Eosinophil associated genes in the inflammatory bowel disease 4 region: Correlation to inflammatory bowel disease revealed

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Abstract

AIM: To study the association between inflammatory bowel disease (IBD) and genetic variations in eosinophil protein X (EPX) and eosinophil cationic protein (ECP).

METHODS: DNA was extracted from ethylene diamine tetraacetic acid blood of 587 patients with Crohn's disease (CD), 592 with ulcerative colitis (UC) and 300 healthy subjects. The EPX405 (G > C, rs2013109), ECP434 (G > C, rs2073342) and ECP562 (G > C, rs2233860) gene polymorphisms were analysed, by the 5'-nuclease allelic discrimination assay. For determination of intracellular content of EPX and ECP in granulocytes, 39 blood samples was collected and extracted with a buffer containing cetyltrimethylammonium bromide. The intracellular content of EPX was analysed using an enzyme-linked immunosorbent assay. The intracellular content of ECP was analysed with the UniCAP® system as described by the manufacturer. Statistical tests for calculations of results were $\chi^2$ test, Fisher's exact test, ANOVA, Student-Newman-Keuls test, and Kaplan-Meier survival curve with Log-rank test for trend, the probability values of $P < 0.05$ were considered statistically significant.

RESULTS: The genotype frequency for males with UC and with an age of disease onset of $\geq 45$ years ($n = 57$) was for ECP434 and ECP562, GG = 37%, GC = 60%, CC = 4% and GG = 51%, GC = 49%, CC = 0% respectively. This was significantly different from the healthy subject's genotype frequencies of ECP434 (GG = 57%, GC = 39%, CC = 5%; $P = 0.010$) and ECP562 (GG = 68%, GC = 29%, CC = 3%; $P = 0.009$). The genotype frequencies for females, with an age of disease onset of $\geq 45$ years with CD ($n = 62$), was for the ECP434 and ECP562 genotypes GG = 37%, GC =
52%, CC = 11% and GG = 48%, GC = 47% and CC = 5% respectively. This was also statistically different from healthy controls for both ECP434 (P = 0.010) and ECP562 (P = 0.013). The intracellular protein concentration of EPX and ECP was calculated in μg/10⁶ eosinophils and then correlated to the EPX 405 genotypes. The protein content of EPX was highest in the patients with the CC genotype of EPX405 (GG = 4.65, GC = 5.93, and CC = 6.57) and for ECP in the patients with the GG genotype of EPX405 (GG = 2.70, GC = 2.47 and CC = 1.90). ANOVA test demonstrated a difference in intracellular protein content for EPX (P = 0.009) and ECP (P = 0.022). The age of disease onset was linked to haplotypes of the EPX405, ECP434 and ECP562 genotypes. Kaplan Maier curve showed a difference between haplotype distributions for the females with CD (P = 0.003). The highest age of disease onset was seen in females with the EPX405CC, ECP434CC, ECP562CC haplotype (34 years) and the lowest in females with the EPX405GC, ECP434GC, ECP562GG haplotype (21 years). For males with UC there was also a difference between the highest and lowest age of the disease onset (EPX405CC, ECP434CC, ECP562CC, mean 24 years vs EPX405GC, ECP434GC, ECP562GG, mean 34 years, P = 0.0009). The relative risk for UC patients with ECP434 or ECP562-GC/CC genotypes to develop dysplasia/cancer was 2.5 (95%CI: 1.2-5.4, P = 0.01) and 2.5 (95%CI: 1.1-5.4, P = 0.02) respectively, compared to patients carrying the GG-genotypes.

CONCLUSION: Polymorphisms of EPX and ECP are associated to IBD in an age and gender dependent manner, suggesting an essential role of eosinophils in the pathophysiology of IBD.

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Key words: Eosinophil derived neurotoxin; RNase 2; RNase 3; Single nucleotide polymorphism; Inflammation bowel disease; Crohn’s disease; Ulcerative colitis

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INTRODUCTION

The concept of inflammatory bowel disease (IBD) comprises the disorders Crohn’s disease (CD) and ulcerative colitis (UC). The aetiology of the diseases is unknown, but IBD is currently believed to develop in individuals with a hereditary predisposition. This is related to the reaction of the immune system to the bacterial challenge of the gut and/or defects in the gut wall, in addition to external environmental factors. IBD usually has an intermittent course of aggravations with common symptoms such as frequent diarrhoea, blood in stool, abdominal pain, and weight loss. The incidence of IBD has increased in the Western world in recent years, and new cases occur primarily in the group between 20 and 40 years of age[1,2]. CD seems to affect women more than men, while UC appears to be more prevalent among men, but some studies showed no gender differences[3-5]. A bimodal age distribution of disease onset has been reported and debated in population studies of CD and UC[6,8]. Patients with CD and UC have an increased risk of developing colorectal cancer. This risk is more pronounced in patients with UC with an overall 2.7-fold increased risk as compared with the general population[9].

