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The Ectopic Germinal Centre Response in Autoimmune Disease and Cancer

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1. Introduction

1.1 The B-cell response in autoimmune disease
The pathological effects of autoimmune diseases on the target tissues can be mediated by autoantibodies, cell-mediated immune responses, or both. It is increasingly evident that some autoimmune diseases previously thought to be essentially T-cell-mediated also have a B-cell component, which may involve direct effects of autoantibody secreted by plasma cells, pro- or anti-inflammatory cytokines secreted by activated effector or regulatory B-cells, or through the highly efficient antigen presentation function of B-cells enabling them to activate CD4+ T-cells and vice versa. The number of autoimmune diseases known to be mediated partly or largely through autoantibodies has increased markedly in recent times. Systemic lupus erythematosus (SLE), in which the pathology is mediated via Type II & III hypersensitivity reactions involving anti-DNA autoantibodies, has long been known to fall into this category. Many other autoantibodies are produced by these patients, principally against nuclear antigens, but most are not thought to be involved in pathology. Hashimoto’s thyroiditis and Graves’ disease patients produce pathogenic autoantibodies against thyroid antigens, the latter being a rare example of an activating autoantibody inducing signalling via the thyroid stimulating hormone receptor. Myasthenia gravis patients produce autoantibodies against the acetylcholine receptor (AChR), present on the motor muscle endplates, thereby inhibiting muscle contraction. Anti-SS-A and anti-SS-B (anti-Ro & anti-La) autoantibodies are implicated in congenital heart block in children born to mothers with Sjögren’s Syndrome due to transplacental uptake of IgG autoantibodies; autoantibodies against α-fodrin are also believed to be pathogenic in these patients. Rheumatoid arthritis (RA), one member of the group of systemic rheumatic autoimmune diseases that also includes SLE, psoriatic arthritis and the various forms of myositis, has now gone full cycle in views on its pathological mechanisms.

1 Abbreviations: AChR, Acetylcholine Receptor; AID, Activation Induced Cytidine Deaminase; AMC, Arthrogryposis Multiplex Congenita; ARS, Anti-amino acyl-tRNA Synthetase; Bmem, Memory B-cell; CDR, Complementarity Determining Region; DM, Dermatomyositis; EOMG, Early Onset Myasthenia Gravis; FDC, Follicular Dendritic Cell; G.C., Germinal Centre; IBM, Inclusion Body Myositis; IM, Inflammatory Myopathies; LOMG, Late Onset Myasthenia Gravis; MAA, Myositis-Associated Autoantibodies; MAb, Monoclonal Antibody; MIR, Main Immunogenic Region; MSA, Myositis Specific Autoantibodies; PM, Polymyositis; RA, Rheumatoid Arthritis; SLE, Systemic Lupus Erythematosus; TAA, Tumour-Associated Antigen; TIL, Tumour-Infiltrating Lymphocytes; Tfh, Follicular T helper cell; UNG, Uracil Nucleotidyl Glycosylase.
Initially thought to be caused by the anti-IgG Fc antibody (rheumatoid factor), although approximately 25% of RA patients are rheumatoid factor negative, the evidence then swung in favour of a cell mediated autoimmune response involving effector T-cells and cytokines, principally TNFα. Although these are clearly involved in joint pathology, autoantibodies against cyclic citrullinated proteins are a much better diagnostic marker for RA than rheumatoid factor and there is some limited evidence that they may be pathogenic. It is also now recognised that B-cells play an important role in the pathogenic autoimmune response, as clearly demonstrated by the marked clinical improvement in patients treated with Rituximab®, an anti-CD20 chimaeric (human/mouse) monoclonal antibody that suppresses B-cell responses. Other autoimmune diseases with B-cell involvement include autoimmune haemolytic anaemia, idiopathic thrombocytopenia, Type I diabetes, and some subtypes of myositis, although the situation is often confused by the presence of non-pathogenic autoantibodies.

1.2 The germinal centre response to foreign antigens

Germinal centres (g.c.) are the main sites of generation of high affinity, antibody-secreting plasma cells and Ig class-switched memory B-cells during T-cell-dependent immune responses, extensively reviewed by others (Allen, Okada, & Cyster, 2007; Brink, 2007; Hauser, Shlomchik, & Haberman, 2007; Klein & Dalla-Favera, 2008; Leavy, 2010; Minton, 2011). Here we shall summarise briefly the principal features of the g.c. response. The response is initiated by B-cells binding to their cognate antigen on the surface of antigen presenting cells, such as dendritic cells, in a secondary lymphoid organ (lymph node, spleen, Peyer’s patches or human tonsil). The antigen becomes internalised, degraded into peptides which are expressed on the cell surface bound to MHC Class II and presented to a helper T (T\_h) cell that provides costimulatory activation signals, including binding of the B-cell surface molecule CD40 to its ligand, CD154, on the T-cell membrane.

This interaction takes place at or near the interface between the B-cell follicle and the T-cell area and some activated B-cells proliferate outside the follicle and differentiate into short-lived plasma cells secreting IgM antibodies. Others migrate into the B-cell follicle where they proliferate and differentiate into centroblasts expressing low levels of surface Ig. This region develops into a germinal centre with a dark zone of densely packed, proliferating centroblasts, and a light zone of more loosely packed B-cells (centrocytes) interspersed with the processes of follicular dendritic cells (FDC, Figure 1A). These have distinct stromal origins, unlike the bone marrow derived, extra-follicular dendritic cells; almost uniquely, their C’ and Fc receptors trap immune complexes and retain antigens in their native state for months.

The pre-existing IgM⁺,IgD⁺ follicle B-cells are pushed out to form a mantle zone around the developing germinal centre, the whole structure being termed a secondary follicle. Proliferation of the dark zone centroblasts is extremely rapid, with cell cycle times estimated at between 6 and 12 hours. These proliferating clones of B-cells switch on the molecular machinery required for somatic hypermutation of their rearranged, expressed, Ig V-genes, including expression of activation-induced cytidine deaminase (AID). This induces mutations specifically targeted to the Ig V-genes at a frequency of 1 per 1000 base pairs per cell division, although much lower levels of mutation can also occur in some other, non-Ig genes such as Bcl-2 & Bcl-6. AID deaminates cytidine to uracil at C/G base pairs, introducing mismatches in the DNA that can be replaced by T/A base pairs. Uracil nucleotidyl glycosylase (UNG) can remove the uracil leading to insertion of any of the four bases at the abasic site; mismatch repair enzymes also recognise the mismatch and induce
single strand breaks which are repaired by error prone DNA polymerases (Di Noia & Neuberger, 2007). The mutations are targeted mainly to the complementarity determining regions (CDRs) which are intimately involved in binding to the epitope and therefore determine specificity and affinity of the antibody. Combined with the rapid proliferation, this results in clones of B-cells expressing receptors with a variety of affinities for the antigen, some high, some low; some will have lost the ability to bind to the antigen altogether and rare B-cells may cross-react with a self-antigen.

