AUTOANTIBODY RECOGNITION OF COLLAGEN TYPE II IN ARTHRITIS

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ABSTRACT

Autoantibodies against collagen type II (CII), a protein localized in the joint cartilage, play a major role in collagen-induced arthritis (CIA), one of the most commonly used animal models for rheumatoid arthritis (RA). The studies included in this thesis were undertaken to elucidate structural and functional requirements for B and T cells to recognize native CII structures during experimental arthritis as well as in human RA. To reveal in detail how CII-specific autoantibodies recognize CII, we determined the crystal structure of two pathogenic immune complexes occurring in arthritis. The two crystal structures reveal how autoantibodies target triple-helical CII and specifically which epitope sequences and CDR residues that are crucial for binding. Interestingly, although amino acid residues in the hypervariable regions of both autoantibodies were somatically mutated, the majority of the contacts with the epitope involve germline-encoded structures. A recombinant CII peptide library was also generated in order to reveal the CII epitope specificity of polyclonal autoantibodies in CIA. This library was used to confirm already known CII epitopes occurring in CIA but also to investigate new CII epitope regions that could play a role in arthritis. Several new regions targeted by autoantibodies were found and interestingly, antibodies to these regions were also identified in non-human primate species with arthritis. Subsequently, autoantibodies to the major CII epitopes C1, U1, and J1 were analyzed in serum and synovial fluid from RA patients. The autoantibody levels to all three CII epitopes were significantly higher in synovial fluid compared to serum, in particular autoantibodies to the U1 epitope. Finally, we studied the CII-specific antibody responses as well as CII-specific T cell responses during different timepoints of CIA in two mouse strains, B10.Q and a humanized mouse strain B10.DR4.Ncf1+/−. The obtained data showed that the antibody and T cell responses investigated stayed relatively constant during the disease course and were dominated by T cells specific for non-modified (K264) and hydroxylated (HOK264) CII259-273 peptides in the humanized mice and to the galactosylated (GalOK264) CII259-273 peptide in the B10.Q mice. C1 specific antibodies dominated in both strains while antibodies to the U1 epitope was characteristic for the humanized mouse.
LIST OF PUBLICATIONS

I. Analysis of autoantibodies to the joint-specific protein collagen II in experimental and rheumatoid arthritis
Ingrid Lindh, Hüseyin Uysal, Omri Snir, Erik Lönnblom, Ida Andersson, Franziska Lange, Michel Vierboom, Kutty Selva Nandakumar, Vivianne Malmström, and Rikard Holmdahl
Manuscript

II. Crystal structure of an arthritogenic anticollagen immune complex
Doreen Dobritzsch*, Ingrid Lindh*, Hüseyin Uysal, Kutty Selva Nandakumar, Harald Burkhardt, Gunter Schneider, and Rikard Holmdahl

III. Comparative analysis of CII-specific immune responses during development of collagen induced arthritis in two B10 strains
Tsvetelina Batsalova*, Ingrid Lindh*, Johan Bäcklund, Balik Dzhambazov, and Rikard Holmdahl

IV. Immunoglobulin heavy chain V gene polymorphism controls epitope-specific antibody response
Bruno Raposo, Doreen Dobritzsch, Ingrid Lindh, Diana Ekman, Changrong Ge, Michael Förster, Hüseyin Uysal, Gunter Schneider, and Rikard Holmdahl
Manuscript

*These authors contributed equally to the work

Additional publication not included in the thesis:

Tissue transglutaminase enhances collagen type II-induced arthritis and modifies the immunodominant T-cell epitope CII260-270
Balik Dzhambazov, Ingrid Lindh, Åke Engström, and Rikard Holmdahl
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<tr>
<td>Aa</td>
<td>Amino acid(s)</td>
</tr>
<tr>
<td>Ab(s)</td>
<td>Antibody (antibodies)</td>
</tr>
<tr>
<td>ACPA</td>
<td>Anti-citrullinated protein antibody</td>
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<tr>
<td>BCR</td>
<td>B-cell receptor</td>
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<td>C</td>
<td>Constant</td>
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<td>CII</td>
<td>Type II collagen</td>
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<td>CB</td>
<td>Cyanogen bromide</td>
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<td>CAIA</td>
<td>Collagen antibody induced arthritis</td>
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<td>CDR</td>
<td>Complementary determining region</td>
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<td>CIA</td>
<td>Collagen induced arthritis</td>
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<td>COMP</td>
<td>Cartilage oligomeric matrix protein</td>
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<td>D</td>
<td>Diversity</td>
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<td>Fab</td>
<td>Fragment antigen binding</td>
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<td>Fragment crystallizable</td>
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<td>HLA</td>
<td>Human leukocyte antigen</td>
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<td>Interleukin</td>
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<td>MHC</td>
<td>Major histocompatibility complex</td>
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<td>pg</td>
<td>Proteoglycan</td>
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<tr>
<td>RA</td>
<td>Rheumatoid arthritis</td>
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<tr>
<td>RF</td>
<td>Rheumatoid factor</td>
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<tr>
<td>TCR</td>
<td>T cell receptor</td>
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<tr>
<td>Th</td>
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1 INTRODUCTION TO THE STUDY

Rheumatoid arthritis (RA) is an autoimmune disorder characterized by persistent synovitis, systemic inflammation, and autoantibodies. The disease etiology, symptoms, and pathogenesis are complex and multifaceted. This can be demonstrated by the effectiveness of certain biological therapies in some patients but a complete lack of response in a large number of other patients (1). Discovering subtypes of RA patients is therefore today considered a central research area for the improvement of response to therapy (2,3). Several subtypes of RA patients have been identified based on particular clinical and molecular characteristics (4,5). The studies presented in this thesis are limited to a subgroup of RA patients developing an autoimmune response to the cartilage-specific protein collagen type II (CII) but can be extended to immune responses against other autoantigens as well.