Under normal conditions, the intestinal mucosa is in a state of “controlled inflammation”. Different subsets of T-cells are present in the healthy intestine, as are eosinophils and antigen-presenting cells with a predominance of anti-inflammatory and regulatory cytokine responses that keep the mucosal homeostasis intact[10]. However, in IBD the balance between mucosal responsiveness and tolerance towards antigens is disturbed, resulting in an exaggerated immune response to the commensal flora. It is well established that neutrophil granulocytes accumulate and infiltrate in the local processes in IBD. In recent years attention has been paid to the involvement of eosinophil granulocytes, since increased numbers of these cells are found in the intestinal mucosa of IBD[11,12]. Studies now emphasise a key role of the eosinophil granulocyte in the pathophysiology of IBD[13,14].

The eosinophil participate in a number of biological processes such as wound healing, defence against parasites and allergic inflammation[15]. The eosinophil granulocyte are characterised by specific cytoplasmic granules containing four major proteins; eosinophil cationic protein (ECP)[16], eosinophil peroxidase (EPO)[17], eosinophil protein X (EPX)[18], and major basic protein (MBP)[19]. Increased intraluminal release of EPX, ECP and EPO has been shown in patients with UC[19]. The cytotoxic activities of ECP and EPX as well as their potent RNase activities have been well documented[20] and are generally conceived as part of the host defence against viruses, parasites and helminths[21,22]. Furthermore both EPX and ECP have been associated with different types of cancer[23,24]. The presence of eosinophils in the intestinal mucosa of healthy subjects may reflect these host defence activities of the cell. In addition EPX has been identified as a member of the Alarmin family with the capacity to affect various aspects of the dendritic cells[25] and in aged asthma patients, relative to young, the degranulation of EPX was decreased[26]. Larger numbers of activated eosinophils, identified as an increased expression of the cell surface receptor CD44, have been reported both in CD and UC, but even more
pronounced in quiescent than in active UC. Findings that might suggest a role of eosinophils in tissue remodelling and repair in IBD[26,29].

By linkage analysis and genome-wide association studies many genetic variations, linked to IBD, have been found[30]. The nucleotide sequences of EPX and ECP have a homology of 90%, and both genes have been located to chromosome 14q11.2[31,32] within the IBD4 locus, located at the 14q11-12 region[31]. Several single nucleotide polymorphisms (SNPs) were reported in the genes for EPX and ECP[33]. An intron SNP in the EPX gene, EPX405 G > C (rs2013109) was together with a SNP from the 3’ untranslated region in the ECP gene, ECP562 G > C (rs2233860)[33], shown to be closely linked to the eosinophil content of EPX and ECP (Jönsson et al[34], to be published). The C-allele of the nonsynonymous missense ECP434 G > C (rs2073342) polymorphism, gives rise to an arginine to threonine shift at amino acid position 97 in the mature protein[35,36]. As shown with recombinant ECP proteins this amino acid shift resulted in an alteration of the protein and its cytotoxic activity[37], but with no effect on its RNase activity[38]. The loss of cytotoxicity of ECP containing threonine at position 97 was confirmed using purified native proteins from genotyped blood donors[39]. The ECP434(G > C) polymorphism was shown to be associated with the expression of allergic symptoms[40], and with disease severity in Hodgkin lymphoma[41] in population based studies as well as to the prevalence and severity of Schistosoma mansoni infection in a Ugandan population[42].

The aim of this study was to investigate the impact of SNPs in the EPX and ECP genes in a cohort of patients with IBD. The hypothesis was that alterations in the cytotoxic activities of ECP and/or the altered expression of EPX might affect the propensity to acquire IBD and also might affect different features of IBD. The influence of polymorphisms in the EPX and ECP genes was first analysed in a pilot cohort of patients with IBD from Uppsala University Hospital and then in a second step in a bigger Swedish cohort. The possible altered expression of EPX was in addition analysed by measuring the intracellular eosinophil content of EPX in a subset of the patients.