Several clones of proliferating, mutating B-cells are usually present within each germinal centre. These cells differentiate into centrocytes expressing mutated antigen receptors and migrate into the light zone. The centrocytes move through the light zone, acquire antigen for a second time from immune complexes on the follicular dendritic cells, which they internalise, process and present to follicular helper T-cells (T_{fh}-cells) (Patakas et al., 2011), thereby receiving survival signals, probably via costimulatory molecule interactions including CD40/CD154 and CD80/CD28 binding. These signals, together with T_{fh}-cell cytokines (IL-4 and IFN_{γ}) and AID deamination of cytidines, promote induction of class switch recombination (Patakas et al., 2011). Some of these centrocytes differentiate directly into plasmablasts and antibody-secreting plasma cells; others differentiate into Ig class switched memory B-cells, both of which migrate out of the follicle. Competition for limiting availability of antigen results in selection of B-cells expressing high affinity antigen receptors; recent evidence has shown that a broad range of mutations is involved in selection, not only for high affinity receptors but also for stability and expression of the B-cell receptor (Weiser et al., 2011). Cells expressing antigen receptors with low affinity are unable to compete for survival signals and the default response is that they die by apoptosis and are engulfed by macrophages, in which their degenerating nuclei are visible as tingible bodies. Most of this information is derived from studies in mice, in which the germinal centres reach maximum size about two weeks after immunisation and then gradually decline in the absence of further immunisation, disappearing after several weeks. Although the cell composition and structure of secondary follicles appear similar in Man, the kinetics and some of the detailed cellular interactions may differ.

Detailed studies of the kinetics and cellular interactions within germinal centres using multiphoton microscopy of living tissue in combination with B & T-lymphocytes expressing defined antigen receptors from transgenic animals have revealed much more dynamic activity than was previously suspected. It is now recognised that there is less distinction between the dark and light zones than suggested by static immunohistological examination, and there is continual recycling of B-cells both between and within the two zones, although there is net migration from the dark zone to the light zone (Beltman et al., 2011)(Figure 1B). Centrocytes move rapidly through the network of follicular dendritic cell processes, apparently sampling the immune complexes attached to their membranes and some of these cells return to the dark zone for further rounds of proliferation and somatic hypermutation. Migration of B-cells between the zones is controlled by chemokines, possibly secreted by stromal cells within the germinal centre. T_{fh}-cells are present mainly in the light zone and recent data suggest that affinity selection of B-cells may involve competition for signals from cognate T_{fh}-cells via peptide/MHC Class II binding as well as, or instead of, competition for antigen on the surface of follicular dendritic cells (Victora et al., 2010). Anti-self B-cells that have escaped negative selection in the bone marrow, or have arisen in the germinal centre due to somatic hypermutation, are either eliminated at this stage, suppressed by regulatory T-cells, or alter their antigen specificity by receptor revision, a process similar to V-gene rearrangement in
developing B-cells. This involves re-expression of RAG1 and RAG2 and rearrangement of an upstream light chain V-gene to an unused J exon (Nemazee, 2006). Despite the absence of D exons in the rearranged heavy chain locus, we have shown that an upstream heavy chain V-gene can also replace all or part of a rearranged V_{H}-gene, thereby altering the specificity of the receptor away from self antigen (Darlow & Stott, 2005). The architecture, cellular components and processes occurring in a typical germinal centre are summarised in Figure 1.

Fig. 1. Diagrammatic representation of a germinal centre in a lymph node.

A: Showing a dark zone containing proliferating clones of mutating centroblasts and a light zone containing centrocytes in contact with follicular dendritic cells and follicular helper T-cells (Tfh cells). Long-lived memory B-cells, plasmablasts and plasma cells secreting antibody molecules migrate out of the g.c. and leave the lymph node via the efferent lymphatic vessel. Apoptotic B-cells, macrophages containing tingible bodies and the mantle zone are not shown.

B: The same germinal centre showing recirculation of B-cells within and between the dark and light zones.

1.3 The ectopic germinal centre response in autoimmune disease

It has been known for many years that the target tissues of autoimmune diseases contain infiltrating lymphocytes and other immune cells, including T-cells, B-cells, plasma cells, macrophages, dendritic and follicular dendritic cells. In many cases the infiltrating cells organise themselves into structures resembling germinal centres. Some of these have a mantle zone, suggesting that they were formed from a primary follicle whereas, even when absent, it is often possible to distinguish a dark zone, containing few or no CD4^{+} T-cells or follicular dendritic cells, and a light zone containing both. Autoantigens have been identified on the finger-like processes of follicular dendritic cells (Shiono et al., 2003) and, in some cases, autoantibodies have been identified in g.c. B-cells. Separate T-cell areas containing dendritic cells and, sometimes, high endothelial venules, can also be seen. The stage of lymphoid
neogenesis appears to be directly related to the extent of infiltration of lymphoid and other immune cells (Aloisi & Pujol-Borrell, 2006). Examples of autoimmune diseases in which germinal centre-like structures have been identified in the target, or disease-related tissues are shown in Table 1. It is now apparent that ectopic germinal centres, also known as tertiary lymphoid organs, can also develop in other chronic inflammatory diseases, such as the gut in Crohn’s disease and ulcerative colitis patients, in chronic infections (Aloisi & Pujol-Borrell, 2006) and some types of cancer (Table 1). The questions these observations raise are: 1. How do they develop?; 2. How closely do they resemble germinal centres in secondary lymphoid organs?; 3. Are the B-cells within them undergoing a germinal centre response, as described in section 1.2 above?; 4. Are they generating plasma cells secreting pathogenic autoantibodies?; 5. What role do they play in the pathogenesis of autoimmune disease?

