Clinical and genetical aspects of celiac disease

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Heredity

I am the family face;
Flesh perishes, I live on,
Projecting trait and trace
Through time to times anon,
And leaping from place to place
Over oblivion.

The years-heired feature that can
In curve and voice and eye
Despise the human span
Of durance — that is I;
The eternal thing in man,
That heeds no call to die.

Thomas Hardy

(First published in Moments of Vision
and Miscellaneous Verses, Macmillan, 1917)

To all the celiac children
Celiac disease (CD), or gluten-sensitive enteropathy, is one of the most common chronic diseases in childhood but is diagnosed in all ages. CD is a genetically driven immunological intolerance to dietary gluten. The treatment is a gluten-free diet. The diagnostic criteria are the ESPGHAN criteria, which include the histological characteristics of villous atrophy, crypt hyperplasia and increased number of intraepithelial lymphocytes (IEL). The clinical manifestations in CD range from severely affected young children to children and adults with milder symptoms as well as patients with silent CD. There is a strong heredity in CD with the well-known HLA components DQ2 and DQ8. The genetics in CD are believed to confer up to 40% HLA genetics and otherwise non-HLA genetics. The knowledge of the genotype-phenotype association in CD is limited.

The aim of this study has been to estimate the risk of a third sibling being affected in CD sib-pair families, identify the chromosomal region containing susceptibility genes in CD and study the genotype-phenotype association in CD.

Material was collected from 107 families with at least two affected siblings, making a total of 224 CD siblings, as well as their healthy siblings and parents. Screening for CD was performed in these apparently healthy members and the estimated risk for CD in the third sibling and parent was then calculated. Thirteen new CD cases were diagnosed, six siblings and seven parents. The estimated sibling risk was 26.3% and the parent risk was 12.9%. The risk of a sibling of two affected siblings having CD was approximately three times higher compared to siblings of one affected sibling. Considering the high level of knowledge of CD in these families, the number of undiagnosed cases was surprisingly high. We suggested that serological screening should be offered all first-degree relatives of CD patients.

Genome-wide linkage scan was performed in the same material. This work showed significant evidence of linkage to CD with an interesting region on chromosome 5q31-33 and on chromosome 11q. Simplex CD family material was collected for further genetic association studies.

The phenotype-genotype association was examined in two studies. An investigation was made of a possible interaction between the phenotypes and HLA class II risk alleles, the CTLA4 +49 A/G polymorphism, the haplotype MH30*G:-1147*T:+49*A;CT60*G;CT61*A and the 5q31-33 locus, in CD. The patients were grouped according to symptoms at presentation, the age at diagnosis and gender. The heritability of the phenotype was estimated to be 0.45. The AA genotype at the CTLA4 +49A/G polymorphism was associated with clinically silent disease. No other correlations were found between genotypes and clinical presentation, age at diagnosis or gender.

A genotype-phenotype analysis was made of phenotypes in DQ2-negative CD patients in the largest DQ2-negative CD group that has been published compared to DQ2-positive CD controls in a European population. The finding was that the clinical presentation differed significantly between DQ2-negative and DQ2-positive CD patients in Italy and Sweden. In both samples there was an association between DQ2-negative cases and classic symptoms. In the Italian sample there was also an association between silent grade and DQ2-negative cases. Autoimmune disease was significantly overrepresented in DQ8-positive patients. This thesis shows that the risk for third sibling and parents is, as expected, increased in sib-pair families, as the expected risk of being affected in polygenic diseases is higher in families with multiple cases compared to single-case families. The genome scan indicated significant linkage to 11q and 5q, which makes these regions interesting for further fine mapping of these regions using association analysis. Genotype-phenotype analysis of both HLA and non-HLA locus showed some significant correlation between silent CD and both CTLA4 +49 AA genotype and the DQ2-negatives. In addition, an association was shown between classic symptom grade and DQ2-negative cases.

Key words: celiac disease, sib-pair, screening, genome-wide scan, linkage analysis, heritability, genotypes, DQ2-negative, phenotypes, autoimmune disease


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Paper I-IV
List of publications


IV. Gudjónsdóttir AH, Nilsson S, Hugot J-P, Mustalhti K, Clot F, Coto I, Percopo S, Ascher A. Clinical features of DQ2-negative compared to DQ2-positive celiac disease. *In manuscript*
Abbreviations

AGA  Anti gliadin-antibodies
APC  Antigen presenting cell
ARA  IgA-class Reticulin antibody
CD  Celiac disease
cM  centiMorgan
CTLA4  Cytotoxic T lymphocyte-associated antigen-4
DH  Dermatitis herpetiformis
DNA  Deoxyribonucleic acid
ELISA  Enzyme-Linked ImmunoSorbent Assay
EMA  Anti-Endomysium antibody
ESPGHAN European Society of Paediatric Gastroenterology, Hepatology and Nutrition
GWA  Genome-wide association study
GWL  Genome-wide linkage study
GFD  Gluten Free Diet
HLA  Human leukocyte antigens
IBD  Identity By Descent
IEL  Intraepithelial T lymphocytes
LD  Linkage disequilibrium
MYO9B  Myosin IXB
NPL  Non parametric linkage
PCR  Polymerase chain reaction
RCD  Refractory sprue or celiac disease
RR  Relative risk
SNP  Single nucleotide polymorphism
TG2  Tissue Transglutaminase type 2
T1D  Type 1 Diabetes mellitus
TDT  Transmission disequilibrium test

Glossary


Allele: Alternative form of a genetic locus; a single allele for each locus is inherited from each parent.

Association: A tendency of two characters (disease, marker, alleles, gene) to occur together at non-random frequencies.

Candidate gene: A gene from the appropriate chromosomal location that is suspected of being the disease gene.
**CentiMorgan:** The unit of genetic distance. One cM is on average equal to about one mega base-pairs (1cM = 1Mb = 1,000,000 base-pairs).

**Epigenetic:** Heritable changes in gene expression that do not change the DNA sequence but rather provide an “extra” layer of transcriptional control that regulates how genes are expressed.

**Epistasis:** Genetic interaction, genes acting on the same or related biological pathway. The gene whose phenotype is expressed is said to be epistatic.

**Gene:** A unit of heredity, which is equal to a region of DNA.

**Genotype:** Genotype describes the genetic constitution of an individual, which is the specific allelic combination of the two homologous chromosomes.

**Gluten:** Prolamins or storage proteins in wheat.

**Haplotype:** A block of alleles that transmit together.

**Heritability:** The variance in the phenotype caused by genetic factors.

**Heterozygote:** The presence of different alleles at one or more loci on homologous chromosomes.

**Homozygote:** The presence of two copies of the same alleles at one or more loci on homologous chromosomes.

**Innate immune response:** A non-specific immune response to antigens, including anatomic and physiologic barriers, endocytic and phagocytic activity, and inflammatory secretions.

**Linkage:** A relationship between loci. Two loci are linked if they are located on the same chromosome.

**Linkage disequilibrium:** A non-random pattern of association between alleles at different loci within a population. Obtained when a particular marker allele is located so close to the disease susceptibility allele that, over generations, the two will be inherited together.

**Locus:** Is a specific position in the genome or on a chromosome.

**Microsatellite:** Polymorphic loci present in nuclear DNA that consist of repeated units of 1-4 base pairs in length, used as genetic markers.

**PCR:** A laboratory method used to amplify specific regions of a DNA strand.

**Phenotype:** The detectable outward manifestations of a specific genotype. Genotype and phenotype are not always directly correlated. Some genes only express a given phenotype in certain environmental conditions. Conversely, some phenotypes could be the result of multiple genotypes.

**Polygenic:** Polygenic inheritance is when many genes together influence the phenotype.

**Sib-pair families:** Two or more affected siblings.

**Transmission disequilibrium test:** If one allele increases risk of disease or trait, this allele will be transmitted to the affected offspring more often than expected by chance alone.
**Introduction**

CD or gluten-sensitive enteropathy is caused by dietary gluten ingestion in genetically susceptible individuals. Tissue transglutaminase type 2 (TG2) specific auto-antibodies are characteristic of CD, and increased TG2 activity has been observed in the small intestinal biopsies of patients. The immunological reaction that gluten induces causes a chronic immune reaction in the small intestine. CD is probably the best understood HLA disorder, as the environmental factor gluten is known. The non-HLA genetic and other environmental factors are still not fully understood in CD. There is some disagreement as to whether CD is an autoimmune disease or concurrent features typical of allergy and autoimmunity. The definition of autoimmune disease is that the tissue damage must be caused by an adaptive immune response to self-antigens [1-3]. Most authors today consider CD to be an autoimmune disease since TG2 antibody was identified in 1997 [4] as the main autoantigen for the anti-EMA. The approach of investigating genes in diseases is shown in a flow chart in Figure 1.

![Flow chart showing the approach of investigating genes in diseases.](image)

**History – milestones in celiac disease and genetic**

**Disease history**

The modern history of CD is short. Dr Samuel Gee (1839-1911) was an English paediatrician who in 1888 published in his thesis [5] the first complete modern description of the clinical picture of CD, and suggested that diet was important in controlling the disease. He stated: "If the patient can be cured at all, it must be by means of diet.” Christian Herter, an American physician, wrote a book in 1908 in which he called children with CD "intestinal infantilism" [6]. He noticed that their
growth was retarded and that fat was better tolerated than carbohydrate. In 1924, Sydney V. Haas, an American paediatrician, reported positive effects of a diet of bananas [7].

The next breakthrough was in 1950 when the Dutch paediatrician Dr Willem Dicke in his thesis showed that children with CD improved when eating a diet without wheat, rye, corn and oats [8]. Together with CM Anderson he later identified gluten as the harmful protein in CD [9, 10]. Studies in 1995 suggested the non-toxic effect of oats [11]. The first codex standard for a gluten-free product came in 1978 and a revised standard in 2001.

Shiner described in 1956 a method of making small intestinal biopsies on adult patients [12].

Serological markers were first described by E Berger in 1958 [13] but were studied and used more after 1970. The AGA antibodies were first used and connective tissue antibodies were later found, such as ARA and EMA (1983). In 1997, W Dietrich discovered the tissue Transglutaminase type 2, the unknown autoantigen of endemic antibodies [4].

MacDonald first described the hereditary character of CD in 1965 [14].