MATERIALS AND METHODS

Study population

The patient group consisted of 1179 individuals recruited from Uppsala University Hospital, Sweden (95 CD and 100 UC), and an established Swedish cohort recruited at Karolinska University Hospital and Orebro University Hospital (492 CD and 492 UC). Diagnoses of CD and UC was based on clinical, histological and endoscopic findings, according to standardized criteria[43]. Bimodal patterns as to age of disease diagnosis were observed for both CD and UC irrespective of gender (Figure 1). The major peak of age at diagnosis of both diseases was seen at 20-25 years, with a second peak around 40-50 years.

A group of 300 healthy Swedish blood donors from the Uppsala blood bank served as reference population.

The study was approved by the local Ethics Committees of the Medical Faculties, of Uppsala University and Karolinska University Hospital, Stockholm, and all patients gave their informed consent to participate in the study (Table 1).

DNA extraction

Genomic DNA from all IBD patients and the reference group were extracted from whole ethylene diamine tetraacetic acid (EDTA) blood, by the use of the QIAamp DNA Blood Mini Kit from QIAGEN (QIAGEN Inc, Valencia, CA, United States), according to the manufacturer’s description.

Genotyping

All IBD samples and controls were genotyped for the EPX405 (G > C), ECP434 (G > C) and the ECP562 (G > C) polymorphisms by use of the 5’-nucleotide allelic discrimination assay. The assay has previously been described in detail[44]. Oligonucleotide primers were designed by the use of Primer Express® (Applied Biosystems, Foster City, CA, United States) based on the se-
EDTA blood was collected from 39 patient samples with intracellular content of EPX and ECP in granulocytes, EPX and ECP in eosinophils. For the determination of an interest in measuring the intracellular content of EPX content between CD and UC patients, which lead analysed in the Uppsala cohort. There was a difference in the EPX and ECP polymorphisms were first only anal-

Blood eosinophil count (B-Eos) (reference interval 0.0 × 10⁹/L-0.5 × 10⁹/L) and total white blood cell count (reference interval 3.5 × 10⁹/L-9.0 × 10⁹/L) were determined by an automated haematology analyser (Celldyl Sapphire, Abbott Laboratories, CA, United States) at the Department of Clinical Chemistry, Uppsala University Hospital, Uppsala, Sweden.

Whole blood extraction The EPX and ECP polymorphisms were first only analysed in the Uppsala cohort. There was a difference in the EPX content between CD and UC patients, which lead to an interest in measuring the intracellular content of EPX and ECP in eosinophils. For the determination of intracellular content of EPX and ECP in granulocytes, EDTA blood was collected from 39 patient samples with

| Table 1 | Clinical characteristic of study populations |
|----------------|---------------------------------|
|          | Crohn’s disease (n = 587) | Ulcerative colitis (n = 592) | Reference group (n = 300) |
| Female  | 282 | 269 | 132 |
| Male    | 305 | 323 | 168 |
| Median age¹ | 48 (18-95) | 48 (14-102) | 44 (19-73) |
| Age at disease diagnosis² | 28 (6-93) | 28 (4-78) | |

¹Range at sampling; ²Median (range).

| Table 2 | Primers and probes |
|----------------|-------------------|
| EPX405 (G > C): | |
| Forward primer | 5’ AAG AGA GCT GAC GTT AGT GCT TAG G 3’ |
| Reverse primer | 5’ GGT CTT GGT TAT GCA ACA CAC TGC ATG 3’ |
| Probe 1 | 5’ VIC-AGC TTT CAC ACT TT 3’ |
| Probe 2 | 5’ 6-FAM-CCT TCC ACA ACG TTG 3’ |

ECP434 (G > C): |
| Forward primer | 5’ GAG TAG ATT CCG GCT GCC TTT ACT 3’ |
| Reverse primer | 5’ CCT CTA TCC CAG GTG GGA AG 3’ |
| Probe 1 | 5’ VIC-AAA CTG CAG GTA TGA ACA 3’ |
| Probe 2 | 5’ 6-FAM-CCT TCC ACG GTC TGC AG 3’ |

ECP562 (G > C): |
| Forward primer | 5’ GCT TCC AGT TCA CCT GGA TAC C 3’ |
| Reverse primer | 5’ GGT ATG GAG ACT GAT GAG GAC AGT 3’ |
| Probe 1 | 5’ VIC-TCA GCA TCT CTC ATC 3’ |
| Probe 2 | 5’ 6-FAM-CCT TCCA CTC CTC ATC 3’ |

EPX: Eosinophil protein X; ECP: Eosinophil cationic protein.

sequence of the EPX (X16546) and ECP (X16545) genes available in GenBank. Sequences for primers and probes are displayed in Table 2.