<table>
<thead>
<tr>
<th>Autoimmune Diseases</th>
<th>Organ containing Germinal Centres</th>
<th>Antigen(s) Recognised by GC B-cells</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hashimoto’s thyroiditis</td>
<td>Thyroid</td>
<td>Thyroglobulin, Thyroperoxidase</td>
<td>(Knecht, Saremaslani, &amp; Hedinger, 1981) (Armengol et al., 2001)</td>
</tr>
<tr>
<td>Graves’ disease</td>
<td>Thyroid</td>
<td>Thyroglobulin, Thyroperoxidase</td>
<td>(Armengol et al., 2001)</td>
</tr>
<tr>
<td>Myasthenia gravis</td>
<td>Thymus</td>
<td>Acetylcholine receptor</td>
<td>(Yoshitake et al., 1994) (SHIONO et al., 2003)</td>
</tr>
<tr>
<td>Sjögren’s syndrome</td>
<td>Salivary glands</td>
<td>SS-A (Ro), SS-B (La)</td>
<td>(Stott et al., 1998)</td>
</tr>
<tr>
<td>Rheumatoid arthritis</td>
<td>Synovial membranes of joints</td>
<td>IgG Fc, Cyclic citrullinated protein/peptide</td>
<td>(Manzo &amp; Pitzalis, 2007) (Humby et al., 2009)</td>
</tr>
<tr>
<td>Psoriatic arthritis</td>
<td>Synovial membranes of joints</td>
<td>?</td>
<td>(Canete et al., 2007) (Gerhard et al., 2002)</td>
</tr>
<tr>
<td>Cryptogenic fibrosing alveolitis</td>
<td>Lungs</td>
<td>?</td>
<td>(Wallace et al., 1996)</td>
</tr>
<tr>
<td>Uveoretinitis</td>
<td>Choroid of the eye</td>
<td>?</td>
<td>(Liversidge et al., 1993)</td>
</tr>
<tr>
<td>Autoimmune hepatitis</td>
<td>Liver</td>
<td>?</td>
<td>(Mosnier et al., 1993)</td>
</tr>
<tr>
<td>Multiple sclerosis</td>
<td>Meninges (?)</td>
<td>?</td>
<td>(Prineas, 1979) (Serafini et al., 2004)</td>
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<tr>
<th>Chronic Inflammatory Diseases</th>
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<tr>
<td>Crohn’s disease</td>
<td>Gastrointestinal tract</td>
<td>?</td>
<td>(Kaiserling, 2001)</td>
</tr>
<tr>
<td>Ulcerative colitis</td>
<td>Descending colon</td>
<td>?</td>
<td>(Kaiserling, 2001)</td>
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<tr>
<th>Infectious Diseases</th>
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<tr>
<td>Chronic hepatitis C infection</td>
<td>Liver</td>
<td>?</td>
<td>(Mosnier et al., 1993)</td>
</tr>
<tr>
<td>Helicobacter pylori or Campylobacter gastritis</td>
<td>Stomach</td>
<td>Bacterial antigens</td>
<td>(Genta, Hamner, &amp; Graham, 1993) (Stolte &amp; Eidt, 1989)</td>
</tr>
<tr>
<td>Chronic Lyme disease</td>
<td>Synovial membranes of joints</td>
<td>?</td>
<td>(Ghosh et al., 2005)</td>
</tr>
<tr>
<td>Oncocerciasis</td>
<td>Skin</td>
<td>?</td>
<td>(Brattig et al., 2010)</td>
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<tr>
<th>Cancers</th>
<th></th>
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<tr>
<td>Lymphoma of MALT associated with Sjögren’s Syndrome</td>
<td>Lymphoma</td>
<td>?</td>
<td>(Bombardieri et al., 2007a)</td>
</tr>
<tr>
<td>Ductal breast carcinoma</td>
<td>Breast tumour</td>
<td>Epidermal growth factor receptor family</td>
<td>(Coronella et al., 2002) (Nzula, Going, &amp; Stott, 2003a) (Simsa et al., 2005) and section 5.</td>
</tr>
<tr>
<td>Medullary breast carcinoma</td>
<td>Breast tumour</td>
<td>Ganglioside</td>
<td>(Coronella et al., 2001) (Kotlan et al., 2005)</td>
</tr>
</tbody>
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Table 1. Diseases in which ectopic germinal centres have been observed.
It has now been shown by combined immunohistochemistry, identification of antigen specificity of B-cells and plasma cells in and around ectopic germinal centres, and sequence analysis of expressed, rearranged Ig V-genes and their somatic mutations, that germinal centre B-cells in the target tissues of several autoimmune diseases are undergoing clonal expansion, somatic hypermutation and affinity selection, in a similar manner to that seen in the germinal centres of secondary lymphoid organs (Table 1 and section 1.2). This has been demonstrated in Sjögren’s syndrome, rheumatoid arthritis, psoriatic arthritis, myasthenia gravis, multiple sclerosis and also in breast cancer. In some of these cases, expression of RAG1 and 2 have been observed (Armengol et al., 2001), indicating that receptor revision also takes place in ectopic germinal centres and therefore the generation and attempted elimination of self-reactive B-cells. The signals involved in tertiary lymphoid organ neogenesis appear to be similar to those in development of secondary lymphoid organs, although the temporal and causal relationship between appearance of these structures in the target tissue and autoimmune pathology-related tissue damage is unclear. One scenario is that an initial event in the tissue, which could, in some cases, include microbial infection, leads to the release of molecules seen by the immune system as “danger signals” (Matzinger, 2007) thereby inducing infiltration of inflammatory cells and subsequent lymphoid neogenesis, causing further tissue damage with concomitant release of self-antigens, more danger signals and a vicious cycle, perpetuating a chronic autoimmune reaction. Alternatively, initial tissue damage may be caused by an autoimmune response commencing in the secondary lymphoid organs, with subsequent events following a similar course to that described above. Lymphotoxins α, β, αβ2 and TNFα have been shown to be required for development of ectopic germinal centres. Growth-factor receptor-bound protein-2 (Grb2) has recently been shown to control orthotopic lymphoid follicle organisation and the germinal centre response by inducing production of lymphotoxin-α via CXCR5 signalling (Jang et al., 2011). These molecules are secreted by infiltrating B and Tfh-cells and activated NK cells; on binding to their receptor on stromal cells they induce expression of adhesion molecules and secretion of chemokines which induce further lymphocyte infiltration and segregation into B-cell follicles, formation of a follicular dendritic cell network and T-cell areas. It has also recently been proposed that overexpression of costimulatory molecules on Tfh-cells may contribute to overcoming B-cell tolerance (Patakas et al., 2011). This may be a contributory factor in ectopic as well as orthotopic germinal centres. Primary B-cell follicles are rarely seen in autoimmune disease target tissues but this may be because chronic antigen stimulation has been in progress for a considerable time before biopsies are taken. For example, in type I diabetes mellitus there is evidence that the autoimmune response develops long before overt disease is diagnosed. Whether ectopic germinal centres are initiated by naïve or memory B-cells is unclear but recent evidence shows that at least some B-cell clones arise de novo from naïve B-cells (Sims et al., 2001; Nzula, Going, & Stott, 2003b; Nzula, Going, & Stott, 2003a). The frequency of ectopic germinal centres varies markedly between autoimmune diseases; as one might expect, the highest incidence is in diseases where pathogenic autoantibodies are most strongly implicated. Thus, they have been identified in thyroid tissues of 100% of Hashimoto’s thyroiditis patients and 54 – 63% of Graves’ disease cases; in rheumatoid arthritis the figure is 25 – 50% but in Sjögren’s syndrome it is only 17%, although variations may to some extent reflect differences in the difficulty of finding the germinal centres. In Sjögren’s syndrome, the source is usually biopsies of the small labial salivary glands of which there is a large number; as g.c.s are only present in some of the many small labial
salivary glands, they may easily be overlooked. Tissues containing different types of cells respond in a variety of ways to inflammatory signals and this may also determine whether, and to what extent, lymphoid organ neogenesis occurs. The origin of follicular dendritic cells is unclear but it has been proposed that they develop from precursor cells already present in the tissue, either fibroblasts or fibroblast precursor cells (Park & Choi, 2005). Alternatively, the precursor cells may be induced to migrate into the tissue by the same or similar chemokines as those attracting the B and T-lymphocytes. In several autoimmune diseases (Table 1) and animal models of autoimmune diseases (Astorri et al., 2010; Nacionales et al., 2009), it has been demonstrated that ectopic germinal centres are generating plasma cells secreting pathogenic autoantibodies and, almost certainly, memory B-cells bearing anti-self antigen receptors, implying that they aid the diversification of the autoantibody repertoire and contribute to the maintenance of immune pathology. In addition to autoantibody production, self-reactive B-cells generated in ectopic germinal centres may also contribute to autoimmune pathology by secretion of pro-inflammatory cytokines and activation of pathogenic T-cells by presentation of processed self-antigens. B-cells may contribute in this way to immune pathology in autoimmune diseases generally considered to be principally T-cell mediated, and may be one explanation for the efficacy of Rituximab therapy for rheumatoid arthritis.