Genetic history

In 1953, Watson and Crick’s paper described the structure of DNA [15]. Further work by Crick and co-workers showed that the genetic code was based on non-overlapping triplets of bases, called codons, allowing Khorana, Holley and Nirenberg to decipher the genetic code and its function in protein synthesis. These findings represent the birth of molecular biology. In 1972, Cohen and Boyer created recombinant DNA, which is biotechnology that became important in the molecular biology. In 1975, the first generation of DNA markers appeared: restriction enzyme length polymorphism (RFLPs). The first gene to be mapped by positional cloning and linkage analysis was Duchenne muscular dystrophy in 1982. In 1983, Kary Mullis invented the PCR technique, which became very important for future genetic research [16]. Microsatellites came in 1989 and made genetic mapping more effective. The sequence of the human genome was published in 2001 [17, 18]. The same year the first non-HLA gene in autoimmune disease was published, NOD2 in Crohn’s disease [19].

Inheritance of celiac disease and risk

CD is one of the most common chronic diseases in Swedish children, surpassed only by asthma and allergies. In Sweden, childhood CD was increasingly diagnosed in the 1980s and early 1990s [20-22]. However, after 1996 the incidence suddenly decreased [23]. Other countries have seen increased prevalence and concluded that it is because of increased awareness and screening programmes [24].

Screening studies for CD in Sweden, Norway and other countries have shown prevalence figures varying between 2 and 6 per 1,000, in some studies even as high as 10 per 1,000 [25-31]. In an ongoing study, ETICS (http://www.umu.se/phmed/epidemi/celiaci/etics/) in Sweden, a screening study in 12-year-old children, has presented a prevalence as high as 30 per 1000 (95% CI: 25-33), Myléus et al presented
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at an International Coeliac Disease Meeting in Maribor in 2007. In North America, CD has been diagnosed increasingly during the last 15 years and a prevalence of 0.75% is found [32].

In family studies (Table 1, Paper I), sibling prevalence more than ten times higher than the prevalence found in population studies has been reported [33, 34]. Reported parent prevalence has varied between 0 and 6.3% [34-41]. Studies of twins have shown a concordance rate in monozygotic twins of at least 86.5% compared to 23% in dizygotic twins [42].

The average prevalence of CD among children with T1D in 26 reports was 4.5% (0.97-16.4%) [43]. A high prevalence of CD is found in some syndromes: in Downs's syndrome 3.6-8% in Europe [44-47] and as high as 10.3% in the USA [48, 49], in Turner 2.2-6.4% [50-52] and in Williams's syndrome 9.5% [53].

The CD iceberg

The iceberg model is often used to gain an epidemiological understanding of CD. The visible tip of the iceberg represents the diagnosed cases. The first part under the surface represents the undiagnosed or silent cases with gluten-induced enteropathy, CD patients found by screening. The deepest part represents the genetically predisposed individuals without gluten-induced enteropathy, called latent or potential CD, although the knowledge of when or if they will get CD is unknown.

Pathogenesis

The role of gluten

Gluten prolams are storage proteins in grain (wheat, barley and rye). Gluten proteins are a mixture of water-insoluble glutenins and the alcohol-soluble gliadins. Human dietary gluten is poorly digested in the upper gastrointestinal tract because of the high proline content. They are resistant to degradation by gastric, pancreatic and intestinal brush-border membrane proteases in the intestine and remain undigested in the intestinal lumen. TG2 is an enzyme in the intestine, found both at the brush border and just below the epithelium. A stress reaction, as a hypothesis via reduced zinc in the intestinal wall and increased Ca⁺ activity [54], activates TG2 which deamidates gliadin peptides. This results in proline-rich peptides containing negatively charged glutamic acid residues, which increase their immunogenicity. Gluten peptides are rich in the amino acid sequence QXP, which is an excellent substance for TG2. Gluten has many known immunogenic peptides identified in α-gliadins, γ-gliadins and the LMW and HMW-glutenins [55]. In CD patients, immune responses to gliadin fractions promote an inflammatory reaction in the upper small intestine and this cause the inflammation and tissue damage (villous atrophy) seen in gluten enteropathy.

Immunopathogenesis

Two immunological pathways are active in CD, the innate immune and the adaptive immune pathway. The gluten peptides are involved in both of them.

The innate immune system, where gluten peptides (α-gliadin) induce IEL (CD8+
cells) mediate enterocytes destruction, by expressing the natural killer receptors D (NKG2D). Gluten can even induce NKG2D expression by stimulating the expression of the cytokine IL-15.

The adaptive immune system, where TG2-modified gliadin residues connect to HLA-DQ2 and DQ8 molecules on APCs and present it to a T cell receptor on CD4+ lymphocytes, which results in an immune response with the formation of antibodies against both anti-gliadin and TG2 [1], Figure 2.

The IL15 produced by the APCs and enterocytes is also capable of stimulating IELs and the T-cells of the adaptive immune system and is therefore one of the links between the two pathways [56].

The mechanism of how gluten peptide crosses the epithelial barrier is still unknown.

Environmental factors
Breastfeeding duration, and if ongoing when gluten-containing foods are introduced to infants, may protect against the developing of CD or cause a delay in the onset of symptoms [24, 57-60].

Infections have always been suspected of being a risk factor for developing CD, a prospective study has shown that a high frequency of rotavirus infection may increase the risk of CD in DQ2-positive and DQ8-positive children [61]. The mechanism is not clear but could be due to a combination of increased intestinal permeability and increased presence of TG2 in the mucosa, which may facilitate the deamidation excreta or stress reaction. A study showing increased risk of CD in children younger than two years of age at diagnosis, born in the summer compared to the winter, may support the hypothesis that infections are involved in the pathogeneses [62] as their gluten introduction takes place during the winter.

Figure 2. The HLA-DQ2 and DQ8 αβ heterodimer are on the surface of antigen presenting cells. The binding site has a preference for negatively charged amino acids and thereby bind gluten peptides deamidated by TG2 with increased affinities and activate the CD4+ cells.

cis=alleles on the same chromosome
trans=alleles on different chromosomes

![Diagram of antigen presentation](image-url)
Introduction

Genetics in celiac disease

The strongest evidence of genetic factors in CD is the increased risk seen in family-based studies and twin studies as shown above. CD is a complex genetic disease.

HLA

The human leukocyte antigen (HLA) complex is located on the short arm of chromosome 6 (6p21.31). This region is gene-dense, with more than 200 gene loci. Forty per cent of the expressed genes encoded in the region have presumed immune system functions. Three regions are recognized: class II, encoding among others, the DP, DQ and DR molecules, class III, encoding among others the TNF family, and class I. HLA class I molecules are found on the surface of most nucleated cells in the body, while class II are present only on the surface of B cells, T cells and macrophages [15]. There are seven HLA class II DQ variants (DQ2 and DQ4-9). Two of these variants, HLA-DQ2 and DQ8, are associated with CD.

The function of the HLA class II proteins on the APC surface is to bind peptide fragments of processed protein antigens and present them to T-cell receptors on CD4+ lymphocytes of the adaptive immune system, Figure 2. The peptide-binding groove on HLA class II molecules is composed of antiparallel β-sheets as a floor and antiparallel α-helices as walls. Small cavities in the groove bear polymorphic amino acids responsible for the peptide-binding specificity. The binding site has a preference for negatively charged side chains.

HLA association in CD

Genetically, CD is characterised by a strong HLA class II component (CELIAC 1). The HLA component consists of a combination of specific alleles at the DQ locus. The HLA haplotypes and serological typing is shown in Table 1.

Table 1. Haplotypes that predispose to celiac disease, the DQ2 and DQ8 heterodimers.

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<th>Haplotype I</th>
<th>Haplotype II</th>
<th>Serological typing notation</th>
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<tr>
<td>DRB1 DQA1 DQB1</td>
<td>DRB1 DQA1 DQB1</td>
<td>DR3-DQ2</td>
</tr>
<tr>
<td>03 05 02 - - -</td>
<td>03 05 02 03 05 02</td>
<td>DR3-DQ2/DR3-DQ2</td>
</tr>
<tr>
<td>03 05 02 07 0201 02</td>
<td>03 05 02 07 0201 02</td>
<td>DR3-DQ2/DR7-DQ2</td>
</tr>
<tr>
<td>11/12 05 0301 07 0201 02</td>
<td>04 03 0302 - -</td>
<td>DR5-DQ7/DR7-DQ2</td>
</tr>
<tr>
<td>04 03 0302 - -</td>
<td>- -</td>
<td>DR4-DQ8</td>
</tr>
</tbody>
</table>

This HLA DQ2 molecule is either encoded in cis in individuals who have the DR3–DQ2 haplotype, or in trans in individuals who are DR5–DQ7/DR7–DQ2 heterozygous. An increased risk of CD in individuals who are DR3–DQ2 homozygous and DR3-DQ2/DR7-DQ2 heterozygous has been demonstrated [63]. A small proportion of the CD patients have neither DQ2 nor DQ8 and these patients almost
exclusively encode for one half of the DQ2 heterodimer, *i.e.* either DQA1*05 or DQB1*02 [64]. In the European population more than 90% of the CD patients carry HLA-DQA1*05-DQB1*02 encoding the DQ2, compared to as many as 20-30% of the healthy population being carriers of DQ2, Figure 3 [15, 65, 66]. Almost all the patients that are DQ2-negative carry the DR4-DQ8.

![Figure 3. Incidence of HLA DQ2 and DQ8 in the European population and in the CD population.](image)

The possibility of other risk loci in the HLA complex has been studied. The DQ-independent risk effects are relatively difficult to reveal due to the strong linkage disequilibrium in the whole region and this question still remains unsolved [67, 68]. A GWA study showed a strong association with the HLA-DQ2 [69]. Even though the HLA genes are considered necessary for CD some studies have estimated that the HLA DQ genes only contribute with 40% of the genetic risk in CD, indicating the importance of genes outside the HLA complex [70].

### Non-HLA candidate genes and regions

Outside the HLA class II region, some other genes and regions of interest have been studied. The 5q31-33 locus (CELIAC 2), the CTLA4 +49 gene on 2q34 (CELIAC 3), the MYO9B on 19q13.1 locus (CELIAC 4) and the 4q27 (IL2-IL21 genes) have been linked or associated with CD.