1.1 BACKGROUND

RA is a complex disorder that if left untreated, severely affects cartilage and bone. CII-specific cells from the immune system (B and T cells) as well as autoantibodies and cytokines they secrete can be detected in these local areas as well as in serum (6–10) and several lines of evidence confirm their contribution to the abnormal pathogenic events operating in and around the RA joints (6,11–16). The basic mechanisms occurring in such complex disorders can be studied in controlled animal model systems that provide insights into one or another particular aspect or phase of the “entire” disease. One of the models frequently used is collagen-induced arthritis (CIA). This model share immunological and pathological features with RA (17–20) but there are also differences between the two as CIA is a time-dependent model induced with CII. The latter makes it a suitable model for studies aiming at understanding CII-specific autoimmunity.

CIA is characterized by high levels of circulating autoantibodies against triple-helical CII, but the specific epitope location and sequences of the antibody binding sites have yet not been fully characterized. This is mainly due to 1) lack of screening systems that cover all possible epitope regions and 2) that studies are limited to the specificity of monoclonal antibodies established very early post immunization. Anti-CII antibodies are pathogenic in mice (18), and except for their characteristic effector functions such as binding and activation of complement, and the activation of FcγR-bearing cells, some of them can also disrupt articular cartilage without any mediators of inflammation (21). The way cartilage and CII is affected depends on the autoantibody specificity (15,16,21,22), suggesting that epitope locations could play a part in these processes. This is also relevant to RA where these types of autoantibodies occur. However, it might be more difficult to identify pathogenic effects in RA compared to CIA since RA is more complex with other pathogenic events that potentially cover the effects of the CII response.

To fully dissect the CII epitope specific response occurring during both CIA and RA, earlier findings need to be complemented. In addition, although previous studies suggest that a consensus motif (R-G-hydrophobic) present in several of the known CII
epitopes, could be a recognition sequence for autoreactive B cells (23), its relevance in CIA and RA has not yet been confirmed.

Not only B cells but also T cells play a critical role in CIA (24–27). The presence of activated CD4⁺ T cells as well as the correlation between RA and the expression of certain MHC class II alleles (in particular certain subtypes of HLA-DR4) (28,29) has raised the possibility that RA is caused by a T cell mediated immune response towards one or a few joint-specific antigens. Both human DR4 and the CIA susceptible mouse expressing H-2q molecules can present CII to autoreactive T cells (30). The immunodominant T cell epitope on CII (CII259-273) can be post-translationally modified on the lysine (K) residues leading to non-modified, hydroxylated or glycosylated epitopes. In general, the T cell response to the glycosylated T cell epitope dominates in RA and in H-2q restricted CIA (6,31) whereas the non-modified epitope dominates in the DR4 transgenic mouse (32). Although specific B and T cell responses can be measured at a certain disease timepoint those might not give the complete picture of how CII specific B and T cell responses occur over time nor their relevance to the disease. Thus, studies focusing on CII specific responses over a longer period of time could reveal how B and T cell responses relate to different disease phases and to each other.

1.2 PURPOSE OF THE STUDY

This thesis was undertaken in order to further dissect autoimmunity to CII in CIA and RA. The main purposes as well as relevant research questions are outlined below.

The main purposes of the study were to:

I. Validate and build a more comprehensive understanding of earlier discoveries of biologically significant CII epitopes in CIA

II. Investigate the anti-CII antibody specificities in different species and mouse strains with arthritis in order to reveal that this model is relevant and matches the events operating in the human disease.

III. Perform long-time studies of the CII epitope specific response in order to reveal if epitopes could be categorized into different phases.

IV. Determine how pathogenic autoantibodies recognize triple-helical CII in order to elucidate how specificities and diversities operate in such immune complexes.

V. Investigate the magnitude and specificity of the CII specific B and T cell responses over time in different mouse strains and investigate the dissimilarities between sick and healthy mice.

Relevant research questions:

1) Could a more comprehensive CII epitope screening library be generated to complement earlier findings?

2) Could such a system be useful to screen for all potential CII epitopes detected during different disease phases and help us to understand whether some CII regions are more exposed and susceptible for antibodies to bind?
3) Could such a system also be used to investigate CII epitope specificities in different species?

4) Could we get a more structured overview of recognition motifs on CII and reveal the existence of early and late phase epitopes?

5) Can the CII specific antibody responses vary in different compartments in the body? If so, is that relevant to the disease pathogenesis?

6) Could we crystallize anti-CII antibodies together with triple helical CII peptides?

7) Could such crystal structures clearly describe the crucial immune complex interactions?

8) Do we see the same magnitude and specificity to the native and modified variants of the T cell epitope over time in CIA compared to RA? Will the response differ between mouse strains?
2 INTRODUCTION TO THE IMMUNE SYSTEM

2.1 INNATE AND ADAPTIVE IMMUNITY

If your body is affected by an infection (could be virus, bacteria, fungi, protozoa, etc.) the components of the innate immune system will recognize structures on the invading pathogens so-called pathogen associated molecular patterns (PAMPs). This system also recognize endogenous (self) molecules created upon tissue injury called damage-associated molecular patterns (DAMPs) (33–35). These structures are recognized by pattern recognition receptors (PRRs) of which the family of Toll-like receptors (TLRs) has been studied most extensively (36,37). The intracellular signaling cascades triggered by these PRRs lead to expression of inflammatory mediators that coordinate the elimination of pathogens and infected cells. In some cases processes that control the inflammatory response work inefficiently, which can lead to immunodeficiencies and induction of autoimmunity (i.e. an immune response against the body’s own cells and/or tissues). The primary cells of the innate immune system are concentrated at epithelial surfaces including the skin, respiratory tract, and gastrointestinal tract. Dendritic cells (DCs) are most important because of their ability to take up and process foreign material, then present antigens for a primary immune response (38–41). Further cells in the innate immune system are macrophages, neutrophils, natural killer (NK) cells, basophils, mast cells and eosinophils. Also γδ-T cells (42) and NK T cells (43) might be considered part of the innate immune system. Not only cells but also the complement system, comprising plasma and membrane proteins plays a key role in innate and adaptive response (44,45) where they have an important role to eliminate pathogens and apoptotic cells. Contribution of complement to the development of RA have been described (46), as well as to the development of CIA in mice and other similar rodent models (47–56).