2. Methods

2.1 Identification and cellular composition of ectopic germinal centres
The methods we used to identify ectopic germinal centres, characterise their cellular composition, analyse the rearranged Ig V-gene sequences expressed by germinal centre B-cells and identify their antibody specificity have been described in detail in previously published papers (Nzula, Going, & Stott, 2003a; Sims et al., 2001). Briefly, sections were cut from snap frozen tissue biopsies and every tenth section stained for B-cells with anti-CD20. Sections containing germinal centre-like structures or B-cell aggregates were further characterised by staining for T-cells (anti-CD3, CD4, CD8), regulatory T-cells (anti-FoxP3), follicular dendritic cells (anti-FDC (DAKO) or anti-CD35), plasma cells (DAKO), macrophages (anti-CD68) and proliferating cells (anti-Ki67). Double immunofluorescent staining with the above cell subset-specific antibodies and Ki67 was used to identify dividing cells. Acetylcholine receptor-specific B-cells in germinal centres from the thymus of myasthenia gravis patients were identified by $^{125}$I-α-bungarotoxin-labelled acetylcholine receptor and autoradiography (Shiono et al., 2003; Hill et al., 2008); other autoantibody-producing cells were identified by immunofluorescence staining with the relevant antigen.

2.2 Cloning and sequence analysis of rearranged Ig V-genes
Ectopic germinal centres and B-cell aggregates were excised by microdissection, digested with proteinase K and the released DNA used as a template for amplification of the rearranged Ig V-genes by nested PCR. Details of the method and the primers are described in Sims et al. (2001) and Nzula et al. (2003). Amplified DNA was purified by agarose gel electrophoresis, ligated into plasmid DNA and cloned in E. coli. Cloned plasmid DNA was purified and the Ig V-genes sequenced in both directions using primers complementary to sequences flanking the cloning site. The best matching germline V, D & J sequences were identified initially by comparison with the VBASE directory of human Ig V-genes and later, after VBASE ceased to be updated, using the Immunogenetics (IMGT) Database of Human
Immunoglobulin Sequences (http://www.imgt.org/). Sequences were analysed using JOINSOLVER (http://join solver.niaid.nih.gov/) and IMGT-QUEST. Silent and replacement somatic mutations were identified by comparison with the germline gene sequence; in early experiments the ratio of replacement to silent mutations was considered to be evidence of affinity selection if significantly higher than 3:1. To correct for the inherent bias towards replacement mutations in CDRs, we have more recently applied the method of Hershberg to determine whether affinity selection has occurred in B-cell clones from ectopic germinal centres (Hershberg et al., 2008). This method employs an algorithm that allows for the effects of microsequences in the complementarity determining regions (CDRs) and the bias towards transition mutations. Clonally related sets of rearranged V-genes were identified by their use of the same germline V, (D) and J exons and shared junctional sequences. Genealogical trees were constructed by analysis of shared and unshared mutations using the parsimony method of phylogenetic analysis (PAUP, (Swofford, 1993)), enabling the assignment of sequences from parent and daughter cells that have been produced during clonal proliferation, thus providing clear evidence of the presence of clonally proliferating, somatically mutating B-cells within the germinal centre.

2.3 Cloning antigen-specific autoantibodies from germinal centre B-cells by ‘phage display’

In order to confirm the antigen specificity of B-cells generated in ectopic germinal centres, and to analyse in detail the relationship between their mutations and antigen specificity, we reconstituted the rearranged Ig V-genes as single chain Fv (scFv) or Fab antibodies by ‘phage display. Single chain Fv antibodies comprise the heavy and light chain variable region domains linked by a short peptide. Although linked together by a short additional peptide sequence, the V_H and V_L domains are able to fold into their natural 3-dimensional conformation and pair correctly, as the antibody produced by a B-cell or plasma cell. They contain the antigen binding site, and therefore mimic the antigen specificity of the original antibodies from which they were derived. A caveat that must be born in mind is that the original H and L chain pairings are unknown, except when both genes are cloned from a single cell. The detailed methodology has been described elsewhere (Stott & Sims, 2000; Matthews et al., 2002). Rearranged V_H and V_L-genes amplified either from microdissected germinal centres or pooled V-genes from the same B-cell clone, were used to construct scFvs using a (Gly4Ser)3 linker DNA. The resulting scFv library, comprising a pool of randomly linked V_H-V_L genes, was then inserted into the phagemids pCANTAB6 or pHEN2 continuously with the gene encoding the bacteriophage coat protein P3, and grown in E. coli in the presence of helper phage. The resulting scFv-P3 fusion protein was expressed on the surface of bacteriophage or as soluble scFv by transfection into a non-permissive, or permissive, strain of E. coli respectively. Alternatively, Fab libraries were constructed using whole light chain cDNA and DNA encoding the V_H region and the first constant region domain of the heavy chain by similar techniques (Matthews et al., 2002). An advantage of amplifying directly from genomic DNA is that the distribution of cloned V-genes reflects the usage of those genes by B-cells and plasma cells more accurately than amplification from cDNA, which is biased towards plasma cells. The library of scFvs or Fabs attached to bacteriophage by the P3 ‘phage coat protein was then panned on plastic plates coated with either a whole extract of the target tissue, or purified recombinant antigen, to identify self-reactive antibodies. Bound ‘phage were washed, eluted, re-grown in E. coli and panning
repeated until the eluate had become enriched with a small number of ‘phage clones. These were recloned and their H & L chains sequenced and used to investigate the specificity and properties of their antigen-binding sites.