Polymerase chain reaction (PCR) cycling was carried out in an ABI PRISM 7000 (Applied Biosystems) with the recommended thermal profile of: 50 °C for 2 min, 95 °C for 10 min followed by a total of 40 cycles of 95 °C for 15 s and 60 °C for 1 min. After the PCR cycling the genotypes were determined according to the application Allelic Discrimination of the ABI Prism 7000 SDS software (Applied Biosystems).

Blood cell count
Blood eosinophil count (B-Eos) (reference interval 0.0 × 10⁹/L-0.5 × 10⁹/L) and total white blood cell count (reference interval 3.5 × 10⁹/L-9.0 × 10⁹/L) were determined by an automated haematology analyser (Celldyl Sapphire, Abbott Laboratories, CA, United States) at the Department of Clinical Chemistry, Uppsala University Hospital, Uppsala, Sweden.

Whole blood extraction
The EPX and ECP polymorphisms were first only analysed in the Uppsala cohort. There was a difference in the EPX content between CD and UC patients, which lead to an interest in measuring the intracellular content of EPX and ECP in eosinophils. For the determination of intracellular content of EPX and ECP in granulocytes, EDTA blood was collected from 39 patient samples with

CD. The subjects were selected based on their EPX405 (G > C) genotypes (GG = 15, GC = 16, and CC = 8). The method for whole blood extraction using cetyltrimethylammonium bromide has previously been described in detail (Jönsson et al[35], to be published). The supernatants were stored at -20 °C until assayed for EPX and ECP concentrations.

Protein assays
EPX whole cell extract was analysed using an enzyme-linked immunosorbent assay (Diagnostics Development, Uppsala, Sweden), according to the recommendations of the manufacturer with minor modifications. The concentration of ECP was analysed with the UniCAP® system, (Phadia AB, Uppsala, Sweden) as described by the manufacturer.

Statistical analysis
All the statistical analyses were performed with the Statistica 9.0 software for Windows (StatSoft, Tulsa, United States), and MedCalc software (V.10, Mariakerke, Belgium). The statistical tests in use for the calculations of results were χ² and Fisher’s exact test, ANOVA, Student-Newman-Keuls test, and Kaplan-Meier survival curve with the Log-rank test for trend. P < 0.05 were considered statistically significant.

RESULTS
Genotype distributions in healthy population vs patients with IBD
The distribution of the EPX405, ECP434 and ECP562 genotypes was investigated in the population of 1179

Table 3  Primers and probes

| Genotype and allelic frequency distribution among 300 healthy individuals and the total cohorts of Crohn’s disease and ulcerative colitis |
|----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
|                | GG-genotype (%) | GC-genotype (%) | CC-genotype (%) | G-allele (%)    | C-allele (%)    |
| Healthy        | 68              | 29              | 3               | 83              | 17              |
| ECP434         | 57              | 38              | 5               | 76              | 24              |
| EPX405 M¹      | 64              | 32              | 4               | 80              | 20              |
| EPX405 F¹      | 53              | 38              | 9               | 72              | 28              |
| CD             | 64              | 33              | 4               | 80              | 20              |
| ECP434         | 53              | 40              | 7               | 73              | 27              |
| EPX405 M       | 55              | 38              | 6               | 74              | 26              |
| EPX405 F       | 58              | 34              | 8               | 75              | 25              |
| UC             | 64              | 33              | 4               | 80              | 20              |
| ECP434         | 55              | 38              | 7               | 74              | 26              |
| EPX405 M       | 62              | 31              | 7               | 78              | 22              |
| EPX405 F       | 57              | 38              | 5               | 76              | 24              |

For ECP434 and ECP562 there was no difference between males and females. In EPX405 there was a difference between the sexes according their genotype (P = 0.049) and allelic distribution (P = 0.021). UC: Ulcerative colitis; CD: Crohn’s disease; EPX: Eosinophil protein X; ECP: Eosinophil cationic protein; F: Female; M: Male.

For EPX405 there was a difference between the sexes according their genotype (P = 0.049) and allelic distribution (P = 0.021). UC: Ulcerative colitis; CD: Crohn’s disease; EPX: Eosinophil protein X; ECP: Eosinophil cationic protein; F: Female; M: Male.
subjects with CD ($n = 587$) or UC ($n = 592$) (Table 1).