The adaptive immunity provides additional protection due to its capability to respond more specifically and diverse. The molecules responsible for this specific and diverse recognition of antigens are antibodies expressed and secreted by B cells, and T cell antigen receptors (TCRs) expressed on T cells. The process describing how antibody specificity and diversity operate will be discussed further in the sections below as well as in study II and IV. Adaptive immunity can be divided into humoral immunity and cell-mediated immunity. Secreted antibodies are the effector molecules of humoral immunity, capable of neutralizing and eliminating extracellular pathogens by activating various effector mechanisms. The cell-mediated immunity becomes active when pathogens survive inside cells and T cells mediate this type of immunity.

2.2 B AND T CELL DEVELOPMENT

B cells (bursa or bone marrow-derived) express B cell antigen receptors (BCRs), which recognize a wide variety of macromolecules and small chemicals. Their origin can be traced back to the evolution of adaptive immunity in jawed vertebrates more than 500 million years ago (57). T cells (thymus-derived) are a population of cells expressing T cell receptors (TCRs) recognizing small peptides loaded onto major histocompatibility complexes (MHCs) molecules.
B and T cells originate from multipotent hematopoietic stem cells in the bone marrow and go through a well-controlled development and educational process before they become fully mature cells ready to be activated and perform their effector functions.

After exiting the bone marrow progenitor T cells enter the thymus as TCR-CD4-CD8- cells. In the thymic cortex, these T cells will rearrange their TCR genes and differentiate into double positive CD4+CD8+ cells (58). It is not completely understood how the division into single positive CD4+ and CD8+ T cells occurs. However it is believed that if double positive cells is divided between two options; either it dies due to neglect if thymic epithelial cells (TEC) do not interact with it (59), or it interacts with MHC I or MHC II on a TEC initiating its development into a CD4+ or CD8+ cell (60). The minor portion of T cells that survive thymic selection exit the thymus as single positive CD4+ or CD8+ T cells expressing cell surface receptors that facilitate their entry into secondary lymphoid organs in the periphery (61) where they recognize antigens. Naïve CD4+ T cells or T helper (T_H) cells exert their effector functions mainly through secreting cytokines and chemokines that activate and/or recruit target cells. T_H cells have four (and possible more) distinct fates that are determined by the pattern of signals they receive during their initial interaction with antigen. These four populations are Th1, Th2, Th17 and induced regulatory T cells (iTreg) (62–64).

The immature B cell expresses IgM and IgD antibodies on its surface as it migrate from the bone marrow and enters the secondary lymphoid organ, the lymph node. In this organ B cells are activated by T helper (T_H) cells, follicular dendritic cells (FDC), or antigens. Following activation it will start to proliferate. The activation induced (cytidine) deaminase (AID) will introduce point mutations on the variable region of the DNA to increase their collective affinity for a particular antigen and any of its close variants (also called somatic hypermutation). AID is also involved in the class-switch recombination in which B cells change their expression from IgM to IgG or other isotypes. The B cell will then differentiate into a plasma cell or a memory B cell with a different type of antibody class.

2.2.1 Antibody structure

From the earliest studies of antibody structure it was clear that all antibody domains, whether variable or constant, form compact globular structures with a characteristic fold, termed the immunoglobulin fold (65–68). Each domain consists of a stable arrangement of hydrogen-bonded, anti-parallel β-strands that forms a bilayer structure, which is further stabilized by a disulfide bond between the two layers. Immunoglobulins (Igs) or antibodies (abs) can be divided into five major classes (isotypes): IgM, IgG, IgA, IgD, and IgE. The IgG class has been most extensively studied and is also the most abundant in the circulation of normal individuals. The IgG consists of two identical “light” chains (L) and two identical “heavy” chains (H) (Figure 1). Both the L-chain and the H-chain consist of variable and constant regions (\(V_L, C_L, V_H, \text{ and } C_H \) domains). Disulphide bonds (S-S) are covalently linking the four individual chains together. Enzymatic digestion of the molecule results in three fragments, each with a molecule weight of approximately 50kDa. Two of these fragments are identical and are called the Fab fragments and the third is called Fe.
Each Fab fragment consists of a complete L-chain and the N-terminal half of a H-chain. The Fab fragment retains the antigen binding activity of the parent molecule, although it can only behave as a monovalent antibody. The Fabs are linked to the Fc by the hinge region, which varies in length and flexibility in the different antibody classes and isotypes.

**Figure 1.** Simplified illustration of an immunoglobulin recognizing its antigen. The IgG consists of two identical “light” chains (L), colored in pink and two identical “heavy” chains (H) colored in green.

The antigen binding sites (paratopes) are located at the tips of the Fabs. Comparative studies on H-chains of the same class have shown that the sequence of the constant H-chain domains C\(_{\text{H1}}\), C\(_{\text{H2}}\) and C\(_{\text{H3}}\) remain constant whereas the V\(_{\text{H}}\) regions display variability. Just as in the L-chain, the variable region of the H-chain occurs at the N-terminal end of the molecule, is approximately 110 amino acid residues in length and contains hypervariable regions (69,70). The antigen-binding site is confined to the V\(_{\text{L}}\) and the V\(_{\text{H}}\) regions and these hypervariable regions specify the conformation of the antigen binding site of immunoglobulin molecules (antigen complementarity regions, CDRs) (69–71).

2.2.2 Antibody development

The development process starts in the bone marrow where DNA encoding the heavy and light chains of the antibodies is rearranged. Diversity (D) and joining (J) gene segments are recombined (D-J joining) followed by adding a variable (V) joining segment to the heavy chain DJ complex. Diversity of the antibodies is produced by the use of different combinations of V, D, and J gene segments in different clones of B cells. The recombination of V and J or V, D, and J, gene segments is mediated by a group of enzymes collectively called VDJ recombinase. The B and T cell-specific VDJ recombinase is composed of the recombinase-activating gene (RAG)-1 and RAG-2 proteins, and they are responsible for these recombination steps (49–51). Additional diversity is produced by sequence changes, and this generates more sequences than are
present in the germline genes. This “junctional” diversity remove or add nucleotides and as a result of these mechanisms, the nucleotide sequence at the site of V(D)J recombination in antibody or TCR molecules made by one clone of B and T cells differs from the sequence at the V(D)J site of antibody or TCR molecules made by every other clone. The junctional sequences encode the amino acids of the CDR3 loop, which are the most variable of the CDRs and the one most important for antigen recognition. Thus, junctional diversity maximizes the variability in the antigen-binding regions of antibodies and TCRs.
3 RHEUMATOID ARTHRITIS

Although the name *rheumatoid arthritis* (RA) was introduced in the 1850s (75), classification criteria were only developed 50 years ago (76,77). RA is a complex autoimmune disorder characterized by inflammatory damage to joints but also by systemic features such as inflammation of blood vessels, lungs, and other tissues. RA is best considered as a clinical syndrome covering several disease subsets (78). These subsets involve several inflammatory cascades (79), which all lead towards a final common pathway.