2.4 Statistical analysis
The method of Hershberg et al. (2008), described in section 2.2 above, was used to determine the significance of replacement mutations in rearranged Ig V-genes cloned from germinal centre B-cells, as evidence for affinity maturation of the antibodies expressed by them. The distribution of V\textsubscript{H} gene families and individual V, D and J exons was assessed using two-tailed \( \chi^2 \) analysis, corrected for multiple comparisons.

3. The ectopic germinal centre response in myasthenia gravis

3.1 Pathology of myasthenia gravis
Myasthenia gravis is an organ-specific autoimmune disease characterised by weakness of striated muscles and thymic hyperplasia (Vincent, 2002). Patients are generally divided into subgroups with early-onset (EOMG, pre-40 years) or late onset (LOMG, post-40 years) forms of the disease, or with thymoma in about 10% of patients. It is a classic autoantibody-mediated autoimmune disease, caused by autoantibodies directed against the postsynaptic nicotinic acetylcholine receptor (AChR) at the neuromuscular junction. Many thymoma patients and some late onset patients also have serum antibodies against striated muscle antigens, interferon-\( \alpha \) and IL-12. Loss of functional AChRs leads to muscle weakness, usually first evident in weakness of eye movement. This can progress to other striated muscles of the body, causing problems with breathing due to effects on the diaphragm, swallowing difficulties and paralysis. These effects can be life-threatening if untreated. Evidence that the effects are mediated by autoantibodies against the AChR include induction of similar symptoms by: their transfer from mother to baby in utero; passive transfer from patients to mice; immunisation of animals with AChR; and marked improvement of symptoms in patients after removal of circulating IgG antibodies by plasmapheresis. Several pathogenic mechanisms are involved (Vincent, 2002; Drachman, 1994): (i) Cross-linking of the receptor by autoantibodies causes loss of AChR by antigenic modulation, leading to internalisation and degradation of the receptors; (ii) The majority of anti-AChR antibodies are of the IgG1 and IgG3 subclasses, which are particularly efficient at complement activation, resulting in lysis and damage to the muscle membrane; (iii) Less commonly, some antibodies cause direct inhibition of the ion channel function of the AChR; (iv) Antibody-dependent cell-mediated cytotoxicity has also been implicated, although there is little direct evidence for this mechanism. The IgG autoantibodies can cross the placenta of pregnant mothers with myasthenia gravis by an active transport mechanism involving the neonatal Fc receptor, FcRn, resulting in transient symptoms of myasthenia gravis in the newborn infant. The symptoms gradually ameliorate as the maternal antibodies are catabolised and replaced by the infant’s own antibodies. More rarely, the autoantibodies produced by multiparous mothers can induce severe, often fatal, developmental abnormalities, termed arthrogryposis multiplex congenita, due to paralysis of fetal muscles in utero (see section 3.4.5).

3.2 Structure and epitopes of the acetylcholine receptor
The AChR is a pentameric transmembrane glycoprotein found almost exclusively at the muscle endplate, comprising two \( \alpha \) polypeptide subunits, one \( \beta \), one \( \delta \) and, in the adult, one
ε subunit; in the fetus there is also one γ subunit, which is gradually replaced by an ε from the third trimester onwards (Fig. 2) (Vincent, 2002). The five subunits are combined into a cylindrical structure with a central cation channel that is closed in the inactive conformation. There are two binding sites for acetylcholine, formed at the interfaces between one α and δ subunit and the second α and ε or γ subunits. Electrical impulses passing down the motor nerve trigger release of acetylcholine molecules at the nerve termini. When these bind to the two receptor binding sites, they cause the central cation channel to open and sodium ions to flood into the muscle resulting in local membrane depolarisation. When this reaches threshold the resulting action potential spreads across the muscle triggering it to contract. Loss of at least 50% of receptors is required to produce overt muscle weakness.

Fig. 2. Diagrammatic representation of the structure of the acetylcholine receptor:
(a) the complete pentameric molecule in the cell membrane; (b) the topology of the subunits, illustrated for the α subunit that contributes to the acetylcholine/α-bungarotoxin binding site, the main immunogenic region (MIR) and the very immunogenic cytoplasmic epitope (VICE); It is doubtful whether the latter plays any significant role in pathogenesis; (c and d) the fetal and adult subtypes of AChR. Reproduced with permission from (Vincent et al., 1997), Plenum Press.

Since the patients’ autoantibodies are almost exclusively specific for the complex native conformation of the extracellular AChR subunit domains, and not short peptides or even whole subunit polypeptides, mapping of the autoantibody epitopes has proved to be difficult. The antibodies are mainly IgG1 or IgG3, of high avidity and heterogeneous in their sequences and fine specificity. Disease severity correlates poorly with autoantibody titre, suggesting that pathogenicity may depend upon precise epitope specificity. Many of the antibodies bind to a region of the extracellular domain of the α chain, the main immunogenic region or MIR. Its conformation is affected by the ε ↔ γ interchange, as
demonstrated by the observation that some antibodies bind better to the MIR of the fetal AChR than the adult form, even though the \( \gamma \) and \( \varepsilon \) subunits do not contribute directly to the MIR (Fostieri, Beeson, & Tzartos, 2000). Titres of MIR antibodies vary considerably between patients and other antibodies may play an equally important role in some individuals. Some patients also produce autoantibodies against the acetylcholine binding sites (that also bind \( \alpha \)-bungarotoxin); these are the blocking antibodies described above.

3.3 Role of the thymus

Early onset myasthenia gravis is associated with thymic hyperplasia characterized by secondary lymphoid organ-like structures in the medulla of >90% of patients. These include T-cell areas containing AChR-specific helper T-cells and large numbers of germinal centres with clearly defined mantle, dark and light zones. Plasmablasts and plasma cells secreting autoantibodies against AChR are also detectable within and around the germinal centres (Hill et al., 2008; SHIONO et al., 2003). Approximately 20% of germinal centres contain plasmablasts positive for antibodies against AChR and AChR is trapped on follicular dendritic cells in c.50% of thymic germinal centres (SHIONO et al., 2003). Anti-AChR-secreting hybridomas and AChR-specific Fabs have been cloned from thymic B-cells and thymectomy results in a reduction in the serum anti-AChR titre and reduced clinical symptoms in some patients, although the benefits of thymectomy have never been rigorously proved (Cardona et al., 1994; Farrar et al., 1997; Graus et al., 1997). The relative contribution of the thymus to production of anti-AChR autoantibodies compared with the secondary lymphoid organs is unknown, but it appears to play a significant role. Therefore, using the methods described in section 2, we tested the hypothesis that thymic germinal centres are sites of ongoing autoimmune responses driven by autoantigen, i.e. sites of activated B-cells, clonally proliferating, somatically mutating their expressed Ig V-genes and undergoing affinity maturation, driven by the acetylcholine receptor.