For comparison the distributions of the same genotypes were investigated in a group of apparently healthy blood donors ($n = 300$). In the healthy reference population males had a higher prevalence of the EPX405 GG-genotype ($P = 0.049$) and the G-allele ($P = 0.021$) compared to females (Table 3). All continuing data has been calculated gender stratified for the EPX405 genotype. No difference was seen between genders for the ECP434 and ECP 562 genotype in the control cohort (Table 3). No genotype differences were seen between younger ($n = 150$; median age: 31 years; range: 19-43 years) and older ($n = 150$; median age: 54 years; range: 44-73 years) healthy references. At first only the Uppsala cohort with CD ($n = 95$) and UC ($n = 100$) was examined. Here there was a difference between patients with CD and UC ($P = 0.048$) with more EPX405 GC and CC in CD. These results could not be verified in the whole cohort where no general differences were found between the healthy and the patient cohorts, or between the CD and the UC cohorts, for any of the genotypes. When dividing the patient cohorts by their median (28 years) age at disease diagnosis (ADD) and comparing them, no difference was seen in any of the genotypes for younger vs older patients with CD (median ADD: 20 years; range: 6-28 years vs median ADD: 40 years; range: 28-93 years) or UC (median ADD: 20 years; range: 4-28 years vs median ADD: 40 years; range: 28-78 years). In the total cohort, no difference was seen in the genotype frequencies related to gender in patients with CD or UC. When the patient populations were separated by age at disease diagnosis, according to the Montreal classification$^{46}$ (< 16 years, 17-40 years and > 40 years) and gender, a pattern with significant differences was observed. Since the incidence curve (Figure 1) of women with CD indicated a second peak of disease onset around 45 years the age range > 45 years was also included in the study. Thus, a lower prevalence of the ECP434 and ECP562-GG genotypes and higher prevalence of the ECP434 and ECP562-GC genotypes were found in female patients with CD, compared to the healthy reference population (Table 4). In males similar differences in the prevalence of the ECP434 and ECP562 genotypes were observed, but only in patients with UC (Table 4).

For both genders the genotype differences were observed in the older cohort of the population with an age at diagnosis above 40 years (Table 4). In the cohort of males with UC a higher prevalence of the EPX405 GC-genotype, compared to healthy controls, was observed in patients with an ADD of > 40 years ($P = 0.048$).

**Genotype distribution within different cohorts**

Among females with CD and ADD of > 45 years, the GC-genotype of the ECP434 ($P = 0.020$) and ECP562 ($P = 0.014$) gene polymorphisms were more common than in females with ADD < 45 years. In females with UC, no such differences were observed. In males with UC there was a difference between those with an ADD over and under 40 and 45 years, concerning all three genotypes (40 years; EPX405 $P = 0.003$, ECP434 $P = 0.003$ and ECP562 $P = 0.004$). The differences were, in all genotypes, due to a higher amount of the GC-genotypes among males > 40 years and > 45 years.

In Figure 2, a comparison of the ECP434 genotype distributions between patients with CD and UC and an age of disease diagnosis of > 45 years is shown. In the comparison the population is separated by gender. Among female patients with CD there was a significantly lower prevalence of the ECP434GG genotype and a prevalence of the heterozygote genotype almost twice as high as in those women having UC ($P = 0.010$). This was contrasted by the findings in males, since those patients having CD had a predominance of the ECP434 GG-genotype, as compared to patients with UC, who had a predominance of the GC-genotype ($P = 0.004$). Highly significant differences were found in ECP434 genotype distributions between females and males (> 45 years) with CD ($P = 0.001$) and with UC ($P = 0.001$). Similar patterns were found for the ECP562 genotypes, but not for the EPX405 genotypes (not shown).

**Relationships of age at disease diagnosis to genotypes and haplotypes**

The relationships of the EPX405, ECP435 and ECP562 genotypes and haplotypes (Table 5) to ADD (females with CD and males with UC) are illustrated in Figure 3. The haplotypes analysed were chosen to include the wild type (WT) (haplotype 1; GG in all three genotypes) and

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**Table 4 Gender specific difference**

| Genotype distribution | Allelic frequency | Genotype distribution | Allelic frequency |
|-----------------------|-------------------|-----------------------|-------------------|
| EPX                  | ECP               | ECP                  | EPX               | ECP               | ECP                   |
| CD/UC ($n$)           |                   |                      | CD/UC ($n$)       |                   |                       |
| Female                |                   |                      | Female            |                   |                       |
| $> 40$                | 80/71             | NS                   | $> 40$            | 80/71             | NS                    |
| $> 45$                | 42/62             | NS                   | $> 45$            | 42/62             | NS                    |
| Male                  |                   |                      | Male              |                   |                       |
| $> 40$                | 77/85             | NS                   | $> 40$            | 77/85             | NS                    |
| $> 45$                | 57/52             | NS                   | $> 45$            | 57/52             | NS                    |