RA affects about 0.5-1% of the population (80,81), and the prevalence varies geographically (82,83). The disease is common in northern Europe and North America compared with parts of the developing world, such as rural west Africa (84,85). RA is more common in women than in men with a ratio of about 3:1 (86). The primarily inflammatory site in RA is the synovial membrane ([Figure 2](#image-2)) and as RA proceeds hydrops, hyperemia, cartilage destruction and bone erosion and at a later stage malalignment of the connected bones occur.

![Figure 2](image-2) Simplified illustration of a normal joint and an RA-affected joint

3.1 RELEVANT RISK FACTORS FOR DEVELOPING RA

RA is a complex disorder with disease susceptibility being linked to genetic, environmental, and immunological factors as well as to stochastic events. The most important genetic risk predisposing to RA is ascribed to the MHC class II region, especially to genes encoding HLA-DR molecules. It has been found that all HLA DRBI alleles that are associated with RA risk (28) encode a conserved sequence of 5 amino acids (positions 70-74) that surrounds the peptide-binding pocket of the antigen-presenting molecule. The presence of this so-called “shared epitope” suggests that the molecules containing it might bind the same antigen (29) or alter the peptide affinity that could have a role in promoting autoreactive responses. One of the strongest genes
linked to autoimmunity outside the HLA is the tyrosine phosphatase non-receptor, type 22 (PTPN22) (87). The first reported association of PTPN22 with RA was found in 2004 (88). Other genes that have been proposed to be associated with RA are cytotoxic T lymphocyte-associated antigen-4 (CTLA-4), peptidyl arginine deaminase, type IV (PADI4), and macrophage migration inhibitory factor (MIF) (89).

As earlier mentioned not only genes, but also environmental factors are believed to play an important role in the susceptibility to developing RA. One of the highest environmental risk factors in RA is smoking (90–92), which like other forms of bronchial stress (e.g. exposure to silica) increases the risk of developing RA among patients with susceptible HLA-DR4 alleles (93). The influence of other environmental factors such as infections, hormones, and dietary factors on RA is still unclear.

### 3.2 AUTOANTIBODIES IN RA

Autoantibodies are commonly observed in autoimmune diseases. **Rheumatoid factor (RF)**, the first autoantibody recognized in RA, was described by Waaler in 1940 (94). Identification and characterization of RF as an autoantibody that binds the Fc portion of IgG was the first direct evidence that autoimmunity might play a role in RA. Long-time studies show that production of RF and other autoantibodies can precede the onset of RA by many years (95). But although they can be detected in 60–80% of RA patients (IgM-RF) (96), there is no clear evidence that RFs are involved in the initial events triggering the disease process of RA. However, it has been shown that RF seropositive patients have more severe clinical disease and complications than seronegative patients included cardiovascular problems (97). RF is not RA-specific and occurs in various infectious diseases (98), other autoimmune diseases most notably Sjögren’s syndrome (99) (transiently) and in healthy individuals following immunization (100,101).

Citrullination of proteins is a physiological process that also occurs in inflammation and is therefore not specific for RA. However, presence of **anti-citrullinated protein antibodies (ACPAs)** is highly correlated with classical RA (102,103) whereas RF is more common but less specific since it is present in other diseases. In similarity with RF, the formation of ACPAs precedes the development of RA (95,104–106). ACPAs are found in 70–90% of RA patients and have high disease specificity and sensitivity (107,108). As with RF, they are associated with more erosive RA (78,109,110). ACPAs are locally produced in RA joints, where proteins are citrullinated during the inflammatory process (111). The major citrullinated protein in the joint was found to be fibrin (112). Additionally, various other synovial and non-synovial proteins such as collagen type II, vimentin, nuclear proteins and stress proteins have been shown to be targets of citrullination in vivo (113).

**Autoantibodies to collagen type II (CII)**, a fibrous collagen protein located in the joint cartilage can be detected in serum (7,8,14,114,115) and synovial fluid (7,9,10,116) of patients with RA. The reported incidence of these autoantibodies have varied greatly (3-71%) and this is believed to be dependent on the immunological technique used as well as types and species of native or denatured collagen (114,117–122). Support for a pathogenic role of these antibodies is obtained by that they can precede radiological changes (13,123) or the appearance of rheumatoid factor (RF) (124) and their
frequency has been correlated in RA with the presence of HLA alleles that confer susceptibility to disease (14). Furthermore, injection of human anti-CII antibodies into arthritis susceptible mice induce a mild inflammatory arthritis (12). Similarly, recent studies have also demonstrated the arthritogenicity of plasma/serum from active RA patients in FcγRIIb-deficient mice, and the IgG rich fraction has been identified as the pathogenic factor (11). Moreover, studies have shown that anti-CII positive patients can have a distinct clinical phenotype characterized by an early acute phase response that might be driven by anti-CII containing immune complexes in joint cartilage (125).

3.3 TREATMENT OF RA

Advances in understanding the pathogenesis of the disease have encouraged the development of new therapies, with improved outcomes. To prevent damage to joints, maintain functional status, and decrease pain, treatment guidelines recommend the use of disease-modifying anti-rheumatic drugs (DMARDs) immediately after the RA diagnosis has been made (126,127). One of the most common DMARDs, methotrexate, remains the first-line medication for moderate-to-severe RA (128). Biological agents have also been introduced that specifically target cytokines in the inflammatory cascade in RA. The primary target of this type of therapy is tumour necrosis factor alpha, TNF-α (129) and there are currently five approved biological (antibody or soluble receptor) for the treatment of RA, which specifically neutralize TNF-α (Etanercept, Infliximab, Adalimumab, Certolizumab, and Golimumab). Anakinra, an IL-1 receptor antagonist, and Tocilizumab, a neutralizing monoclonal antibody against human IL-6, are also approved biologicals for the treatment of RA, and in particular Tocilizumab, has been shown to be a good alternative for the treatment of nonresponders to TNF-α inhibitors (130,131). Also Rituximab, a chimeric monoclonal antibody against the CD20 antigen on all B cells except plasma cells, is used as an effective treatment of RA patients (132,133).
4 COLLAGEN INDUCED ARTHRITIS (CIA)

Collagen induced arthritis (CIA) is one of the most common animal models for RA. It can be induced in rodents and non-human primate species by immunization with CII emulsified in adjuvant, generally complete Freund’s adjuvant (CFA) (134–136).

