound healing and maintains high levels of expression for several weeks after injury. As healing progresses, collagen type III is replaced by collagen type I, leading to increased tissue strength.43

A number of COL5A1 mutations cause Ehlers–Danlos syndromes type III (EDS-III) and type IV (EDS-IV), both being autosomal dominant connective tissue disorders. Depending on the individual COL5A1 mutation, the severity of these diseases can vary from mild to life threatening. Symptoms include dermal manifestations, joint hyper-mobility and, in severe cases, spontaneous rupture of the bowel or large arteries.44 COL5A1 mutations similar to those underlying EDS are unlikely to be disease-causative in patients with GORD due to the milder phenotypic expression seen in GORD compared with EDS. One would rather expect mutations resulting in more subtle effects on gene function.

The genetic associations we see for haplotypes using the assayed SNPs (table 2) are much stronger than the genetic association seen for single SNPs (table 3). We conclude that none of the SNPs used in this study are causative but rather in linkage disequilibrium with the different disease alleles. We investigated the presence of causative mutations by sequencing the COL5A1 gene in 48 patients from the trio cohort. No disease-associated mutation, apart from the SNPs assayed, could be identified. Our interpretation is that the disease-causative mutations reside in regulatory sequences.

In our genetic analysis we identify several gender differences in the COL5A1 associations, both for GORD and HH. In the paediatric Trio cohort, only males showed association with GORD. In the adult case–control cohort, COL5A1 was associated with GORD without a gender bias, whereas only males showed association with HH. Finally, elevated collagen type III protein staining of the oesophageal biopsies from GORD patients was only found to be significant in males.

These differences can be attributed to differences between our materials such as: age, population origin, gender composition and disease definition. In fact, COL5A1 itself, is partly regulated by oestrogen.45 This is also shown by the decline of collagen type III expression in post-menopausal women.46 Additionally, a gender difference in collagen deposition during wound healing has been reported in disease models.47 Furthermore, male gender and increasing age has been shown to be a risk factor for elevated wound healing times.48 Hence, the differences in association we observe in our analyses may be a consequence of the biological regulation of collagen type III.

From the data we have generated in the extended Kalixanda cohort it is clear that two different mechanisms are at work in COL5A1. One allele is associated with GORD while another allele confers male susceptibility to HH. The SNP that shows the strongest association with GORD (p_corr = 0.02) shows absolutely no evidence for association with HH (p_corr = 0.94). In this study we have an overlap of 98 patients where both GORD and HH are present. When testing if the male HH-associated SNP is also associated to GORD in males, the

Table 3

<table>
<thead>
<tr>
<th>Number</th>
<th>SNP</th>
<th>Gastro-oesophageal reflux disease</th>
<th>Case-control, males</th>
<th>Hiatus hernia</th>
<th>Case-control, males</th>
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</thead>
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<tr>
<td></td>
<td></td>
<td>p Value</td>
<td>p corr</td>
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</table>

Only the males from the Trio cohort show significant genetic association for SNP number 2 after multiple testing adjustments with a p_corr of 0.018. There is no gender bias in the adult case control cohort. Significant association for GORD in the case–control cohort was identified for SNP number 5 with a p_corr of 0.022. COL5A1 is also associated with hiatus hernia in the adult case–control cohort. The associated SNP, number 8, has a p_corr of 0.019. The identified association is male specific. The patient overlap is 98 individuals, eg, having both GORD and hiatus hernia.
association is further weakened, underlying that there are two separate disease alleles for GORD and HH.

We hypothesise that an altered collagen type III expression in the oesophagus might lead to altered oesophageal tissue strength and flexibility, making the oesophageal lining prone to mucosal damage and wounds. Attenuated COL3A1 function may impair wound healing in the oesophagus. Mutations in COL3A1 have been shown to delay the wound healing response. An altered collagen type III expression could also contribute to an increased risk for developing HH. An altered collagen I/III ratio has been discussed in conjunction with HH. Hiatus hernia might be a risk factor for developing GORD through a mechanism involving increased distensibility of the oesophagogastric junction, resulting in GORD as a secondary event.