**CTLA4 +49** is shown to be associated with CD in several studies and populations [71-74] but not in others [75]. In the GWA study there was a weak association with the CD28-CTLA4-ICOS region [69].

As CTLA4 +49, a co-stimulatory molecule of the T cells, plays an important role in maintaining immunological tolerance to self-antigens, it is an interesting candidate gene. Some studies have implicated a functional role of CTLA4 +49. The G allele of the CTLA4 +49 A/G dimorphism has been shown to be associated with reduced control of T cell proliferation [76]. The role of CTLA4 +49 in CD is not clear, but more likely minor.

The 5q31-33 region on chromosome 5 was the strongest region shown in the meta-analysis and pooled analysis of four genome scans [77]. Further association studies have, however, failed to find a strong susceptibility candidate [78]. There are 200
genes in this region, many of immunological importance. Some have been analysed, e.g. IL4, IL5, IL12, IL13 and IL14, without significant association in CD [79, 80]. On 19q13.1 the MYO9B gene was shown to be associated with CD in two independent cohorts in the Dutch population [81]. However, this finding has been difficult to replicate in other populations included in the Swedish/Norwegian study or in the GWA study [69, 82-84]. MYO9B might play a role by affecting the tight junction and epithelial barrier function of the gut and increase permeability of the gut, allowing the gluten peptides to enter the lamina propria more easily [85].

The strongest association in the recent published GWA study was shown to the KIAA1109/Tenr/IL2/IL21 region on 4q27 [69]. This finding has been verified by our group in Swedish and Norwegian patients [86]. This region is of immunological interest since IL2 is a cytokine for T-cell activation and proliferation and IL21 is a cytokine that enhances B-, T- and NK-cell proliferation as well as interferon-γ production.

**Clinical aspects**

**Diagnostic criteria**

The first diagnostic criteria were discussed at the second Annual Meeting of ESPGHAN (ESPGA at that time) in Interlaken in 1969. The Interlaken or ESPGHAN 70 criteria were published [87] and contain three small intestinal biopsies, characteristic histological appearance of the mucosa, its normalization on a gluten-free diet (GFD) and a histological relapse within two years on reintroduction of gluten. A new round table discussion on the diagnostic criteria took place at the 22nd Annual Meeting of ESPGHAN in Budapest in 1989. The revision of the Interlaken criteria or ESPGHAN 90 criteria was published in 1990 [88]. The ESPGHAN 90 criteria abandoned the previous obligatory gluten challenge and the requirement of three biopsies, in cases with typical first small intestinal histology and a clear remission on GFD and serological normalization. Otherwise the investigation must be continued until the CD diagnosis is confirmed or can be excluded. However, a gluten challenge with a subsequent biopsy has a role in establishing the diagnosis in select clinical settings, such as in patients with a high suspicion of CD who have started with a GFD without biopsy confirmation of the disease, or of a negative serologic test result.

**Serological markers**

The first serological screening tests were described in the 1960s. Anti-gliadin antibodies (AGA) in serum were the most used antibodies in the earlier research and clinical practice and are still used in clinical practice in combination with the newer antibodies [89].

Later, anti-Endomysium antibody (IgA-EMA), directed against intermyofibril connective tissue of smooth muscle fibres, became available [90]. From the studies IgA-EMA demonstrated sensitivities of 97.4% (95% CI: 0.957-0.985) in adults and 96.1% (95% CI: 0.945-0.973) in children. The specificity of IgA-EMA was 99.6% (95% CI: 0.988-0.999) in adults. In studies of children, the specificity of IgA-EMA was 97.4% (95% CI: 0.963-0.982) [91].
TG2, the main auto-antigen for EMA, was identified in 1997 [4]. TG2 is a Ca\(^{2+}\)-dependent enzyme that is responsible for converting glutamine residuals to glutamic acid by deamidation when the amine is replaced by water [92]. An ELISA method using guinea pig IgA-TG2 was used earlier but now human IgA-TG2 is the method of choice. It has been shown to have a sensitivity of 96-100% and a specificity of 96-100% [93, 94] although in a systemic review the sensitivity of human TG2 was similar: in adults 98.1% (95% CI: 0.901-0.997) and in children 95.7% (95% CI: 0.903-0.981) with a specificity of 98.1% (95% CI: 0.958-0.991) in adults and 99% (95% CI: 0.946-0.996) in children [91].

Estimates of the sensitivity of the IgG class antibodies of EMA and TG2 suggest that these tests have poor sensitivities, around 40% (95% CI: 0.363-0.543), although the specificities were high at around 98.8% (95% CI: 0.935-0.998) [91].

Recently, a new ELISA test of antibodies against deamidated gliadin peptides (DGP) has been developed [95]. A few studies have been published, indicating a sensitivity and specificity for IgA-DGP of 83.6-91% and 90.3%-95%, and for IgG-DGP of 84.4-92% and 98.5%-99% respectively. The sensitivity for a combination of IgAG-DGP/IgA-TG2 was 100% [96, 97].

However, all tests have overlapping confidential intervals [89, 91] that need to be taken into account when using them. When screening studies of risk groups, priority must be given to sensitivity although when screening unselected population tests with a high specificity should be given priority.

**Histopathology**

The histological criteria for celiac diagnosis includes villous atrophy, crypt hyperplasia, an increased number of intraepithelial T lymphocytes (IEL) [98], or >20-30 lymphocytes/100 enterocytes, and inflammation of the lamina propria. Different classifications have been used to make the examination as objective as possible. The Alexander classification has four grades based on stereomicroscopic appearance and histological characteristics [99], where grades 3 and 4 fulfil CD criteria. Table 2 shows the modified version. The Marsh classification has types 0-III, where type III fulfils the CD criteria. The subgroups are a, b and c as seen in Table 3 [100, 101].

<table>
<thead>
<tr>
<th>Grade</th>
<th>Histological characteristics</th>
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<tr>
<td>1</td>
<td>Normal</td>
</tr>
<tr>
<td>2</td>
<td>Normal villous length and crypt depth (villous/crypt ratio ≥ 2). Increased inflammation in lamina propria. Increased IEL.</td>
</tr>
<tr>
<td>3</td>
<td>Subtotal villous atrophy, elongation of crypts (villous/crypt ratio &lt; 2). Increased inflammation in the lamina propria. Increased IEL.</td>
</tr>
<tr>
<td>4</td>
<td>Total villous atrophy, elongation of crypts. Increased inflammation in the lamina propria. Increased IEL.</td>
</tr>
</tbody>
</table>
Introduction

Immunohistochemistry examination could be used in uncertain cases. In CD, 20–30% of IELs bear \( \gamma\delta \) T-cell receptor-bearing cells, which comprise less than 10% of the IELs in non-celiac subjects [102].

Clinical manifestation

CD has many clinical manifestations, ranging from severely ill young children to children and adults with milder symptoms as well as asymptomatic patients or silent CD [103]. The clinically silent patients are defined as if they had not complained of illness before screening was performed, but some can in a later examination have a low grade of the illness, mainly diagnosed by screening risk groups or in screening studies. Silent CD is an interesting manifestation and more studies are presenting changing features in CD and more silent patients [104-107].

As a result of increased knowledge of the different clinical manifestations, awareness of the disease and widespread screening, more children and adults are now being diagnosed. Despite the fact that undiagnosed cases are many, population screening studies show that the majority of individuals with CD are undiagnosed [25, 108, 109].

The main symptoms of CD in children [110] are gastrointestinal, such as chronic diarrhoea, steatorrhea and abdominal pain, constipation, vomiting and bloating. Growth retardation is usual in children.

The more “classic” CD, characterised by weight loss due to malabsorption, diarrhoea, fatigue and development stagnation, are more common in younger children.

CD in school age presents more often with abdominal pain, abnormal linear growth and lack of puberty [104]. The clinical manifestation in children has changed to milder symptoms and later diagnosis in Sweden [105], Italy [107], the UK [106] and the Netherlands [24].

In adults, the clinical manifestations are usually milder and more systemic. Presentation in the form of diarrhoea has decreased (73% to 43%) and the time for onset of symptoms has decreased (9 years to 4.4 years). Common non-gastrointestinal symptoms are weight loss, fatigue or tiredness. Adults often have atypical symptoms from different organs [111]. CD is twice as common in females as it is in males [23].

Table 3. Marsh classification for histology of the small intestinal biopsies.

<table>
<thead>
<tr>
<th>Type</th>
<th>Histological characteristics</th>
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<tr>
<td>0</td>
<td>Normal</td>
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<tr>
<td>I</td>
<td>Normal villous architecture. Normal crypt depth. Increased IEL.</td>
</tr>
<tr>
<td>II</td>
<td>Normal villous architecture. Crypt hyperplasia. Increased IEL.</td>
</tr>
<tr>
<td>III</td>
<td>Villous atrophy. Crypt hyperplasia. Increased IEL.</td>
</tr>
<tr>
<td>a.</td>
<td>Partial villous atrophy</td>
</tr>
<tr>
<td>b.</td>
<td>Subtotal villous atrophy</td>
</tr>
<tr>
<td>c.</td>
<td>Total villous atrophy</td>
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</table>

Immunohistochemistry examination could be used in uncertain cases. In CD, 20–30% of IELs bear \( \gamma\delta \) T-cell receptor-bearing cells, which comprise less than 10% of the IELs in non-celiac subjects [102].

Clinical manifestation

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**Introduction**

*Dermatitis herpetiformis* (DH) was first described in 1884 by Duhring, although Marks et al [112] described the association with gluten-sensitive enteropathy in 1966. Skin symptoms, with itching blisters, appear on the elbows, knees and buttocks. Studies have shown that 25% of DH patients have only increased intraepithelial lymphocytes or normal mucosa and a lack of antibodies [113, 114]. GFD is the treatment of choice [115]. DH even occurs in children [114] but is probably often missed.

Disorders associated with CD are many, listed in Table 4.