4.1 CIA IN MICE

The CIA model share several immunological and pathological features with RA (17–20,134,136) such as synovitis, cartilage breakdown and bone deformities but there are also differences between the two as CIA is a time-dependent model induced with CII. CIA, like human RA, is MHC dependent and only occurring in animal strains with the proper class II. The susceptibility is linked to the H-2d and H-2r haplotypes (27). These haplotypes are also permissive for the induction of arthritis with homologous mouse CII, and development of a strong autoantibody response to CII occurs (137). Furthermore, transgenic mice expressing the human HLA-DR4 (DRB1*0401) or DR1 (DRB1*0101) are susceptible to CIA (138–140). Some of these susceptible mouse strains will be discussed in more detail in study III. Both the human MHC II (DRB1) and the mouse H-2d bind an immunodominant CII peptide located at position 259-273 on triple helical CII (CII259-273) (30,140–142). Immune response to this peptide has also been detected in RA patients (6,143,144). The CII259-273 peptide binds to the DR4*0401 molecule with the major MHC anchors shifted three amino acids as compared with the binding of the same peptide to the H-2r molecule but the T cell receptor recognition sites are partly shared (140,145,146). Thus, these humanized mouse strains provide us with a CIA model that shares molecular structures of possible critical importance for the human disease.

4.1.1 Anti-CII antibodies in CIA

The need of B cells in CIA has been demonstrated in B cell deficient mice (147) and a strong B cell response to CII occur after CIA induction (23,148,149). Several studies have revealed detailed CII epitope analysis using the technique of monoclonal antibody production (23,150–155). To date, the most charaterized and dominant CII epitopes recognized by autoantibodies in CIA are denoted C1, U1, and J1. Additional CII epitopes have been discovered and are discussed in more detail in study I. The antibodies bind to conformation-dependent epitopes at distinct positions along the native molecule (156) and do not crossreact to either denatured CII or to other collagens. The structural importance of the anti-CII recognition has also been shown, as CIA cannot be induced in rats and mice by denatured CII (157,158).

There are several studies demonstrating the pathogenic nature of autoantibodies to CII directly in vivo (55,159–165). These studies show that anti-CII antibodies can induce arthritis in mice. As an example, injection of a cocktail of anti-CII monoclonal antibodies induce an very acute arthritis denoted collagen antibody induced arthritis (CAIA) (161,164). There is also evidence that the epitope specificity of the antibodies could have a major role in disease development since it previously was
shown that co-injection of a monoclonal antibody specific for epitope F4, denoted CII-F4 reduced the induction of arthritis in mice. Interestingly, antibodies to the F4 epitope were in fact negatively associated with RA whereas antibodies to epitope C1, J1 and U1, were positively associated (166). Autoantibodies to C1, U1, and J1 are well-characterized and dominates the response in CIA in mice (167,168) and rats (169). Monoclonal antibodies against these major CII epitopes show unique arthritogenic capacities. For instance the monoclonal antibody CIIC1 (discussed in detail in study II) show degradative effects in vitro on cartilage synthesis, and disorganization of fibrils in the extracellular matrix (16). Another pathogenic monoclonal antibody denoted M2139 (discussed in detail in study I and IV) cause thickening and aggregation of CII fibrils in the extra cellular matrix and abnormal chondrocyte morphology (170). Furthermore, a monoclonal antibody against the U1 epitope, UL1 induced proteoglycan depletion in vivo independent of inflammation (22), discussed more in study I.

4.1.2 T cell epitope on CII

The T cell epitope on CII was discovered by Michaëlsson et al (30) and consists of a 12 amino acid long sequence GIAGFKGEQGPK located at positions 259-270 on triple helical CII. The CII molecule can be posttranslationally modified (at position 264 and 270) by lysine hydroxylation and glycosylation. It was previously shown that the K at position 264 is the key amino acid residue for CII-specific T cell recognition in H-2d and HLA-DR4 mouse models (32,171). The modifications generate many possible T cell epitopes that seem to be recognized by specific T cell clones (171,172). Most of the T cells activated after immunization with heterologous CII react with the CII259-273 peptide.

4.2 CIA IN NON-HUMAN PRIMATES

CIA can also be induced in non-human primate species, most known and characterized is the rhesus monkey (Macaca mulatta). The animals are immunized with CII emulsified in CFA and swelling of joints develops after 3-4 weeks (173). The target tissue of disease onset is the synovium. Removal of the synovium by gene therapy in rhesus monkeys with clinically active CIA abolishes joint inflammation (174). Rhesus monkeys (of Indian/Burmese origin) positive for the Mamu B26 MHC class I marker are resistant to the development of clinical arthritis (175). The resistance is associated with the incapacity to produce an adequate IgM antibody response against the immunizing CII. By selecting animals lacking the Mamu B26 marker, a reproducible induction of the disease at an incidence > 95% can be obtained. CII-specific antibodies of the IgM isotype play a pivotal role in rodent models of CIA (176) and also in rhesus monkeys (177) and common marmosets (178).

Recently a new CIA model was introduced in the New World primate species, the common marmosets (178). This model shares many characteristics with the rhesus monkeys but at the level of histopathology, less cartilage and bone destruction is observed compared with the rhesus monkeys. Anti-CII antibody responses in these two strains will be discussed in study I.
Collagens are a large group of proteins that can be divided into several subgroups characterized by structure and function. Collagen type II (CII) belongs to the group called fibril-forming collagens. The group of fibril-forming collagens includes type I-III, V, and XI collagen. They are all similar in size and, they all contain large triple helical domains with about 1000 amino acids per chain (179). Already in the intracellular space they are going through modification and assembly, events involving post-translational hydroxylation and glycosylation, association of polypeptide chains, and folding of the triple helix. Collagens of this group are first synthesized as procollagen, characterized by non-helical extensions and the N- and C-termini that are then cleaved by specific proteinases after secretion (Figure 3). Extracellularly, they will self-assemble into fibrils followed by crosslinking of the fibrils. Another shared characteristic is that each molecule is displaced about one-quarter of its length relative to its nearest neighbor along the axis of the fibril (180).