Difference ($\chi^2$, $P$-values displayed) between the genotype distributions and allelic frequencies, related to age at diagnosis, in individuals with CD or UC compared with the healthy population. UC: Ulcerative colitis; CD: Crohn’s disease; NS: Not significant; EPX: Eosinophil protein X; ECP: Eosinophil cationic protein.
In females, age at disease diagnosis (ADD) > 45 years, a significant difference between Crohn’s disease (CD) and ulcerative colitis (UC) concerning ECP434 (P = 0.008) (Figure 3B). In the study UC cohort there were several patients (n = 27; females: 8; males: 19) that through continuous endoscopic examination were shown to have developed dysplasia and/or cancer (D/C) (median age at cancer diagnosis: 49 years; range: 27-67 years). The D/C cohort was compared to all UC patients, females and males who had not developed D/C, since no difference was shown between older and younger UC patients in the mixed cohort. The distributions of the ECP434 and ECP562 alleles in patients with UC, with and without findings of D/C of the intestine are shown in Figure 5. The C-allele of both genotypes was more prevalent among patients with D/C (ECP434 P = 0.034 and ECP562 P = 0.019). The D/C patients also differed from the healthy controls (ECP434 P = 0.027 and ECP562 P = 0.007) (Figure 5). The relative risk (RR), for UC patients with the ECP434 GC or CC genotypes, to develop D/C was 2.5 (95%CI, 1.2-5.4) compared to other patients with UC. In patients with the ECP562 GC or CC genotypes the RR was 2.5 (95%CI, 1.2-5.8) as compared to UC patients. Considering the whole cohort, no associations to the EPX were shown.

As noted in the introduction the eosinophil content of the alarmin EPX is associated to the EPX405 genotype with the highest content in the subjects carrying the EPX405CC genotype (ANOVA P = 0.009) (Figure 4). The eosinophil content of ECP was also linked to the EPX405 genotype (ANOVA P = 0.022), but in a reverse mode (Figure 4).

### Relationship to dysplasia/cancer

In the study UC cohort there were several patients (n = 27; females: 8; males: 19) that through continuous endoscopic examination were shown to have developed dysplasia and/or cancer (D/C) (median age at cancer diagnosis: 49 years; range: 27-67 years). The D/C cohort was compared to all UC patients, females and males who had not developed D/C, since no difference was shown between older and younger UC patients in the mixed cohort. The distributions of the ECP434 and ECP562 alleles in patients with UC, with and without findings of D/C of the intestine are shown in Figure 5. The C-allele of both genotypes was more prevalent among patients with D/C (ECP434 P = 0.034 and ECP562 P = 0.019). The D/C patients also differed from the healthy controls (ECP434 P = 0.027 and ECP562 P = 0.007) (Figure 5). The relative risk (RR), for UC patients with the ECP434 GC or CC genotypes, to develop D/C was 2.5 (95%CI, 1.2-5.4) compared to other patients with UC. In patients with the ECP562 GC or CC genotypes the RR was 2.5 (95%CI, 1.2-5.8) as compared to UC patients. Considering the whole cohort, no associations to the EPX405 (G > C) genotypes were detected.
DISCUSSION

Inflammatory bowel disease, including CD and UC, are idiopathic diseases with chronic inflammation of the gut. A key role of the eosinophil in the pathophysiology of IBD has been suggested. Our findings in this report, of intriguing associations between gene polymorphisms of the two major eosinophil granule proteins ECP and EPX, emphasize this notion. The primary results gained from the Uppsala cohort gave a hint that there was some association between the EPX and ECP polymorphisms and IBD. The associations, however, were in the larger cohort mainly found when the patients were separated into gender, age groups and diagnosis, since the associations were most apparent in older women with CD and in older men with UC. Another interesting observation was the higher

![Figure 3 Kaplan-Meier Curve of four groups haplotypes and EPX405, ECP562 genotypes. A: Kaplan-Meier Curve of four groups of haplotypes in male patients with ulcerative colitis (UC) (P = 0.0009); B: Kaplan-Meier Curve of four groups of haplotypes in female patients with Crohn’s disease (P = 0.003); C: Kaplan-Meier curve of the EPX405 genotype in men with UC (P = 0.007); D: Curve of the ECP434 genotype in men with UC (P = 0.002); E: Curve of the ECP562 genotype in males with UC (P = 0.01). Dotted line: GG genotype; Dashed line: GC genotype; Continuous line: CC genotype. ECP: Eosinophil cationic protein; EPX: Eosinophil protein X.](image-url)
prevalence of the ECP434C-allele in those patients with dysplasia/cancer, since this allele codes for the non-cytotoxic variant of ECP. We also confirmed the relationship between the EPX405 (G > C) polymorphism and the eosinophil content of EPX, since an increased content of EPX was found in subjects carrying the C-allele. The C-allele was also related to the age of disease diagnosis suggesting an impact of the alarmin EPX on the course of the disease in IBD.