Moreover, HH formation may lead to delayed gastric emptying, increasing the clearance time for acid refluxes in the oesophagus. We suggest that the associated alleles of COL3A1 are conferring increased susceptibility to both GORD and HH, respectively.

We welcome a replication of our results from other groups to verify our findings of COL3A1 and, hopefully, identification of the causative mutations. We also raise the question whether there are differences in the wound healing response in controls versus individuals with HH due to an altered collagen I/III ratio.

Acknowledgements: We are grateful to all the patients with GORD and their relatives for participating in this study. We thank FISRA (Friends of Infant Reflex Association) in South Australia for their support. We thank T Storskrubb, P Aro and J Ronkainen for the patient collection and characterisation of the extended Kalanka cohort.

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Competing interests: None.

Ethics approval: Informed consent was obtained before enrolment. All data and DNA/tissue samples were coded. Ethics approval was obtained for all patient cohorts. DNA samples were collected from 312 patients in a random sample of the Swedish adult general population: the Kalixanda study. DNA/tissue samples were coded. Ethics approval was obtained for all patient collections. Extended Kalanka was approved by Umeå University with reference number 321/03, diary nr 03-285. Families and Tors was approved by WER Research Ethics Committee with reference numbers REC1281/11/2004 and REC1340/6/2005. EsoNerd was approved with reference number S 499-3. Decision 2003-11-18 Studiekod: DPM-ESEP-0002.

REFERENCES

ANSWERS

From the questions on page 1047

The air pockets were also traced to the splenic vein (fig 1A, below), inferior mesenteric vein (fig 1B) and perirectal space (fig 1C). Enhancement of the small bowels was intact. Emergency exploratory laparotomy on the same day as the computed tomography (CT) scan was performed revealed turbid ascites but intact peristalsis and vital colour of the small bowels and colon; therefore, bowel resections were not performed. Haemorrhagic and inflammatory change of pancreas was found. Rubber drains were inserted and after normal saline irrigation they drained dark bloody fluid. Subsequent colonoscopy disclosed a short-segment mucosal tag and white-plaque coated and bullae formationed mucosa at rectum and sigmoid colon, compatible with ischaemic colitis. Acute necrotising pancreatitis complicated with ischaemic colitis with aeroportia, pneumatosis coli and renal cortical necrosis was diagnosed. Oral feeding was resumed 24 days after operation and the patient was discharged with normalised amylase and lipase level after a 70 day admission.

Colonic complications were more frequent in severe pancreatitis, occurring in 15%. The principal presentations were necrosis, fistula and stricture. All episodes of colonic necrosis complicated severe acute pancreatitis, were diagnosed operatively, presented at a median of 25 (1–55) days into the episode and were associated with a mortality of 54%. Surgical resection is the mainstay of management of necrosis. In this case, the portal vein air could be traced to the inferior mesenteric vein via the perirectal space and ischaemic colitis of rectosigmoid colon was also confirmed in the subsequent colonoscopy. The enhancement of small bowels and colon was intact in the CT scans and confirmed by exploratory laparotomy.

Renal cortical necrosis is usually extensive and bilateral owing to decreased blood supply within the renal cortex but sparing of medulla, juxtamedullary cortex and a thin rim of subcapsular cortex. The majority of cases are identified following complications of late pregnancy and septic abortion. Rarely, bilateral cortical necrosis complicating acute necrotic pancreatitis is observed.

Acute pancreatitis is a commonly encountered condition in the emergency department but its complications of portal venous air and acute renal cortical necrosis are rare. We encountered this case with two concurrent rare complications. Acute renal cortical necrosis was ignored due to her underlying end-stage renal disease. The portal venous air combined with acute and massive bowel necrosis usually has a dismal result. When we had a careful perusal, the air was from the rectal area via the inferior mesenteric vein into the splenic vein and portal vein. The small and large bowels were still well-enhanced. The surgical intervention could be reserved for any further deterioration in symptoms or signs.

References

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