The collagen triple helix is formed from three identical polypeptide chains that are each coiled around each other to form a right-handed super helix resembling a rope-like rod (181,182). Formation of this super-helical structure is facilitated by the presence of a glycine as every third amino acid. Due to the absence of a side-chain only glycine is small enough to fit in at specific positions in the center of the triple helix where the three chain come together. The amino acid proline is frequently in the X-position of the -Gly-X-Y- sequence and 4-hydroxyproline is frequently in the Y-position. The rigidity of these two amino acids favors specific backbone conformations and limits rotation of the polypeptide chains. A regular hydrogen-bonding network between the chains further stabilizes the triple helix. The conformation of the triple helix places the side-chains of amino acids in the X- and Y- positions on the surface of the molecule. This arrangement explains the ability of many collagens to polymerize, since the multiple clusters of hydrophobic and charged side-chains directly self-assemble into precisely ordered structures. The triple helix is relatively rigid. In some contexts, the resistance of the molecule to extension or compression is important for the biological function of the protein. In many collagens, the triple helix is interrupted by globular sequences that make the molecule more flexible.
Figure 3 Simplified illustration of how CII self assembles into fibrils and fibers. Three pro α-chains (A) self assemble and the procollagen triple helix forms (B). Specific proteinases cleave the non-helical ends of the triple helix (C) and the triple helical molecule self assemble into fibrils that can covalently crosslink to each other (D). Finally the fibrils aggregate to form CII fiber (E).
6 PRESENT INVESTIGATION

6.1 METHODOLOGIES

Methods used in paper I-IV are described in detail in the respective “Materials and Methods sections. Here follows an overview of the methods that mainly were used in this thesis with reference to the papers in which they were applied:

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<tr>
<th>Method</th>
<th>Description</th>
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<td>Induction and assessment of CIA, organ dissection, serum collection</td>
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<tr>
<td>Circular dichroism (CD) (II)</td>
<td>Analysis of the secondary structure of proteins</td>
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<tr>
<td>Fab preparation (II, IV)</td>
<td>Antibody digestion using papain and purification of Fab fragments</td>
</tr>
<tr>
<td>Gel filtration chromatography (I, II)</td>
<td>Purification technique to separate proteins according to size</td>
</tr>
<tr>
<td>Monoclonal antibody production (I, IV)</td>
<td>Process by which large quantities of antibodies can be produced by fusion of single antibody producing cells with tumor cells grown in culture</td>
</tr>
<tr>
<td>Polymerase chain reaction (PCR) (II, IV)</td>
<td>Amplification of DNA generating thousands to millions copies of a particular DNA sequence</td>
</tr>
<tr>
<td>Radioactive proliferation assay (III)</td>
<td>Assay to measure the proliferation rate of cells in vitro by incorporation of $^3$H-thymidine into the cell DNA</td>
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<tr>
<td>Recombinant protein production (I-II, IV)</td>
<td>Cloning, expression and purification of recombinant proteins</td>
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<td>Sequencing (II, IV)</td>
<td>Determining the nucleotide sequence of a gene using labeled probes</td>
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<td>Site-directed mutagenesis (II, IV)</td>
<td>Methods to introduce mutations at specific sites in plasmid DNA</td>
</tr>
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<td>Sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) (I-II)</td>
<td>Analysis of protein size and purity</td>
</tr>
<tr>
<td>Solid phase enzyme-linked immunosorbent assay (ELISA) (I-IV)</td>
<td>Analysis of antibody binding capacities to various targets in vitro</td>
</tr>
<tr>
<td>Statistical analysis (I-IV)</td>
<td>One-way Anova (Kruskal-Wallis test), Dunn’s multiple comparison tests, Wilcoxon singed rank test, Mann-Whitney test</td>
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T cell in vitro assays (III) Assay based on re-stimulation of cell cultures with various agents, followed by measuring cytokines in the supernatant
<table>
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<th>X-ray crystallography (II, IV):</th>
<th></th>
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<td>Protein crystallization by hanging and sitting drop vapour diffusion</td>
<td>Technique to generate protein crystals for use in X-ray crystallography</td>
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<tr>
<td>Molecular replacement (MR)</td>
<td>X-ray crystallography method using diffraction data from crystals of a protein in combination with an already solved crystal structure of a homologue for structural determination</td>
</tr>
<tr>
<td>Model building and refinement</td>
<td>Computer analysis using the obtained electron density maps to build a first model of the structure. The model is then refined (improved) through statistical adjustment of the atomic coordinates</td>
</tr>
</tbody>
</table>
6.2 STUDY I

Analysis of autoantibodies to the joint specific protein collagen II in experimental and rheumatoid arthritis

This study was undertaken in order to obtain a more comprehensive understanding of CII epitope specific responses in mice and monkeys with CIA and in patients with established RA. The work was based on the generation of a recombinant CII peptide library covering overlapping sequences from the entire triple-helical region of CII. CIA was induced in B10.Q (BALB/c x B10.Q) N2 developing a chronic relapsing disease and in B10.Q developing an acute arthritis. Sera were collected at different disease timepoints and analyzed for autoantibody specificity. Autoantibodies specific for several hitherto unknown CII epitopes were discovered and earlier discovered CII epitopes were confirmed. The epitope predominance varied during the disease course. The specific autoantibody response detected in mice was also apparent in non-human primate species (rhesus monkeys and common marmosets) with CIA. Furthermore, immunodominant CII epitopes with known biological significance in mice were tested for autoantibody reactivity in serum and synovial fluid of patients with established RA. All three investigated CII epitopes (C1, U1, and J1) were detected in a subgroup of patients and interestingly they were all significantly increased in the synovial fluid, in particular autoantibodies against epitope U1 with frequencies reaching close to 70%.