<table>
<thead>
<tr>
<th>Table 4. Disorders associated with celiac disease.</th>
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<tbody>
<tr>
<td><strong>Oral manifestation</strong></td>
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<td><strong>Bone and connective tissue</strong></td>
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<td><strong>Skin</strong></td>
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<td><strong>Haematological</strong></td>
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<td><strong>Liver diseases</strong></td>
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<td><strong>Endocrinological</strong></td>
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<td><strong>Gynaecological</strong></td>
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<td><strong>Neurological and psychological disturbances</strong></td>
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<td><strong>Other diseases</strong></td>
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Multiple haematological manifestations are associated with CD as anaemia with iron deficiency (46%), B12 deficiency (8-41%) or folic acid deficiency, often without anaemia in children, as well as thrombocytosis (60%), thrombocytopenia (rare), leucopenia (rare), coagulopathy (18.5% prolonged prothrombin time), venous thromboembolism (rare), hyposplenism (21%), and IgA deficiency (2-3%) [116]. IgA-deficiency patients have around a 10% risk of developing CD. The clinical forms are not different and other autoimmune diseases are also increased [117, 118]. Bone mineral density has been found to be lower in untreated CD patients, and in older children, especially these of short stature [119, 120]. Complete recovery is shown on GFD after one year in children [121] and in most adults [122]. Oral manifestations have been reported. Dental enamel defects are the oral lesions most closely related to CD [123, 124]. If systematic, the patients should be screened for CD even in the absence of gastrointestinal symptoms. There are conflicting data on the association between CD and recurrent aphthous stomatitis [125, 126]. A correlation of CD to atrophic glossitis may occur but is seen more often with other conditions, such as B12, iron, folic acid deficiency [127]. Neurological and psychological disturbances are associated with CD, the most common being ataxia, peripheral neuropathy, encephalopathy and myopathy, but also epilepsy without or with the described cerebral calcification, headache and depression. Gluten-related neurological disease is treated with GFD and has a better prognosis if diagnosed early [128, 129].

Autoimmune diseases have been associated with CD [130]. Ventura et al presented a study showing a prevalence of 34% for autoimmune diseases in adults CD and correlation to the duration of gluten exposure [131], although other studies have not been able to confirm this correlation to the duration of gluten exposure [132-134]. Many autoimmune diseases have a CD preponderance to females. The mechanism of this association of autoimmune disease to CD is still a subject for discussion – whether it is secondary to linkage disequilibrium of genes predisposing for both or if CD leads to the onset of other autoimmune diseases in genetically susceptible individuals.

Type 1 diabetes (T1D) is associated with CD in many studies. The average prevalence of CD among children with T1D varies widely, with an average in different studies of 4.1% (0.97-16.5%) [43]. The risk is regardless of which disease is diagnosed first but more common, in 90% of the cases, is that T1D is diagnosed first [135]. The symptom form of CD is often mild or silent in T1D. Meta-analysis for T1D has shown the well-known increased risk of DR3-DQ2/DR4-DQ8, with predisposing to DR4-DQ8 subtypes [136]. T1D is associated with the gene-encoding CTLA-4 +49 [137].

Autoimmune thyroid diseases are associated with CD with a risk of 2-5%. These conditions share similar HLA haplotypes and are associated with CTLA4 +49 [138]. Autoimmune Addison’s disease is increased in CD both in children and adults [139, 140]. Liver diseases, such as primary biliary cirrhosis, autoimmune hepatitis and autoimmune sclerosing cholangitis, are associated with CD although isolated hypertransaminasaemia with non-specific histological changes in a liver biopsy is found.
in 9% of CD as well as more severe liver damage [141, 142].

Pancreas diseases, such as pancreatitis and pancreatic exocrine dysfunction, are found to be associated with CD [143].

Rheumatoid arthritis has been found to be associated with CD even if other studies have failed to show an association [144, 145].

CD is associated with an increased risk of developing malignancy, especially “enteropathy-associated T-cell lymphoma” (EATL) and other gastrointestinal cancers, such as small bowel adenoma. The overall risk of non-Hodgkin’s lymphoma is much lower than previously thought (relative risk of 2-4%) [146, 147]. In ESPGHAN’s inventory of combined paediatric CD and malignancy only 22 cases were found, including three cases of thyroid carcinoma and five of small intestinal lymphoma (four B-cell lymphoma of the Burkitt type and one sarcoma) [148]. There is compelling evidence to suggest that the GFD protects against the development of CD-associated malignancies, especially if started early in life [149].

Refractory CD or refractory sprue, defined as persistent symptoms and villous atrophy despite scrupulous adherence to a GFD, has a rate of occurrence of approximately 5% in adults. RCD can be categorized into type I or type II. Type I RCD has a more favourable prognosis compared with type II. Type II RCD carries a poor prognosis and is more likely to progress to life-threatening malnutrition or intestinal T-cell lymphoma [150].

**Treatment**

**Gluten free diet**

GFD is the current treatment for CD. The Codex Alimentarius Commission was created in 1963 by the FAO and WHO to develop food standards, guidelines and related texts, such as codes of practice, under the Joint FAO/WHO Food Standards Programme. Gluten-free products consist of ingredients that do not contain any prolamin from wheat, rye or barley, with a gluten level not exceeding 20 ppm or consisting of ingredients that contain any prolamin from wheat, rye or barley, with a gluten level not exceeding 200 ppm.

The knowledge of the non-toxic effect of oats came with the first studies in 1995 [11] and follow-up studies, both clinical and immunological, have shown that eating oats is not harmful to adults [151-153]. After studies on children [154] oats were allowed in the GFD for CD children in Sweden in 2004 [155].

**Future treatments**

Living under the regime of a life-long gluten-free diet is burdensome as wheat and related cereals are very commonly used in the food industry. Moreover, the presence of gluten in food is not always obvious. Patients ask for alternative treatment. Future treatments demand to be as safe and effective as GFD. Alternative therapies are therefore of interest. Ongoing studies deal with different types of “blockers” of TG2, the DQ2-mediated antigen presentation or IL15 [156] and the efficiency of gluten degradation by post-proline cutting enzyme [157]. Attempts to generate wheat varieties have not been successful [158].
**Molecular genetics**

**The human genome**

In 1980, the human genome project started. The work increased in 1995 and it was completed in 2001 [17, 18].

Our genetic information is stored in **chromosomes**, which are made up of DNA (deoxyribonucleic acid) [10] and genes, are special units of chromosomal DNA. A human has 23 pairs of chromosomes that vary widely in size and shape. Chromosome 1 is the largest and is over three times bigger than chromosome 22. The 23rd pair of chromosomes determines our sex: females XX and males XY. Each chromosome is a very long molecule and so it needs to be wrapped tightly around proteins for efficient packaging of the DNA double helix. Near the centre of each chromosome is its centromere, a narrow region that divides the chromosome into a long arm (q) and a short arm (p). The chromosomes can be divided further using special stains that produce stripes known as a banding pattern. Each chromosome has a distinct banding pattern, and each band is numbered to help identify a particular region of a chromosome. This method of mapping a gene to a particular band of the chromosome is called cytogenetic mapping. The CTLA4/CD28 region, for example, is found on 2q33 and the HLA is found on 6p21.31 [159].

The **DNA** is about 2 metres long, and 3 cm or 1.5% of the human genome consists of protein-coding exons. However, DNA sequences that do not code protein may still encode functional non-coding RNA molecules, which are involved in the regulation of gene expression. Genomic DNA is located in the cell nucleus of eukaryotes, as well as small amounts in mitochondria (37 genes). A **gene** is a unit of heredity and is a region of DNA, and the complete set or parts of this information in an organism is called its **genotype**. Humans have paired homologous chromosomes in their somatic cells and these contain two copies of each gene. A person who has two copies of the gene that are identical, i.e. have the same **allele**, is described as being **homozygous** for that gene. A person who has two different alleles of the gene is described as being **heterozygous**.

The DNA double helix is stabilized by hydrogen bonds between the bases attached to the two strands. The four bases found in DNA are adenine (abbreviated A), cytosine (C), guanine (G) and thymine (T). These four bases are attached to the sugar/phosphate to form the complete nucleotide. These bases are classified into two types: adenine and guanine are fused five- and six-membered heterocyclic compounds called purines, while cytosine and thymine are six-membered rings called pyrimidines [159]. Of the three billion base pairs we have 1/1000 SNPs or three million SNPs that have been located. In genetic studies we look at different variances. Only a small part of the DNA has a known function although our knowledge has increased more rapidly during the past year than ever before. Consortium researchers have confirmed the existence of 19,599 protein-coding genes in the human genome and identified another 2,188 DNA segments that are predicted to be protein-coding genes [160].
Recombination

Chromosomal crossover is the process by which two chromosomes, paired up during prophase 1 of meiosis of a germ cell, an egg or a sperm, exchange some portion of their DNA. Crossover usually occurs when matching regions or matching chromosomes break and then reconnect to the other chromosome. The result of this process is an exchange of genes, called genetic recombination, Figure 4.

This process leads to offspring having different combinations of genes from their parents. Recombination between two loci on chromosome has occurred if one is of maternal and the other of paternal origin. Genetic recombination is necessary for linkage analysis to be able to locate genes on different parts of a chromosome.

The probability of recombination is called recombination fraction; it is a measure of the genetic distance between two loci, measured in centiMorgan (cM). One cM is equal to a 1% chance that a marker at one genetic locus on a chromosome will be separated from a marker at a second locus due to crossing over in a single generation.

Markers

A genetic marker is a known DNA sequence. It can be described as a variation, which may arise due to mutation or alteration in the genomic loci, which can be observed. A genetic marker may be a short DNA sequence, such as a single base-pair change as SNP (single nucleotide polymorphism), or a long one, such as microsatellites that are 1-4 nucleotide long segment repeats. Microsatellites vary between humans and occur throughout the genome. SNP genotyping is the process of determining the single nucleotide polymorphisms of an individual. Tag SNP is a representative single nucleotide polymorphism in a region of the genome with high linkage disequilibrium to other SNPs in the region.

PCR can amplify the markers and fluorescing molecules can make these visible. The Hap Map project, http://www.hapmap.org, now makes it possible to select a limited number of SNPs or CNVs (copy number variant) to capture most of the variation in a given segment of the genome [161].

Figure 4. Recombination. Marker A and B can make crossover, in sibling II there is recombination and in sibling III, non-recombination.
Introduction

Complex traits

When the genome was mapped the possibility of carrying out a genome-wide scan for diseases that do not follow the Mendelian segregation or pedigree patterns became a reality. This requires interdisciplinary co-operation between clinicians, geneticists and statisticians.