In our study, we show that older females with CD had a relatively higher amount of the ECP434 GC and ECP562 GC genotypes as compared to healthy individuals, and younger females with CD. Our novel findings show that in healthy individuals and females diagnosed with CD at a young age the GG genotype of ECP434 and ECP562 was more dominating than in older women with CD, implying a more cytotoxic ECP and a higher content of ECP. This interesting finding could have a bearing on granuloma formation in CD. Granuloma formation is one of the pathological hallmarks of CD and is detected in the gut wall in 37%-60% of the patients\(^{14}\). Patients with granulomas, and particularly females, are diagnosed early in their course of disease\(^{14}\). In CD, the production of tumor growth factor (TGF)-\(\beta\) appears critical in CD-associated fibrosis and in the formation of granulomas\(^{16,17}\). ECP is known to promote TGF-\(\beta\) production and migration of fibroblasts\(^{18,19}\) and the ECP434 GG genotype has been linked to increased fibrosis\(^{20}\). All together this suggests a role of ECP in the production of granulomas in younger females with CD. Furthermore high numbers of bacteria, and mainly gram negative bacteria as *Escherichia coli* (*E*. coli), can induce secretion of ECP\(^{20}\). By the use of laser capture microscopy and PCR it was shown that DNA from *E*. coli was present in 80% of granulomas from patients with CD\(^{31}\).

There are few studies associating genetic risk factors and gender to falling ill with IBD. For patients with UC, there is a gender-specific risk associated with the interleukin-10 promoter. Females with a particular genotype and haplotype have an increased risk of contracting the disease earlier in life\(^{22}\). In patients with CD the DLG5 R30Q variant was shown to be associated with an increased risk for male patients to develop CD. This risk has been described both in the paediatric\(^{33}\) and the adult age group\(^{34}\). We showed that age at diagnosis in females with CD is linked to the EPX405 genotype. Females
with haplotype 13, including EPX405 GC, are diagnosed earlier in life. The longest span of years before onset of disease was observed in females carrying haplotype 4, including EPX405 GG. The C-allele of EPX405 is linked to a higher content of EPX in the eosinophils. The increased availability of EPX (in combination with unchanged amount of ECP) may result in an earlier disease onset in life. EPX was previously shown to present characteristics of an endogenous alarmin, which strengthens the immune systems Th2 immune response. CD has often been regarded as a disease with a dominant Th1 immune response, but a Th2 response in the early stages of the disease with active ileitis has been shown. EPX uses the toll-like receptor 2 (TLR2)-MyD88 signal transduction pathway to induce dendritic cell maturation and activation of NF-κB and also acts as a chemotactic activator of dendritic cells. Few of the intestinal dendritic cells in healthy controls express TLR2, whereas in patients with IBD the TLR2 receptor density is increased. NOD2 also induces nuclear factor κB (NF-κB) activation through the MyD88 pathway. The potentiated activation of NF-κB through the co-operation of EPX and NOD2 in CD, therefore, is an interesting possibility and may explain the differences found between CD and UC, since the 405C-allele was more common in CD in patients with shorter onset of disease.

In the cohort of older men with UC, the ECP434 and ECP562 GC genotypes were more common than in men with CD, females with UC, younger men with UC and the healthy population. Compared to younger men and the healthy reference population the EPX405 genotype was also significantly different, with more GC genotype in the older cohort. Thus, older men had a lower content of less cytotoxic ECP and a higher content of EPX. These results are similar to the cohort of older females with CD. Thus, from a genetic point of view, females with CD show similar distributions of the ECP and EPX SNPs as men with UC. These are interesting findings and suggest major gender differences in the pathogenesis of the two diseases.