6.3 STUDY II

Crystal structure of an arthritogenic anti-collagen immune complex

Little is known about the structural specificity of a pathogenic autoantibody when interacting with CII. This study was therefore performed to investigate how one of the most pathogenic autoantibodies in mice, CIIC1 recognizes its epitope (C1) on cartilage CII. The CIIC1 fab was produced and mixed with the triple-helical C1 peptide and the crystal structure of the complex was determined to 2.2 Å resolution, revealing how antibody and epitope interact in molecular detail. The importance of the observed interactions was further investigated using site directed mutagenesis of a recombinant C1 peptide. The antibody binding to the C1 peptide did not introduce any significant conformational changes in the antibody although the peptide, which retained its typical triple-helical conformation upon antibody binding, penetrated rather deep into the pocket. All three peptide chains forming the epitope were in contact with the antibody. The antibody contacts the CII peptide primarily via germline-encoded residues. Interestingly the antibody clearly recognized the sequence “R-G hydrophobic”, a motif that has been suggested to be a consensus recognition sequence in CII epitopes and potentially a recognition site for autoantibodies. The arginine at position 360 (R360) from all three chains contributed significantly to the binding. Its crucial role was further confirmed by site directed mutagenesis, showing that substitution of R360 by alanine completely destroys the interaction between the antibody and the peptide. All six CDRs interacted with the triple-helical peptide. CDRs H2 and L3 contribute most to the area buried upon peptide binding, while CDR L2 contributes less, mainly because the loop residues are located rather far away from the epitope.
6.4 STUDY III

Comparative analysis of collagen type II-specific immune responses during development of collagen-induced arthritis in two B10 mouse strains

In this study we wanted to understand the magnitude and specificity of how CII-specific T cells respond to modified variants of the T cell epitope in CIA, but also how autoantibodies recognize major CII epitopes over the course of arthritis. The specific T and B cell response was investigated in two mouse strains, B10.Q and a humanized transgenic strain B10.DR4.Ncf1*/* after CII immunization. Serum and draining lymph nodes were collected at four timepoints during the disease course and used for analysis of the CII-specific T cell as well as antibody response in the two mouse strains. The analyses revealed no clear shift in either antibody specificity or T cell reactivity between the phases in either of the two strains. Instead, the T cell recall response stayed rather constant with dominant response to the non-modified (K264) and the hydroxylated (HOK264) CII259-273 peptide in the humanized mice and to the glycosylated CII259-273 (GalOK264) peptide in the B10.Q mice. One was noted at the initial stage of CIA in B10.Q mice when a relatively equal T cell reactivity to non-modified (K264) and to the galactosylated (GalOK264) was detected. The C1 epitope was the most recognized of the epitopes investigated in both strains. Antibodies to epitope J1 were almost negligible and the concentration of antibodies to epitope U1 was elevated at all timepoints investigated in the humanized mouse model. In addition, we revealed that the T cell response had much higher magnitude in B10.DR4.Ncf1*/* compared to B10.Q. Moreover, titers of anti-CII antibodies as well as anti-C1 antibodies were significantly higher in arthritic mice compared with healthy mice, and so was also the antibody response to U1 and J1 in the transgenic mice.

6.5 STUDY IV

Immunoglobulin heavy chain V gene polymorphism controls epitope specific antibody response

It was recently shown that production of anti-CII antibodies, in particular antibodies specific for the J1 epitope, strongly associate with the IgHv locus on mouse chromosome 12 (168). These studies were performed using high resolution mapping in a heterogeneous stock (HS) mouse cohort (183). In order to better understand this finding we investigated the specific anti-CII response to the arthritogenic J1 epitope on CII in detail.

Sequencing of J1-specific monoclonal antibodies revealed a common IgHv gene usage. Comparing this sequence with those encoded by the same gene in all strains comprising the HS, two major polymorphisms were found in the CDR1 domain (S31R and W33T). In order to investigate the impact of these two polymorphisms upon antibody binding a pathogenic monoclonal antibody M2139_fab were crystallized together with the triple helical J1 peptide. The crystallized M2139_fab–J1 complex confirmed a strong interaction of these two amino acid residues with the triple helical J1 peptide. By modeling, almost all interactions were lost after replacing the two polymorphic sites in the CDR1 region suggesting that these positions are major contributors to the M2139_fab -J1 affinity. This was further confirmed using recombinant single chain fragments (ScFv) and site-directed mutagenesis.
The crystal structure was also used to compare the immune complex with the CIIC1\textsubscript{fab}-C1 antibody complex described earlier (184). We observed clear differences in how the collagen peptides are oriented in the antibody-combining site, with C1 remaining relatively straight and the J1 peptide bending into the pocket. Although the structures of both fab fragments superimpose remarkably well and the CDR sequences show a considerable degree of structural and sequence conservation, none of the crucial interactions with the peptide are conserved. M2139\textsubscript{fab} -J1 makes only about half as many polar interactions to the J1 peptide than CIIC1 forms with C1. The major recognition motif also markedly differed with M2139\textsubscript{fab} compared to CII-C1 was detected. The major recognition motif also markedly differed with M2139\textsubscript{fab} primarily binding to I_{26}A_{27}G_{28} and CIIC1\textsubscript{fab} recognizing the R_{18}G_{19}L_{20}. 
7 DISCUSSION AND FUTURE PERSPECTIVE

Unraveling CII autoimmunity is an important task in order to understand the basic mechanisms operating in autoimmune diseases such as RA. The research I have performed during my PhD have answered several important questions and provided new insights into these areas but they have of course also raised new questions. I will below outline my views of the result obtained and further suggest how to continue with the research within this field.