CD is a disease with a complex inheritance pattern, complex trait or polygenic trait, where it is difficult to see the pedigree patterns. It is then difficult to know anything in detail about disease allele frequencies and penetrances. The phenotypes, clinical manifestation, are many. A complex trait disease is likely to be associated with the effects of multiple genes in combination with lifestyle and environmental factors. The correlation between genotype and disease phenotype can be weak.

Up until last year the history of human genes showed that a few genes of complex traits have been found [162]. Now it seems to be changing. New techniques have made it possible to perform genome-wide association (GWA) studies using SNPs with a 1000-fold greater density compared to the older genome-wide linkage studies. GWA studies use arrays that can examine some 500,000 SNPs at a time, in very large study materials. By tallying which SNPs co-occur with symptoms, the risk associated with each SNP can be determined. This new technique has meant that last year researchers linked variants of more than 50 genes to an increased risk of a dozen diseases. Identifying the relevant genes has been difficult, partly because each causal gene only makes a small contribution to the overall heritability [160]. The new studies have raised the hope that genes influencing complex trait diseases are about to be discovered, such as for diabetes, heart diseases, many autoimmune diseases and infection diseases such as AIDS. The first GWA study on CD has been published, showing evidence that the IL2 and IL21 regions is involved in CD [69].

Genetic analysis

Linkage describes the relationship between different loci on the chromosomes. It is a method used to map genes as searching though the entire genome base by base is not possible in practice. The principle of linkage is the proximity of two or more markers on a chromosome; the closer the markers, the lower the probability that they will be separated during meiosis, and hence the greater the probability that they will be inherited together. That means that if a marker is located beside an inherited disease gene it will more often than not be inherited together with the gene. Genome-wide linkage studies can identify chromosome regions that bring susceptibility genes by examining segregation at several 100 to 1000 loci across the genome.

Association between genetic polymorphisms on the other hand describes the relationship between alleles at different loci and is a statistical statement about the co-occurrence of alleles or phenotypes. Two alleles are genetically associated if they appear together more frequently than we would expect from the allele frequencies. The most common reason for genetic association is Linkage Disequilibrium (LD), which is the
Introduction

Non-random association of alleles at two or more loci. LD describes a situation in which some combinations of alleles or genetic variants occur more or less frequently in a population than would be expected from a random formation of haplotypes from alleles based on their frequencies. Non-random associations between alleles at different loci are measured by the degree of LD. LD is generally caused by genetic linkage and the rate of recombination, the rate of mutation, the random drift, or non-random mating, and the population structure. It is not the same as linkage, which describes the association of two or more loci on a chromosome with limited recombination between them.

HLA-DQ2, for example, is found in 20-25% of the general Scandinavian population but in over 90% of people with CD and is therefore strongly associated with CD.

An association could have many possible causes, not all of them genetic. Linkage, on the other hand, is a specific genetic relationship between loci (not alleles or phenotypes). Linkage does not in itself produce any association in the general population. The B locus, for example, is linked to the C locus. Within a family where a C mutation is segregating, we would expect affected people to have the same allele of B, but over the whole population the distribution of B alleles is just the same in people with and without C. Linkage thus creates associations within families, but not among unrelated people. However, if two supposedly unrelated people with disease D have actually inherited it from a distant common ancestor, they may well also tend to share particular ancestral alleles at loci closely linked to D. Where the family and the population merge, linkage and association merge.

**Summary:** Linkage studies localize chromosomal regions containing disease genes by investigating co-inheritance of genetic markers and disease in the families. Association studies assess if a specific allele is more or less frequent in affected individuals. A marker associated with a disease is either the causative gene itself or it is in LD with the causative gene [163].

**Genotype-phenotype association**

Humans give their children their environment as well as their genotypes. A genotype is often the largest influencing factor in the development of its phenotype although it is not the only one. Monozygous twins share the same genotype but they have never exactly the same phenotype. Phenotype describes mostly as alternative A in Figure 5. But our knowledge is changing and maybe alternative B is closer to the reality.

Epigenetic is heritable changes in gene expression that do not change the DNA sequence but rather provide an “extra” layer of transcriptional control that regulates how genes are expressed. Our understanding of the mechanism of the interplay between epigenetic gene expression and the environment is still finite. Epigenetic research aims to understand heritable gene regulation that is not directly encoded in the DNA sequence. Epigenetic mechanisms, such as DNA methylation and histone modifications, modulate the packaging of the DNA in the nucleus and thereby influence gene expression. Patterns of epigenetic information are faithfully propagated...
Introduction

over multiple-cell divisions, which make epigenetic regulation a key mechanism for cellular differentiation and cell fate decisions. In addition, incomplete erasure of epigenetic information can lead to complex patterns of non-Mendelian inheritance. Stochastic (when two persons do not react the same to treatment for the same diagnosis) and environment-induced epigenetic defects are known to play a major role in cancer and ageing, and they may also contribute to mental disorders and autoimmune diseases [164].

The knowledge of the interaction between genotype and phenotype in CD is limited. Several studies have failed to establish associations between different HLA-genotypes and phenotypes. In an Italian study of 145 CD patients, including 27 silent patients, no correlation was found between HLA-genotypes and phenotypes [165]. In this study there were ten DQ8-positives, two of them asymptomatic, and five DQ2-negative/DQ8-negatives, all of them symptomatic. In other 28 symptom-discordant sib-pairs no association between symptoms and DR-DQ haplotypes was found. Of the patients, 25% were DQ2 homozygous. The symptomatic CD seemed to have earlier onset than silent CD [166]. In two studies however, a gene dose effect of DQB1*02 on the phenotype was found. Homozygous patients had a moderate overrepresentation of classic presentation, female gender, lower age at diagnosis and a shorter delay between onset of symptoms and diagnosis. In this study four patients were DQ8-positive patients, all had classic symptoms, and of four DQ2-DQ8-negative patients one had classic symptoms, two had atypical symptoms and one was asymptomatic [167]. In the other study DQB1*02 homozygous were correlated to severity of the villous atrophy, lower age at diagnosis and lower hemoglobin values at diagnosis [168].

In the clinical presentation of 25 DQ2-negative patients in a study of Finnish and Spanish patients [169], all were assessed as typical of CD. Of the 25 DQ2-negative patients there were two patients without any part of DQ2 or DQ8. They did not differ clinically from the other patients.

One study showed no association between IL12B and IRF1 genes, in the 5q31-33 region, and enteropathy grading according to the Marsh criteria [170].

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**Figure 5.** The phenotype of human is influenced directly by its genotype and its environment. In addition, these two factors indirectly influence phenotype via the epigenotype [164].
The role of epigenetic in CD is unknown although Megiorni et al showed a major distortion in the DR3-DQ2 transmission from fathers to daughters and suggested a possible different effect of parent-specific epigenetic modifications in the two genders [171].
The role played by environmental factors in CD is still not clear – if, for example, breastfeeding prevents or simply delays symptoms or the role of infections. See above.
Aim of the study

The aim of the thesis was to:

Screen for CD in apparently healthy members of nuclear families with two affected siblings and to estimate the risk of CD in the remaining siblings and parents.

Perform a genome-wide scan in a population of Scandinavian families with CD, to identify non-HLA candidate chromosomal regions showing linkage to CD.

Investigate the heritability of the phenotype in CD and the influence on the phenotype of different genes associated with the disease.

Compare the clinical presentation between DQ2-negative and DQ2-positive CD patients in a European population.
Patients and Methods

Patients

Multiplex families (Paper I, II, III and IV)
Families with at least two affected siblings were collected from Sweden and southern Norway in order to perform a genome-wide scan for celiac genes and genotype-phenotype association studies (Papers I-IV).
The aim was to collect blood samples and clinical data from at least one hundred affected sib-pairs, their parents and their healthy siblings if any.
Three lines of action were used for recruiting families: 1. Advertisement in the journal of The Coeliac Society of Sweden. 2. Inquiries to all Swedish paediatric departments. 3. Invitation to all CD sib-pair families of the registries of our own paediatric departments. By collecting the families in these different ways more sib-pair families were recruited and the range of clinical presentation in the material was wider.
A semi-structured telephone interview was performed when entering the study. The interview included questions concerning the diagnosis, the gluten consumption, general health and possible symptoms of parents and siblings without CD diagnosis.
In order to avoid inclusion of falsely diagnosed cases, strict inclusion and exclusion criteria were applied. Medical records of individuals with diagnosed CD were collected and scrutinized. Diagnostic data and clinical manifestations were recorded. Only families where all CD siblings were diagnosed according to the ESPGHAN 90 criteria [88] were included. Children under the age of two years at the time of diagnosis were included in the study only if they fulfilled the original ESPGHAN 70 [87] with three biopsies on different diets. Individuals on GFD without CD diagnosis were excluded.
Blood samples for DNA extraction and serological screening were collected from all CD patients, the healthy parents and healthy siblings available.
Of the 152 families recruited, 113 families met the inclusion criteria. Six families dropped out and the remaining 107 families with two to four affected siblings met the diagnostic criteria and were included in the study (Paper I, Figure 1 and Table 2). Of 102 additional siblings, eight were not included in the study: seven were living abroad and one was excluded because he was on a gluten-free diet without diagnosed CD.
In the screening study (Paper I), families without healthy siblings available for participation in the study were excluded as well as individuals without CD diagnosis not eating a gluten-containing diet. Altogether, 65 families (56 with two affected, 8 with three affected and one with four affected) with 94 siblings were included in the calculations of the risk of CD in a third sibling. A total of 187 of the 192 non-affected parents were available for screening. Twenty-two parents had a previously diagnosed CD, one family had no parents available and three families had only one parent available.
The median age at diagnosis in the 224 affected siblings (76 males and 148 females)
Methods

Serologic markers
IgA-EMA was the antibody of choice used as a screening method for CD in both the multiplex and simplex materials [89]. IgA-EMA was used in clinical practice at the time of the study [172]. Sera with IgA-EMA antibodies detected at a dilution of 1:10 or more were considered positive. All IgA-EMA positive siblings and parents

Simplex families (Paper IV)
For confirmation studies a simplex material consisting of 135 families with one affected child was collected from south-west Sweden. An invitation was send to CD families of the registries at our own paediatric department in the Gothenburg region. The same strict inclusion and exclusion criteria were applied as in the multiplex material. Medical records of individuals with diagnosed CD were collected and reviewed. Diagnostic data and clinical manifestations were recorded. Blood samples for DNA extraction were collected from all CD patients and the healthy parents available as well as samples for serologic screening.