3.4 The thymic germinal centre response in myasthenia gravis

3.4.1 Germinal centres in the thymus

Thymi from 5 EOMG patients were examined by immunohistology. All 5 contained large numbers of germinal centres with typical mantle zones within the thymic medulla, histologically indistinguishable from germinal centres in human tonsil controls. The mantle zones contained densely packed CD20\(^+\) B-cells surrounding the germinal centre B-cells (Fig. 3A). These were interspersed with a network of follicular dendritic cells extending throughout the dark and light zones (Fig. 3C) and a crescent of T-cells can be seen at the apex of the light zone (Fig. 3B). Proliferating B-cells were distributed throughout the germinal centre but in larger numbers within the dark zone (Fig. 3D). Autoradiography with \(^{125}\)I-\( \alpha \)-bungarotoxin alone, which binds to AChR, diffusely labelled c.50% of germinal centres and appeared to be associated with the follicular dendritic cell processes. No labelling was seen in human tonsils or thymi from two seronegative myasthenia patients and bungarotoxin binding was blocked by the cholinergic drug, carbamyl choline, which is structurally similar to AChR, indicating that the follicular dendritic network contained membrane-bound antigen or immune complexes.

In contrast, \(^{125}\)I-\( \alpha \)-bungarotoxin-labelled AChR bound to individual cells in 20% of germinal centres, including large numbers of moderately labelled centrocytes in the light zone, smaller numbers in the dark zone, and heavily labelled plasmablasts/plasma cells in and around the germinal centres (Fig. 3E & F).
Fig. 3. Immunohistochemically stained serial sections through thymic germinal centres.

A & B: Germinal centres stained (red) with anti-CD20 and anti-CD3 for B and T-cells respectively. The arrow (B) shows a crescent of T-cells in the light zone. C: A network of follicular dendritic cell processes is spread throughout the germinal centre, including both the light and dark zones. D: Germinal centre cells stained with an antibody against proliferating cell nuclear antigen, revealing dividing B-cells in both areas but more concentrated in the dark zone. E & F: \(^{125}\text{I}-\alpha\)-bungarotoxin-labelled AChR reveals individual AChR-specific plasmablasts/plasma cells in and around germinal centres (detected by autoradiography). Diffuse labelling in the light zone probably indicates binding of free \(^{125}\text{I}-\alpha\)-bungarotoxin to AChR trapped on follicular dendritic cells (see text). Reproduced from Sims et al (2001).

3.4.2 Ig V-gene expression by thymic germinal centre B-cells

In order to determine whether thymic germinal centre B-cells are subjected to antigen-driven clonal proliferation, somatic hypermutation and affinity selection, as seen in the orthotopic germinal centres of secondary lymphoid organs, we cloned and sequenced rearranged Ig heavy chain V-genes from multiple sections through four thymic germinal centres (A – D) and the follicular mantle surrounding one of them (A). 216 rearranged \(V_{\text{H}}\)-genes, derived from 61 independently rearranged, functional sequences, were obtained from the four germinal centres and 46 \(V_{\text{H}}\)-genes from the follicular mantle were derived from 24 functional \(V_{\text{H}}\)-genes. Since the PCR error rate in control experiments was estimated to be less than one base per four \(V_{\text{H}}\)-genes, only sequences using the same V, D & J exons, the same junctional sequences and a minimum of one base difference per gene, were accepted as mutated members of a clonally related set of B-cells. This conservative assessment almost certainly discards some B-cell clones with low frequencies of somatic mutation and therefore underestimates the true B-cell diversity. When calculating the number of \(V_{\text{H}}\)-genes used, members of the same B-cell clone were counted only once. However, this reflects the number of individual B-cell clones and non-dividing B-cells rather than the total number of B-cells using a particular gene.
The distribution of \( V_H \)-gene families was similar in the follicular mantle and all four germinal centres, with predominant use of members of the \( V_H3 \) gene family compared with the number of germline \( V_H \)-genes in this family (Fig. 4A). However, since this gene family is also predominantly used by the peripheral blood B-cells of healthy individuals, it was not considered to be significant. No \( V_H6 \), \( V_H7 \) or \( J_H2 \) genes were isolated and \( J_H1 \) was under-used, but the rarely used \( V_H5-51 \) gene and the \( J_H4 \) exon (Fig. 4B) were over-represented, both being used by rearranged V-genes from three different germinal centres in combination with different D exons (\( D_H2-2 \), \( D_H5-12 \) and \( D_H6-19 \)). In many cases the D exon could not be identified due to junctional diversity and removal of bases during recombination. Of those that could be identified, \( D_H3 \) and \( D_H6 \) were the most commonly used. These data strongly imply selection for B-cells expressing antigen receptors using particular combinations of V, D and J segments, despite the heterogeneity of the germinal centre B-cells. This would be even more apparent if individual members of the same B-cell clone were counted separately.

![Fig. 4. Frequency of usage of \( V_H \) and \( J_H \) germline genes by thymic germinal centre B-cells.](image)

**A** The frequency of \( V_H \) gene family usage in all germinal centres analysed differs significantly from predictions from the number of members of each family in the germline. Members of clonally related sets were only counted once. **B** The frequency of individual \( J_H \) exon usage in all germinal centres analysed differs significantly from the number of \( J_H \) exons in the germline (*p<0.01*). Reproduced from Sims *et al.* (2001).

### 3.4.3 Somatic hypermutation and clonal proliferation of B-cells in thymic germinal centres

All the germinal centres and the follicular mantle contained B-cells expressing both mutated and unmutated Ig V-genes, the latter presumably coming from naïve B-cells. The majority of V-genes from the germinal centres and the follicular mantle were mutated, with considerable variation in the number of mutations, ranging from 0 – 52. Some of the rearranged V-genes in clonally related sets had high ratios of replacement to silent mutations from 4.7:1 to 7.0:1 in the CDRs, which form the antigen-binding site, suggesting affinity selection of mutated antigen receptors is taking place in the germinal centre. Some other sequences had low numbers of replacement mutations, suggesting selection against replacement mutations, as also found by (Zuckerman *et al.*, 2010). 18 different sets of functionally rearranged \( V_H \)-genes included two or more related sequences sharing the same V, D and J segments and junctional sequences but with significantly more than one mutation per V-gene and, in three cases, they were cloned in separate amplifications from different sections through the same germinal centre and therefore could only be from different B-cells.
Fig. 5. Examples of clonally related sets of rearranged $V_H$ genes isolated from thymic germinal centres C and D.