One of the purposes of this thesis was to build a more comprehensive understanding of biological significant CII epitopes in CIA and RA. As earlier discoveries were limited to analyses using large CII fragments and specificity of monoclonal antibodies established from the very initiative phase of CIA, we decided to generate a library consisting of 70 overlapping CII peptides covering the entire triple-helical region of CII that are fused with a foldon trimerization domain (185). This library allowed detection of specific epitope binding and revealed the autoantibody response at different time points in a chronic mouse model of CIA. The results indicate that the three CII epitopes C1, U1, and J1, known to be the most common epitopes, become “challenged” by several others that appear as strong or stronger. The majority of the newly identified epitopes are localized on the cyanogen bromide (CB) fragment 11, a region where several important epitopes are located (23). This region seems to more easily attract arthritogenic antibodies. The study further shows the epitope specificity during different stages of the disease. Clearly, we show that some CII epitopes are more frequently recognized in early CIA whereas others appear later in the chronic phase of the disease. This could perhaps depend on the epitope location on CII. Some parts of the CII molecule might not be accessible for binding until cartilage breakdown has been ongoing for a while whereas others are more easily recognized and accessible. If this is the case, different epitopes could be used as markers for different stages of the disease. In order to get a better picture of CII epitope specificities in arthritis it is of course very important to continue discovering the new epitope regions and to study the cartilage degradation effect/s of autoantibody binding to these epitopes. The CII-specific studies were continued by analysis of the epitope specific response (C1, U1, and J1) in serum and synovial fluid from RA patients. A clear overrepresentation of all three CII epitopes was found in the synovial fluid with surprisingly high frequencies of the U1 epitope suggesting that there is local antibody production. To date, we do not have a direct explanation for the elevated levels measured for epitope U1. But these findings could just reflect the importance of investigating CII-specific responses using smaller peptides rather than the entire CII or CB fragments. Earlier results using CII have given inconsistent results partly depending on the not 100% pure CII batches used, leading to contaminants such as other matrix molecules surrounding CII. These molecules could potentially mask the epitopes. So maybe the pure peptide antigens give the more correct response? The library could also be used to reveal CII specific responses in non-human primate species although the assay was partly modified due to crossreactivities to the foldon domain attached at one end of the triple-helical peptides. Although the library generated could be used for many purposes, we suggest introducing an even more comprehensive library including additional CII epitopes that are posttranslationally modified (glycosylated, citrullinated, and hydroxylated).
Moreover it is important to generate even better and more sensitive assays to exactly reveal how RA patients respond to the CII determinants. The way CII peptides are synthesized should also be taken into account to reach the best results.

In the next two studies we wanted to structurally determine how pathogenic autoantibodies recognize native CII in its triple-helical state. These studies were initiated using recombinant CII peptides attached to the foldon domain. However, the resolution reached was not enough to clearly see the interactions between antibody and antigen, thus we used synthetic triple helical CII peptides also for the crystallographic studies. M2139\textsubscript{Fab}-J1 and CII\textsubscript{C1}-C1 complexes share some common characteristics but have completely different recognition sequences. It has earlier been proposed that the so called “R-G-hydrophobic” motif found in almost all discovered epitopes could be a recognition sequence for autoreactive B cells (23). From the two crystal structures and from finemapping analysis it is clear that anti-CII antibodies do not necessarily share recognition residues in their epitopes. The postulated recognition motif could be of importance for most but not all CII autoantibodies, as indicated by the completely different recognition sequence of antibody M2139. However, although differences exist between the two complexes our finemapping studies still argue for that “R-G”, not necessarily followed by a hydrophobic residue, is a common and crucial recognition motif for at least some anti-CII antibodies. This amino acid selection could partly be explained by the surface exposure of arginine (R) due to its long side chain and charged guanidinium group. We already have projects ongoing that aim at the structure determination of more pathogenic anti-CII antibodies, and hopefully this will increase our knowledge base and understanding of these characteristics in even more detail. In this study we also revealed that contribution of the CDRs to the interaction with the peptides differs: in the M2139 complex VH1 and VL3 provided most to the area buried upon peptide binding, in CII\textsubscript{C1} VH2 and VL3. Although the CII epitopes are not identical it has earlier been shown earlier that anti-CII antibodies commonly derive from the same V-gene family. That could be based on the similarity of CII epitopes with regard to being located on a conserved triple helical molecule with high prevalence of repeated sequences.

The main purpose of this thesis was to describe structural requirements for how autoreactive B cells recognize CII. However, since both B and T cells are important for CIA we undertook one additional study aiming to analyze both CII-specific B and T cell responses. CIA was induced in two mouse strains B10,Q and the humanized DR4.Ncf1\textsuperscript{+/+} on a B10 genetic background and autoantibodies and IFN-\(\gamma\) levels were measured at different timepoint during the disease course. We show in this study that both T and B cell responses are clearly present but both the magnitude and specificity of the T cell response differ remarkably between the two strains. The B10.DR4.Ncf1\textsuperscript{+/+} shows higher levels of IFN-\(\gamma\) at all phases investigated compared with B10,Q. The reason for this difference is entirely clear but one could suggest that the Ncf1 mutation has an effect of the results since it regulates susceptibility and severity of CIA (188). Another explanation could be differences in the number of T cells expressing CD4\textsuperscript{+} co-receptors between the two strains. In the humanized mouse model all T cells including CD8\textsuperscript{+}, express CD4\textsuperscript{+} co-receptor on the cell membrane leading to that also CD8\textsuperscript{+} T cells could be responsible for the observed response. Therefore the magnitude in T cell responses between these transgenic and non-transgenic mice needs to be optimized.
before results can be conclusive. One way to circumvent this bias could be by establishing a DR4 (and human CD4) knock-in mouse instead of using a transgene. This would lead to a more physiological expression of the proteins. When investigating the extent to which T cells recognize posttranslational modifications of CII in DR4 and B10.Q mice during the disease course we revealed that the non-glycosylated CII259-273 peptide was clearly immunodominant in DR4 at all phases investigated whereas the galactosylated epitope dominated in H-2\textsuperscript{b} expressing mice. Thus the H-2\textsuperscript{b} mice show more similarities to the CII-specific T cell response observed in human RA (6). The explanation for this biased result is currently unknown but maybe also in this case non-physiological expression of the DR4 transgene might influence which CII peptides that will become immunodominant.
Det är en speciell och intensiv period i mitt liv som är på väg mot sitt slut. En tid som gett mig oerhört mycket på många plan, både forskningsmässigt och privat. Nu börjar en ny spännande resa och det är ofta just i detta läge man tittar tillbaka på alla fina människor man har och har haft omkring sig och som varit så fantastiska lärare, alla på olika sätt. Det finns vissa som har varit extra betydelsefulla och som jag vill säga ett speciellt stort tack till. Ni är:

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