At the start of the study there were two CD parents (both females, one with DH) although seven new CD parents were found through screening (three males and four females).

The mean age at diagnosis in the 135 Swedish affected CD (44 males and 90 females) was 1.9 years and the median age 1.4 years (range 7 months – 12 years).

DQ2-negative cases and DQ2-positive controls (Paper IV)
The patients in this study were among those collected by the different European partners in the EU financed research consortium on CD, The European Cluster on Coeliac Disease (QLKT – 1999-00037) [64]. The cases were DQ2-negative from France, Italy, Finland and Sweden and the DQ2-positive controls from the same materials were matched only by country. A total of 85 DQ2-negative cases (38 DQ8-positive and 47 DQ2-negative/DQ8-negative) and 102 DQ2-positive controls were collected. This material is described in detail in Paper IV. The same diagnostic criteria as described previously in multiplex and simplex materials were used for the Swedish patients and at least the ESPGHAN 90 [88] for the patients from Finland, France and Italy.

A questionnaire was used to collect clinical data retrospectively from the medical files. Distribution of age at diagnosis and gender are presented in Table 2, Paper IV.

Ethics
The ethics committee of respective centre approved the study.
without previously diagnosed CD underwent small intestinal biopsy. IgA deficiency was excluded in negative individuals using routine laboratory methods.

**Small intestinal biopsy**
The biopsies were done according to the routine of the home clinic. The biopsy routines at clinics in Sweden and Norway are quite similar. Children were mainly investigated using capsule biopsy [173] and adults sing endoscopic biopsies. The histology reports were evaluated and classified by me according to a modified Alexander scale (Table 2) [99, 174]. When the written reports were difficult to assess or evaluate the histological slides were re-evaluated by a pathologist specialised in paediatric gastro-enterology (Walter Ryd, Department of Pathology, Sahlgrenska University Hospital, Göteborg, Sweden). Alexander grade 3 or more was considered compatible with CD, used for the multiplex and simplex family materials.

In Paper IV MARSH 3 (Table 3) [100, 101] was considered indicative of CD.

**Statistics**
*Linkage analysis uses a non-parametric approach.* A study of allele-sharing between relatives. Relatives concordant for disease should have increased allele-sharing close to a susceptibility locus, while relatives who are discordant should have decreased allele sharing. No special model of inheritance is assumed and it is therefore a suitable method for complex diseases such as CD. One common non-parametric approach is the affected sib-pair method, which counts the alleles that a sib-pair has inherited in common, IBD at a marker locus. According to the Mendel’s first law of segregation, the siblings would have, at any non-linked locus, a 25% probability of inheriting 0 alleles IBD, a 50% probability of inheriting one allele IBD (1 maternal or paternal shared allele) and a 25% probability of inheriting two alleles IBD (both the maternal and paternal alleles shared).

In the calculation of the risk of CD in the members of nuclear families with two affected siblings, we denoted $K_{2S}$ as the risk that an additional sib to an affected sib-pair is affected and $K_{2O}$ as the risk of a parent of an affected offspring-pair being affected (page 340, Paper I).

**Bootstrapping** is a computer method [175] that we used for estimating confidence intervals for the risk in Paper I. New samples of the same size are drawn with replacement from the original sample. The idea behind bootstrap is to produce a variety of values whose variability reflects what would be obtained if samples were repeatedly taken from the whole population.

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<th>Median</th>
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<td>Original sample: 11, 22, 33, 44, 55, 66, 77, 88, 99</td>
<td>55</td>
</tr>
<tr>
<td>Bootstrap 1: 22, 33, 55, 55, 33, 99, 88, 44, 11</td>
<td>44</td>
</tr>
<tr>
<td>Bootstrap 2: 44, 55, 22, 11, 11, 66, 99, 88, 88</td>
<td>55</td>
</tr>
</tbody>
</table>
This is repeated 1000 times. The medians are then ordered by increasing value. The 25th and 975th values out of 1,000 give the lower and upper estimates of the 95% confidence interval.

The GENEHUNTER computer program [176] was used for linkage and linkage disequilibrium analyses of the results of the fragment analysis of the microsatellites in Paper II. **Allegro** is another computer program for multipoint linkage analysis, which can analyse pedigrees of larger size [177]. To analyse possible associations between IBD sharing at CTLA4 and 5q31-33 on the one hand and the phenotype on the other, we used the non-parametric linkage (NPL) statistics. Families were divided into two groups according to symptom gradation, one containing symptom-concordant sib-pairs and one containing symptom-discordant sib-pairs (Table II, Paper III). For each group, NPL was calculated using **Allegro** 1.2, thus obtaining two statistics, NPL\textsubscript{conc} and NPL\textsubscript{disc}.

**Permutation analysis** with 10,000 iterations was further used to assess the significance of the difference between NPL\textsubscript{conc} and NPL\textsubscript{disc}. Permutation analysis shows the range of statistics possible under the null hypothesis – that there is no association between marker genotype and phenotype. The idea is to keep the original statistic T\textsubscript{obs}, e.g. a correlation between two variables, and then by repeatedly shuffling either variable destroy all but random associations between them and calculate a statistic T\textsubscript{i} for each permutation i. Finally, the statistic T\textsubscript{obs} is compared to all T\textsubscript{i} (i = 1 to n) n typically large (often 10,000 or more). All the T\textsubscript{i} form an empirical distribution from which an empirical p-value can be estimated.

The heritability of the symptom grade was established using the software SOLAR (Sequential Oligogenic Linkage Analysis Routines) a software that has the general pedigree variance component and IBD estimation methods [178]. The one-sided TDT was used to analyse transmission of A and G alleles in the high and low HLA risk groups in Paper II.

**Spearman’s rank** correlation coefficient does not require the assumption that the relationship between the variables is linear with normally distributed residual, nor does it require the variables to be measured on interval scales; it can be used for variables measured at the ordinal level as in our case age at diagnosis.

**Fisher’s exact test** or Chi-square test was used for comparing two groups.

**Mantel-Haenszel test** was used for comparing data from several 2x2 tables.

**Statistical stratification**

Only one sibling from each sib-pair family was used (Papers III and IV) to avoid bias, as the siblings’ phenotype can be familiar for genetic as well as non-genetic reasons.

The phenotype distribution as well as the genotype distribution differs between countries in Paper IV. Therefore, the groups were compared first with Fisher’s exact test or chi-square tests within each country and secondly, if the odds ratios were found to be homogenous between countries, a Mantel-Haenszel statistics was used.
Methods

Genetic analysis

Analysis strategy

A two-step strategy was used for the analysis. First we selected 70 families, which we considered to be the most informative, i.e. those with a large number of children and affected individuals (Group A). The remaining 36 families (Group B) were added to group A in a second step and were genotyped over selected regions based on one of three different criteria. First criterion: chromosomes implicated by the previously published CD genome-wide screens: chromosomes 5, 11 and 15 [179-181]. Second criterion: novel chromosomes identified from Group A, which showed NPL-values of above 2.0. Third criterion: chromosome 2 and 20 because of the location of two candidate genes, suggested as being involved in CD: CTLA4 +49 and TG2. (Figure 1, Paper II). In total, 137 markers across nine chromosomes: 2, 5, 6, 9, 11, 14, 17, 20 and X were analysed for all 106 nuclear families (Groups A and B). The remaining 261 markers on the other chromosomes were only analysed for the initial 70 families (Group A).

Microsatellite genotyping

In Paper II, 106 sib-pair were analysed with total of 398 microsatellites. Weber screening set version six from Research Genetics, containing 390 microsatellite markers with an average distance between the markers of 10 cM (ftp://ftp.resgen.com/pub/mappairs/humanset), was used. Additional markers were used in two regions, typed over a 2 - 5 cM region: in the HLA class II region on chromosome 6 and in the CTLA4/CD28 region on chromosome 2q [72, 182]. We amplified the microsatellite marker regions using an ABI 877PCR robot under standard conditions. PCR products were separated by electrophoresis on an ABI 377XL sequencer and a 5% denaturing polyacrylamide gel, Figure 6. Genotyping was performed using GENESCAN ANALYSIS 2.1 and GENOTYPER 2.0 software (http://www.appliedbiosystems.com).

Figure 6. Two microsatellite markers fluorescently labelled.
The linkage analysis was done with the non-parametric statistic NPLall using the software GENEHUNTER version 2.0 for autosomes and version 1.3 for the X chromosome. NPLall was used in multipoint mode and uses the disease status from all family members and takes into account, for example, if one parent is affected and produces higher scores if an affected parent transmits a chromosome IBD.

In our studies the following genotypes have been found to have an association and/or linkage to CD and were used in Paper III: association to CTLA4 +49 A/G by TDT and linkage using NPL analysis [72]. Analysed SNPs on MH30, -1147, CT60 and CT61 observed strong LD. A haplotype of this region marked by the alleles -1147*T: + 49*A:CT60*G:CT61*A was associated significantly with CD [183]. IBDs from linkage on 5q31-33 [184]. The HLA genotyping was done previously [68].

For the case-control association study in Paper IV, the previously performed HLA genotyping was used [64].

**Phenotypes**

I did the clinical classification, retrospectively from the medical records (Papers III and IV) or from a questionnaire by the person responsible for each country (Paper IV). The patients were classified into three symptom grades: Grade 1: Patients with “classic” celiac symptoms such as diarrhoea, vomiting, abdominal distension malabsorption and growth failure. Grade 2: Patients with milder presentation of symptoms typical of CD. Grade 3: Originally we had few patients with atypical symptoms (three patients in Paper III and five patients in Paper IV) but for statistical analysis purposes they were grouped as Grade 2. Patients with clinically silent CD are defined as asymptomatic individuals with positive antibodies and gluten enteropathy.

Age at diagnosis, age at onset of symptoms, gender, being a proband, presence of gastrointestinal symptoms and presence of autoimmune disorder or DH were recorded.

**Genotype-phenotype association (Paper III-IV)**

In Paper III the genotyping was classified into four groups: 1. HLA-risk groups: high when a patient had either of the two genotypes, DR3-DQ2/DR3-DQ2 or DR3-DQ2/DR7-DQ2, intermittent when DR3-DQ2/X (X is any other genotype) or DR5-DQ7/DR7-DQ2 and low when the patient had any other HLA genotype. 2. The CTLA4 +49 A/G allele (AA, AG, GG). 3. The haplotype MH30*G:-1147*T:+49*A:CT60*G:CT61*A. 4. The 5q31-33 region, IBD status at locus D5S436.