In men with UC the Kaplan Meier analysis showed the lowest ADD in subjects carrying haplotype 27 (EPX405 CC, ECP434 CC and ECP562 CC) which codes for elevated production of EPX, less cytotoxic ECP and reduced production of ECP protein. The highest ADD was seen in those carrying haplotype 13 (EPX405 GC, ECP434 GC and ECP562 GG) which codes for increased production of EPX and cytotoxic ECP. These findings suggest a protective role of ECP, putatively against the microbial challenge, in the development of UC, whereas the role of the increased availability of EPX may be involved in the initiation of the disease. In contrast to the findings in men with UC, haplotype 13 was associated with the lowest ADD in women with CD, which should result in the increased production of cytotoxic ECP. As noted above this finding may relate to early granuloma formation. Our findings emphasize major differences in the pathophysiology and role of eosinophils in CD and UC. A notion supported by previous studies showing differences in the morphotype and function of eosinophils in CD and UC.

Our findings of an association between the ECP434 and 562 SNPs and UC with complication of dysplasia/cancer suggest a role of ECP in the defence against malignant cell transformation, since the results indicate a lower production of ECP and that the produced protein is non-cytotoxic and unable to kill malignant cells. The relative risk for a patient with UC and a C-allele in ECP434 or 562 to develop D/C is 2.5. When separating the cohort due to gender the EPX405 genotype was also significantly associated to D/C in males (data not shown). It is noticed that the number of patients with D/C was limited in our cohort, but recent studies have pointed out the association of ECP and EPX and cancer.

It is important to remember that the polymorphisms EPX405, ECP434 and ECP562 are in linkage disequilibrium (Jonsson et al to be published). Because of the profound linkage disequilibrium between the alleles, it is difficult to determine which allele that is the determining allele.

In conclusion, the present study has identified a gender and age related difference between polymorphisms in the EPX and ECP genes in CD and UC, and showed that the haplotype distributions are associated to the age of disease diagnosis. We also confirmed a link between the eosinophil content of EPX and the EPX405 gene polymorphism. Finally our results suggested a link between polymorphisms in the ECP gene and the risk in patients with UC of developing dysplasia/cancer. The role of neutrophils in IBD is a hallmark of the disease, and the results presented in this study suggest a key role of eosinophil granulocytes and their major granule proteins in the pathophysiology of inflammatory bowel disease.

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COMMENTS

Background

A role of eosinophils is implicated in inflammatory bowel disease. The intraluminal release of eosinophil derived cytotoxic proteins is increased in patients with ulcerative colitis (UC). Larger numbers of activated eosinophils, identified as CD44-high, are seen in quiescent as opposed to active UC. Gene polymorphisms in eosinophil protein X (EPX)/eosinophil derived neurotoxin and eosinophil cationic protein (ECP) affect the eosinophil content of EPX and ECP.

Research frontiers

The cytotoxic activities of ECP and EPX as well as their potent RNase activities have been well documented and are generally conceived as part of the host defence against viruses, parasites and helminths exerted by eosinophils. Polymorphisms in the genes of EPX and ECP are known to be linked to changes of cytotoxic affect and protein content. Increased release of EPX and ECP has been shown in patients with UC.

Innovations and breakthroughs

There are no previous studies focusing on the genetics of the eosinophil cationic proteins and inflammatory bowel disease (IBD). Studies of IBD and genetics in a gender and age dependent manner are few. With the present study, the
authors have combined the two, enhancing the relation between the eosinophils and IBD, and revealing the possible risk for cancer.

**Applications**

The study results have helped to attain an increased knowledge of the eosinophil's role in IBD with focus on age and gender related differences. The results also suggest that the determination of the EPX and ECP genotypes may contribute in patients risk stratification for cancer development.

**Terminology**

Eosinophil granulocyte: the eosinophil granulocyte is a pro inflammatory cell. The cell participates in wound healing and host defense against parasites and helminths. The cell is also present in the process of the allergic inflammation. Eosinophil cationic proteins: the cationic proteins are granula proteins secreted by activated eosinophils. The proteins are eosinophil specific and are therefore often used as markers for the eosinophil activities. Single nucleotide polymorphism (SNP): this is a variation in the DNA sequence, occurring when a single nucleotide (A, C, G or T) in the genome, differs between paired chromosomes in an individual or between individuals. If a SNP is present in the coding region of a gene this may result in an amino acid change, influencing the phenotype of a specific gene.

**Peer review**

The authors of this paper report that polymorphisms in the EPX and ECP genes are associated with IBD and contribute to the risk for the disease development in a gender-specific manner. Furthermore, these polymorphisms seem to be specifically related to the late-onset disease and to increased risk for colorectal cancer in UC patients. The results of this large study are certainly interesting, especially with the attention paid to gender-and age-stratified analysis which is a typical shortcoming of most of the genetic studies in IBD.

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