The genealogical trees were constructed using the most parsimonious parent-daughter cell relationships of clonally related sets of sequences (section 2.2). The best matching germline gene is depicted as an ellipse. Letters in the circles refer to individual sequences from the same clonally related set. Deduced intermediates are shown as dotted circles. Numbers at the side of the arrows indicate the minimum number of mutations required to generate a daughter cell from the immediate parental cell. The results show that the B-cells from which they were derived are undergoing antigen-driven clonal proliferation and somatic hypermutation from both naïve and memory B-cell precursors. A: B-cell clones from germinal centre C; B: B-cell clones from germinal centre D; C: Frequency of mutations in the $V_H$ genes expressed by the most probable progenitor cells of all 18 B-cell clones analysed. Reproduced from Sims et al. (2001).
These related sets of V-genes were therefore derived from members of the same proliferating B-cell clone, showing that antigen-driven B-cell proliferation and somatic hypermutation are taking place within the thymic germinal centres. All of the isolated B-cell clones were small, containing a maximum of five members.

We do not know the total number of clones proliferating within a single germinal centre. We examined every tenth section but may have missed small B-cell clones, so there are likely to be more than we detected. Genealogical trees of related sets of V-genes were constructed by the most parsimonious relationship on the basis of shared and unshared mutations (section 2.5). Six of the 18 clones isolated from two germinal centres containing AChR-specific B-cells were derived from unmutated precursors, i.e. naïve B-cells, whereas the V\textsubscript{H}-genes of the earliest founder cells isolated from the other 12 clones contained from 5 to >30 mutations. Examples of six of these clones are shown in Fig. 5. Although we cannot rule out the possibility that some of these clones were also derived from unidentified naïve B-cells, it is most probable that the majority were generated from founder memory B-cells. Thus, both memory and naïve B-cells have been stimulated by antigen and are proliferating and mutating their antigen receptors within thymic germinal centres of myasthenia patients.

### 3.4.4 Evidence for selection of AChR-specific B-cell clones in thymic germinal centres

Three B-cell clones from 3 different AChR positive germinal centres expressed the rarely used V\textsubscript{H}5-51 genes. Furthermore, these three independent sets of V-genes exhibited the same amino acid replacements at three positions. Comparison of our V-gene sequences with those of heavy chains from known AChR-specific hybridomas and Fabs revealed some common features. Germline genes used by four of our B-cell clones also encode anti-AChR antibodies cloned from other EOMG patients, and several of the amino acid substitutions in CDR1 and CDR2 from two B-cell clones were also present in an anti-AChR Fab (Sims \textit{et al.}, 2001; Matthews \textit{et al.}, 2002). It is therefore unlikely that these mutations occurred by chance, suggesting that a common selection process for mutants with high affinity for the AChR is in operation.

### 3.4.5 Evidence for immunisation by the fetal form of acetylcholine receptor

Babies born to mothers with myasthenia gravis can develop transient symptoms due to transplacental transfer of the maternal autoantibodies and, in rare cases, they have severe developmental abnormalities, arthrogryposis multiplex congenita (AMC), caused by maternal anti-AChR antibodies that inhibit the ion channel function of the fetal AChR, causing paralysis during fetal development \textit{in utero}, whereas the adult form of the mother’s AChR is relatively unaffected. Fetal AChR-specific antibodies are particularly prevalent in women who have had babies, suggesting that they may be induced by the fetus.

In order to determine the specificity and clonal origins of fetal AChR-specific autoantibodies, combinatorial Fab libraries were constructed from cDNA prepared from thymic cells of two mothers (M2 and M6) of AMC babies. 25 Fab clones were isolated and two clonally related sets were examined in greater detail. All Fabs bound specifically to the γ subunit of fetal AChR, except one that recognised the β subunit also present in the adult receptor. Sequencing of the fetal-specific Fabs revealed clearly restricted usage of V\textsubscript{H}, J\textsubscript{H}, V\textsubscript{K} & J\textsubscript{K} gene segments and convergent coding mutations. All the Fabs from AMC mother M2
used the \( V_{H3-07} \) gene recombined with \( J_{H6b} \) and an unidentified D exon in combination with various \( V_{k} \) genes, suggesting that the heavy chain is the major contributor to AChR binding, at least in this case. The \( V_{H3-07} \) segments were mutated and clonally related. Most of the Fabs from AMC mother M6 used the same combination of mutated \( V_{H3-21} \) and \( J_{H5b} \), with an unidentified D exon, plus a \( V_{k02-12/J_{k4}} \) light chain, which was also used in two of the Fabs from M2. In this case, both the \( V_{H3-21} \) and \( V_{k02-12} \) sets of sequences were clonally related, suggesting that they may both be derived from the same B-cell clone. The clonally related sequences from both sets of Fabs from M2 and M6 contained many shared and unshared coding mutations. The apparent founder member of each set of sequences had a large number of base differences from the best matching germline V-gene, suggesting that the clones were derived from mutated memory B-cells (Fig. 6).

Fig. 6. Clonally related sets of rearranged \( V_{H} \& V_{k} \) genes encoding AChR-binding Fabs cloned from thymic cells from AMC mother M6.

The heavy and light chains were from the same set of Fabs. Genealogical trees were constructed and mutations numbered as described in the legend to figure 5. Fab names are shown in the circles, H referring to the heavy chain and K referring to the \( \kappa \) light chain. Dotted circles represent hypothetical intermediates. Numbers at the side of arrows show the minimum number of mutations required to generate a daughter cell from its immediate parent. Reproduced from Matthews et al. (2002).

Several sequences from both clonally related sets of Fabs had many more replacement mutations than expected by chance, indicating affinity selection. There was also clear evidence of convergent mutations. Many independently rearranged sequences from both patients shared consensus mutations. All the Fabs using \( V_{H3} \) genes contained the same 31S\( \rightarrow \)T mutation and most of the \( V_{k02-12} \) genes contained a 22SRASET28 motif in CDR1. A search of the GenBank database of all human Ig sequences found only two other \( \kappa \) chains containing this motif, suggesting that it is important in determining the specificity of the anti-AChR autoantibodies.
3.5 Conclusions

Since the EOMG thymus contains many typical germinal centres surrounded by mantle zones, including clones of proliferating, AChR-specific B-cells, it is clearly a site of autoantibody diversification. The B-cells are undergoing somatic hypermutation and affinity selection by cognate binding to AChR bound to the membrane processes of follicular dendritic cells, which form a network surrounding B and T-cells in the germinal centres. B-cells expressing high affinity, AChR-specific antigen receptors receive rescue signals from follicular dendritic and helper T-cells inducing them to differentiate into antibody-secreting plasmablasts and plasma cells that migrate out of the follicles and leave the thymus to enter the circulation, in a classical germinal centre type response. The numerous mutations seen in some V-gene sequences suggests that Ig class-switched memory B-cells are also generated and either leave the thymus, or some may recirculate within the germinal centre, undergoing further rounds of somatic hypermutation and affinity selection, consistent with the observations of others that recirculation of centrocytes between the light and dark zones occurs in orthotopic germinal centres. Several of the B-cell clones we identified were derived from highly mutated precursors, which supports this hypothesis. The autoantibodies produced by mothers of AMC babies included antibodies specific for the fetal form of the AChR, directed against the γ subunit.