In Paper IV, the HLA-DQ2-negative cases were sub-grouped as DQ8-positive and DQ2-negative/DQ8-negative patients (Figure 1, Paper IV) and compared to DQ2-positive controls.
Results

Paper I
Fourteen apparently healthy relatives were IgA-EMA positive, six siblings and eight parents. All underwent a small intestinal biopsy, which confirmed the CD diagnosis in all except one parent (Table 4, Paper I). The risk for the remaining siblings and parents in sib-pair families was increased, as expected according to Falconer’s theory [159, 185]. We could confirm a threefold increased risk compared to families with only one child with CD. The sibling risk was 26.3% (95% CI, 13.8-38.7) and the parent risk 12.9% (95% CI, 9.0-17.1).

Table 5. Estimated risk for CD in population and in first-degree relatives in single and sib-pair families.

<table>
<thead>
<tr>
<th></th>
<th>Population risk</th>
<th>Siblings</th>
<th>Parents</th>
</tr>
</thead>
<tbody>
<tr>
<td>Populations studies</td>
<td>0.2 – 1.0 %</td>
<td>2.6 – 12.2 %</td>
<td>0 – 6.3 %</td>
</tr>
<tr>
<td>Single CD case families</td>
<td>26.3 %</td>
<td>12.9 %</td>
<td></td>
</tr>
<tr>
<td>Sib-pair CD families</td>
<td>26.3 %</td>
<td>12.9 %</td>
<td></td>
</tr>
</tbody>
</table>

An unexpected male preponderance was found among the new CD cases (10 males: 3 females) compared to all CD cases (86 males: 160 females) at the start of the study.

Paper II
The genome-wide scan of 106 celiac families and 398 microsatellite markers. In total, chromosomes 2, 5, 6, 9, 11, 14, 15, 20 and the X chromosomes were analyzed in all families and the remaining chromosomes were analyzed in 70 families. Apart from HLA on chromosome 6 (NPL 4.40) our best results were found in eight chromosome regions, 2q11-13, 3p24, 5q31-33, 9p21, 11p15, 11q23-25, 17q22 and Xp11 (Table 2, Paper II). Taking into account that chromosomes 5 and 11 had been reported previously, our results strengthened the notion that these regions could harbour susceptibility genes for CD.

Paper III
The heritability of the phenotype was statistically significant and was estimated at 0.45 (p = 0.02). Association analysis (Table III, Paper III) showed a statistically significant dependence between CTLA4 +49 A/G genotypes and the different symptom grades (p = 0.014), with more AA genotypes than expected in the clinically silent group.
No significant association between symptom grade and the other genotypes was found.
The stratified linkage analysis showed a difference in allele-sharing at CTLA4 +49A/G among symptom-concordant sib-ship families (NPLconc = 2.0) compared to the
Results

Symptom-discordant sib-ship families (NPLdisc = -0.7) (p = 0.04). The 5q31-33 locus, however, had no significant difference (p = 0.13) in allele-sharing between the two groups. No significant correlation was found between age at diagnosis in the probands and the genotypes or between sex and the genotypes. Nor was any significant association between sex and symptom grade found.

Paper IV

First we compared the DQ2-negative cases and DQ2-positive controls (Table 2, Paper IV). The clinical grades were significantly different between the cases and the controls in Italy (p = 0.006) and Sweden (p = 0.014). In both countries there was an association between classic presentation (Grade 1) and DQ2-negative cases, and in the Italian group there was an additional association between DQ2-negative cases and silent presentation. No significant differences between cases and controls were found for age at diagnosis, age of onset of symptoms, gender or gastrointestinal symptoms. The DQ2-negative cases in the Italian group were diagnosed significantly more often by screening (p = 0.006). In the Finnish cases a milder intestinal histology was found (p = 0.001). There were no significant differences between cases and controls for DH or other autoimmune diseases. When conditions for analysing all countries as one group were achieved, a Mantel-Haenszel test (HM) was performed. There were no significant differences between cases and controls for these other phenotypes.

In the next step the cases in the DQ8-positive and DQ2-negative/DQ8-negative subgroups were compared in the same way (Table 4, Paper IV). Autoimmune diseases were significantly (p = 0.007) overrepresented in the DQ8-positive subgroup compared to DQ2-negative/DQ8-negative. There were no significant differences between DQ8-positive and DQ2/DQ8-negative subgroups for the other phenotypes, including clinical symptom grades.

In the DQ2-negative/DQ8-negative subgroup all patients except three encoded half of the DQ2 heterodimer, either DQA1*05 or DQB1*02. These three individuals were from the Italian group and satisfied the ESPGHAN diagnostic criteria for CD [88]. All of them were diagnosed in childhood. One patient was six years old at diagnosis. He was diagnosed in a family screening and had a clinically silent CD. The other two were both two years of age at diagnosis and had classic symptoms such as diarrhea, failure to thrive and vomiting. One of them had a sibling with CD. These cases are previously described in detail [64].

Autoimmune disease was diagnosed in six DQ2-negative patients. All of them were DQ8-positive. Three had type 1 diabetes mellitus (T1D), one thyroid disease, one rheumatic disease and one ulcerative colitis. Thus six out of the 38 (16%) DQ8-positive cases had another autoimmune disease compared to none of the 47 DQ8-negative cases (p = 0.026). Furthermore, five of the six patients were homozygous for DQ8; the patient with ulcerative colitis was the only heterozygous. In the DQ2-positive CD controls only two (one T1D, one rheumatic disease) out of 102 (2%) had another autoimmune disease, which was significantly less compared to the DQ8-positive cases (p = 0.008).
Conclusion and discussion

This thesis was a co-operative endeavour between physicians, geneticists and statisticians to discover the non-HLA genes in CD as well as to increase the knowledge of HLA genetics. Furthermore, performing genotype-phenotype analysis is an attempt to understand the complexity of the phenotypes. This work required collaboration with other centres in Sweden and other European countries.

Some families have stronger genetic susceptibility for CD as in families with two or more affected siblings. These affected sib-pair families are important for genetic studies to find common susceptibility genes.

The strategy for collecting the best material for genetic analysis and genotype-phenotype analysis in CD is as follows: Large sib-pair material, reliable diagnosis in the cases, age at diagnosis, gender and clinical symptoms grades.

The advantage of the Swedish/Norwegian material is that the risk of patients with a false diagnosis is minimal as review of all investigation was done with as much thought as possible. The other patients are part of genetic study materials and are diagnosed according to the ESPGHAN 90 criteria. All healthy relatives were set as unknown because, as in a disease such as CD, one screening cannot eliminate the possibility of being affected later.

Almost all parents’ DNA was available and so the risk of genotype error was minimal as we could check the allele or haplotype. The patient materials are family-based, which can be powerful when combining both genetic linkage and association, and they are also regional population-based.

All the IgA-EMA positive individuals underwent a small intestinal biopsy to confirm the CD diagnosis and minimise the risk of false-positive cases. Latent or potential CD is also an unknown factor here. We are aware of siblings that had their CD diagnosis a few years after the study came to an end and our results are thus probably slightly underestimated. Considering the high level of knowledge of CD in these families, the number of undiagnosed cases is surprisingly high and many had symptomatic CD. We suggested that serological screening should be offered to all first-degree relatives of CD patients. Clinical practice is often otherwise. The time for screening children is always a debate as screening only once in younger children is not enough. The natural course of undetected CD is still a matter for debate – whether the risk of other autoimmune diseases, anaemia, osteoporosis and malignancy is increased to the same extent for patients who are clinically silent as for symptomatic patients. Should those that have positive antibodies without gluten enteropathy receive treatment with GFD or only be offered follow-up biopsy. One parent in our study with positive IgA-EMA and increased IEL cell infiltration in the intestinal biopsy was symptomatic and commenced GFD with good clinical response.

The increased risk for remaining siblings and parents shown in our study is in accordance with another study, which showed a sibling risk of 21.3% (95% CI, 11.0-31.6) where they even calculated an offspring risk of 14.7% (95% CI, 6.2-23.2), second-degree relatives 19.5% (95% CI, 15.1-23.9) and first cousins 17% (95% CI, 6.8-27.7) [186].
For genetic analysis, it could be a good strategy to first analyze genetic information in region- or ethnic-based study material and then make confirmation analysis in other materials. This material, one of the largest sib-pair materials of CD published, is relatively homogeneous with regard to regional ethnicity with mostly Swedish Caucasian families and some south Norwegian Caucasian families. In a complex disease such as CD, all susceptibility genes are not likely to cause disease in all families. Many independent sib-pair families are preferable in GWL studies of complex disease as there is a good chance of finding those common susceptibility genes that cause disease in the majority of cases. The different phenotypes can include age at diagnosis, gender, clinical manifestation, associated diseases and HLA-DQ type. In this material there are patients with differences with regard to age at diagnosis as well as different clinical phenotypes and we also had access to the parents’ genetic information. The two regions on chromosome 5q31-33 and 11q23-25 that our GWL study indicated as true susceptibility regions have been reported in other sib-pair GWL studies, which makes these regions interesting for the further study of susceptibility genes in CD. The GWL study was also included in a meta analysis together with three other European partners (Finland, Italy and UK), where chromosome 5q31-33 was the only significant locus, apart from HLA [77]. Further studies in our research group have been published and some are ongoing on 5q31-3 [78, 184]. The genotype-phenotype study included the susceptible gene and regions that had shown association and or linkage in our material. Paper III is the first study to publish a genotype-phenotype analysis for CTLA4 +49 A/G in CD. Our findings indicate that this gene may influence the phenotype of CD. Surprisingly, we found that the AA genotype was correlated to the silent phenotype. As the genetic association was to the A allele [72] one might think that the AA genotype would be associated more to symptomatic disease. The role of CTLA4 +49 in CD is not known. A few studies have conducted a genotype-phenotype analysis for CTLA4 +49 A/G in other autoimmune diseases. In MS, a genotype-phenotype study showed that AA and AG genotypes are associated with MS progression [187] but in rheumatoid arthritis no CTLA4 +49 association to the phenotypes was found [188]. No association was found with the different HLA risk groups, the MH30*G::1147*T::+49*A:CT60*G:CT61*A haplotype or the 5q31-33 region.

Knowledge of genotype-phenotype can help to stratify the material by sub-phenotype, which selects for those specific individuals who carry a rare gene variant and increase the chance of detecting linkage or association due to rare variants within a given sample set.

HLA is a necessary and single most important genetic factor in CD. The HLA-DQ2 is responsive in 90% of the patients and together with DQ8 predisposes to disease development with a known mechanism (Figure 2). The DQ8-positive and the DQ2-negative/DQ8-negative patients are 5% each of the CD population. We had the opportunity to compare the clinical presentation in DQ2-negative cases with DQ2-positive controls in a large DQ2-negative sample. Silent presentation was correlated to DQ2-negative status but only in the Italian group. The other countries had too few patients with silent presentation to detect such a pattern. However, analysing only symptomatic patients, DQ2-negativity was associated with classic symptoms or
Conclusions

Grade 1, seen both in the Italian and the Swedish groups. Our results differ from a study of 25 DQ2-negative cases in Finland and Spain, where no clinical differences compared to the DQ2-positive controls were found [12].

In clinical practice, HLA, the use of serology tests that test for DQ2 and DQ8-heterodimers, has become more common. However, they need to be used with caution as our results showed that CD could not be excluded by only testing whether or not patients are DQ2 and DQ8-positive. There were more silent patients in the DQ2-negative cases with an even division between DQ8-positive and DQ2-negative/DQ8-negative CD patients. However, most of the DQ2-negative/DQ8-negative patients have one half of the DQ2 heterodimer, except for the three patients that had the non-part of the heterodimer but had classic CD symptoms.

Our effort was to collect two controls for every DQ2-negative and country, which was not accomplished. There were less DQ2-positive controls from Italy than we had planned. We therefore needed to analyse each country separately in case of genetic or clinical differences between the countries.

Other autoimmune diseases were looked at in Papers III and IV and despite the fact that the samples have small amounts there were some interesting findings. T1D was the disease that occurred most frequently. Interestingly, there were no patients with autoimmune disease carrying homozygous for CTLA4 +49 A-allele or who shared 2 haplotype IBD with their co-sibling in the 5q31-33 region. Autoimmune diseases were significantly overrepresented in the DQ8-positive subgroup compared to the DQ2-negative/DQ8-negative subgroup. Genetic analyses are ongoing in most autoimmune diseases associated with CD and many are looking at genes in the same region. It is most likely that there are some common genes in these diseases, probably with some genes with immunological function.

The significance of interaction or epistasis of different genes is not possible to determine from this thesis as no gene-to-gene interaction analysis has been done. Theoretically, there are probably gene interactions and the genotypes could have a different significance in different patients e.g. HLA high-risk patients may not need any or few other candidate genes whereas HLA low-risk patients need other genes or several genes to get CD. The genetic effects may be understood before we understand how different environments may interact with specific genetic components that lead to the disease.
Future genetic and genotype-phenotype studies in CD

This thesis has increased the knowledge of the genetic and the genotype-phenotype association in CD. The results support the theory that there could be different genetic susceptibility in different phenotypes. The main findings were the association shown between CTLA4 +49 A/G genotypes and the different symptom grades with more AA genotypes than were expected in the clinically silent group. DQ2-negative cases were significantly different compared to DQ2-positive controls with regard to clinical grades. Autoimmune diseases were significantly overrepresented in the DQ8-positive subgroup compared to DQ2-negative/DQ8-negative.

The research into the genetics of common or complex diseases is now in an exciting phase. There are new technologies in genetic research, such as GWA studies that have been listed as “the breakthrough of the year” 2007 in Science [160]. More than 50 new disease-susceptibility genes have been found in different common diseases in just one year. This will hopefully offer a completely new view of CD and other immune and autoimmune diseases. Discover new potential candidate genes that could increase the potential for genotype-phenotype analysis. Increase our understanding of the role that genes play in disease pathogenesis and design molecular mechanism-based therapies that will improve clinical outcome or quality of life.

One GWA study on CD has been published [69]. The HLA locus came out with very highly significant P values as expected. The second strongest association was found for a few SNPs located close to the IL2 and IL21 genes on chromosome 4q27. A follow-up study by the same group of >1,000 markers showing association signals in the initial celiac GWA study has identified an additional seven loci associated with CD. The regions are 1q31, 2q11-12, 3p21, 3q25-26, 3q28, 4q27, 6q25 and 12q24. Immune-related genes map to six of these regions, underscoring the possibility that CD has an immunological basis [189].

Fine-mapping the new regions will be required to find potential candidate genes and provides better conditions for further functional and clinical studies. Moreover, there are probably modifying effects between the genetic (epistasis) and environmental factors and interactions between the genetic and environmental factors and epigenetic factors. It may also be that in different patients with identical phenotypes, only some genetic/environmental factors are shared and only some of the pathways leading to disease development are identical. This makes genotype-phenotype analysis difficult.

When gene-gene (epistasis) analysis with true candidate genes and optimal patient material, with all the phenotypes and perhaps larger subgroups as clinical symptom grades, HLA-subgroups, CD patients with other autoimmune diseases etc, we might succeed in our ambition to walk along the research path (Figure 1) towards disease susceptibility proteins that will increase our understanding of CD and hopefully other autoimmune diseases. By associating phenotypes to specific profiles of gene/protein expression pathological mechanisms can be unveiled, predictive instruments
in future diagnosis can be found and candidates for specific therapies can be selected. In summary, this thesis shows that the risk for third siblings and parents is as expected increased in sib-pair families, as the expected risk of being affected in polygenic diseases is higher in multiple-case families compared to single-case families. The genome scan indicated significant linkage to 11q and 5q, which makes these regions interesting for further fine-mapping of these regions using association analysis. Genotype-phenotype analysis of both HLA and non-HLA locus showed some significant correlation between silent CD and both CTLA4 +49 AA genotype and the DQ2-negatives. In addition, an association between classical symptoms and DQ2-negative cases has been shown, when only the symptomatic patients were analysed.
Sammanfattning på svenska

Sjukdomen celiaki (CD) eller glutenintolerans är en kronisk inflammatorisk sjukdom av autoimmun typ och är en av de vanligaste sjukdomarna i barnåren, men diagnostiseras i alla åldrar. Förekomsten av CD i Sverige är runt 1 % men kan vara betydligt högre. Risken för ett syskon (syskonrisken) att bli sjuk är 10 % om man har ett sjukt syskon. CD är mångfaktoriell sjukdom med samspel av troligtvis flera gener och omgivningsfaktorer. CD är unik sjukdom genom att man känner den utlösende faktorn, äggviteämnet i gluten, som leder till inflammation och skada på tunntarmens ludd. För att kunna ställa rätt diagnos krävs tunntarmsbiopsi, men screening med serologisk antikroppstest görs ofta först. Behandling med gluten fri kost utan vete, råg och korn, leder till utläckning. Diagnos kriterier för CD, eller ESPGHAN criteria, grundar sig på att tunntarmslemhinnans morfologi i biopsi preparat karaktäriseras av villusatrofi, kryptyperplasi och ökat antal av intraepitheliala lymphocyter (IEL). Sjukdomsbilder eller fenotypen varierar från svårt sjuka barn till barn och vuxna med mindre symptom och till patienter utan symptom eller symptomfri (silent) CD.

Genetiken har stor betydelse vid sjukdomen. HLA är en typ av antigenpresenterande molekyler som existerar på cellytan hos alla celler i människokroppen. Dess funktion är att presentera utvalda peptider för immunförsvarets celler. HLA klass II på kromosom 6 är väl känt vid CD. HLA-DQ2 är den allra vanligaste molekylen men en mindre del har HLA-DQ8. Andra HLA typer kan förkomsa och kallas här DQ2-negativa/DQ8-negativa. HLA genen kan bara förklara runt 40 % av sjukdomens genetik så resten måste förklaras av non-HLA gener. Dessa kandidatgener är ännu inte kända men eventuella kandidatgen områden har identifierats. Genotyp-fenotyp association vid CD är begränsad.

Målsättningen med detta arbete var att beräkna syskonrisken hos det tredje syskonet och föräldrar till två sjuka syskon, genomföra systematisk screening för att kartlägga misstänkta sjukdomsregioner och undersöka genotyp-fenotyp association vid CD. Material insamlades från Sverige och Norge med 107 familjer med minst två sjuka barn, totalt 224 CD syskon, samt deras friska syskon och de flesta av deras föräldrar. Screening för CD genomfördes hos friska syskon och deras föräldrar. Trettio nya fall av CD diagnostiserades, 6 syskon och 7 föräldrar. Den beräknade syskonrisken var 26.3 % och föräldrarisken var 12.9 % eller nästan tre gånger högre än om ett syskon är sjukt. Det höga antal ny diagnostiserade fall är förvånande med tanke på den höga kunskapsnivån om sjukdomen som finns i dessa familjer. Vi föreslår att man erbjuder alla första grad släktingar serologisk screening för CD.

Systematisk screening av alla kromosomer gjordes med kopplingsanalys och genetiska variabler för att identifiera kromosomregioner som nedärvs tillsammans med sjukdomen. Förutom HLA, visade den signifikant koppling till sjukdomen på kromosom 11 och 5q31-33. Område 5q31-33 har visat koppling i flera andra studier. Ytterligare material med 136 svenska familjer med ett sjukt barn samlades in för genetiska associationsanalyser.

Två genotyp-fenotyp association studier genomfördes. Den första på de kandidatgener eller genområden som hade visat association och/eller koppling i vårt mate-

Genotypen AA i CTLA4 +49A/G polymorfier visade association till den symptomfria sjukdomsbilden. Ingen association fanns mellan de andra genotyperna och symptombilden, ålder vid diagnos eller kön.


Ökad kunskap om genetiska faktorer och omgivningsfaktorer samt interaktionen mellan dessa leder till ökad kunskap om patogenesen t.ex. de immunologiska mekanismerna vid CD och även andra autoimmuna sjukdomar. Detta kan på sikt leda till ökade möjligheter till primärprevention och nya behandlingsformer.
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These studies have required interdisciplinary cooperation between clinicians, geneticists and statisticians. The teamwork has made the research more profitable and pleasant.

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