The reason why large numbers of germinal centres producing AChR-specific B-cells and plasma cells develop in the thymi of myasthenia gravis patients is unclear, but it is possible that the rare thymic myoid cells may be involved. The induction of fetal AChR-specific antibodies in parous women suggests, at least in some cases, that expression of the fetal AChR on thymic epithelial and myoid cells may initiate an immune response, for which the patient’s immune system has not been tolerised. Furthermore, generation of thymic germinal centres may be induced by a pre-existing proinflammatory cytokine environment, including IFNγ and TNFα. These molecules have been shown to upregulate expression of AChR subunits in thymic epithelial cells and on the membranes of myoid cells (Poea-Guyon et al., 2005). The chemokines CXCL13 AND CCL21 are produced by endothelial cells of the afferent lymphatic vessels in the thymus, attracting activated T and B-cells, including naïve B-cells, suggesting that this is the mechanism of induction of thymic germinal centres (Berrih-Aknin et al., 2009; Le Panse et al., 2006; Meraouna et al., 2006; Le Panse et al., 2010).

We therefore propose a two step process for initiation of the autoimmune response in myasthenia gravis (SHIONO et al., 2003). In step 1, there is hyperplasia of thymic epithelial cells expressing linear AChR epitopes, including the α and ε subunits, in the context of the MHC Class II antigen HLA-DR52a, a susceptibility factor for EOMG patients. Whereas these peptides would normally induce tolerance, an imbalance in regulatory factors and expression of costimulatory molecules results in activation of thymic T_{h} cells against AChR epitopes. In step 2, the T_{h} cells induce an early B-cell response against the linear peptides and some of the resulting antibodies cross-react with conformational epitopes of the native AChR expressed on the thymic myoid cells, leading to myoid cell damage, release of AChR/antibody immune complexes, danger signals, and recruitment of professional antigen presenting cells. These stimulate an enhanced B-cell response accompanied by formation of germinal centres with production of high affinity, class switched, pathogenic autoantibodies. Although some aspects of this hypothesis are conjectural, they are also testable.
4. Tissue-infiltrating B-cells in inflammatory myopathies

4.1 Introduction

The inflammatory myopathies (IM), collectively called myositis, are classified into three principal subsets, Dermatomyositis (DM), Polymyositis (PM) and Inclusion Body Myositis (IBM) (Bohan & Peter, 1975a; Bohan & Peter, 1975b; Dalakas & Hohlfeld, 2003). Each of these disorders is characterised by moderate to severe muscle weakness and muscle fatigue with inflammatory mononuclear cell infiltration within the muscle, but each disorder has distinct clinical and pathological features. IM can be associated with various autoimmune and connective tissue disorders as well as malignancies, the latter being associated with up to 45% of adult DM patients.

DM, the most common of the inflammatory myopathies, is a multi-organ disease not only affecting skeletal muscle but, often, the skin as well as other tissues and is more commonly found in women than men; it also accounts for up to c.85% of all juvenile IM (Rider, 2007). DM is characterised by a heliotrope rash on the upper eyelid, face or upper trunk accompanying, or more commonly preceding, proximal muscle weakness. Muscle inflammation is predominantly perivascular and/or perimysial or in the interfascicular septae and around, rather than within, the muscle fascicles. Perivascular atrophy is a characteristic feature of DM patients, often in groups at the periphery of the fascicle. In DM, muscle lymphocytic infiltrates consist largely of B-cells and CD4+ T-cells (Arahata & Engel, 1984; Engel & Arahata, 1984) suggesting that DM may be a humorally mediated immune response.

PM and IBM, though separate disorders, are both characterised by scattered necrotic and regenerating muscle fibres and endomysial inflammation with invasion and destruction of non-necrotic muscle fibres. PM generally becomes evident in adulthood and is best defined as a subacute myopathy that evolves over weeks to months and presents with symmetrical weakness of the proximal muscles. Its clinical onset is hard to define with no early recognition signs such as the rash observed in DM. PM is uncommon as a stand-alone disorder and more commonly associates with other autoimmune and connective tissue disorders.

Onset of IBM is usually after the age of 50 and occurs more frequently in men. Muscle weakness can be both proximal and distal and is often asymmetrical. Despite similarities with PM, distinctive features of IBM include: rimmed vacuoles; groups of atrophic fibres; increased lymphocytic invasion of non-necrotic fibres; less frequent myofibre necrosis; and a more slowly progressing clinical course with patients being unresponsive to treatment. In both disorders, inflammatory infiltrates typically consist of CD8+ T-cells and macrophages (Arahata & Engel, 1984; Engel & Arahata, 1984) which invade MHC Class 1 antigen-expressing muscle fibres, a feature absent in normal muscle tissue, leading to fibre necrosis. The muscle fibre invading CD8+ T-cells can be clonally expanded in both PM and IBM (Dalakas, 2004; Fyhr et al., 1997; Hofbauer et al., 2003; Mantegazza et al., 1993; Seitz et al., 2006), which persists over time (Amemiya, Granger, & Dalakas, 2000).

4.1.1 Autoantibodies associated with myositis

As with most autoimmune disorders, different autoantibody specificities have been described in DM and PM; autoantibodies are generally absent from IBM although they have been detected in a small number of cases (Dalakas et al., 1997). They can either be myositis-specific (MSAs) or myositis-associated autoantibodies (MAAs), which can also be associated
with other autoimmune diseases. Most bind to protein or ribonucleoprotein complexes involved in protein synthesis, translocation or elongation; MAA target antigens are primarily located in the nucleoplasm or nucleolus. The most prevalent MSAs are directed against amino-acyl-tRNA-synthetases (ARS), and are associated with a distinctive clinical phenotype, anti-synthetase syndrome, characterised by myositis, Raynaud’s phenomenon and interstitial lung disease, with a higher mortality. Anti-Jo-1 (anti-histidyl-tRNA synthetase) antibodies are the most prevalent in myositis patients (20-30% of patients), while the other anti-ARS antibodies are only present in 1-3% of IM patients, and are a diagnostic and prognostic marker for disease severity (Mielnik et al., 2006; Zampieri et al., 2005).

4.1.2 Muscle-infiltrating B-cells in myositis
As described above B-cells have been found to be prominent within the muscle infiltrating cell populations of DM patients and are rarely found, or absent, in the inflamed muscle of PM and IBM patients. CD138+ plasma cells have been identified within the infiltrating populations, predominantly in the endomysial areas of muscle tissue of PM and IBM patients (Greenberg et al., 2002; Greenberg et al., 2005). This was confirmed by sequence analysis of immunoglobulin V-genes expressed by laser dissected cells as well as microarray studies which showed an abundance of immunoglobulin transcripts.

The role for B-cells and plasma cells in IM is still currently unresolved, with continuing studies providing further insight into the immune mechanisms. The identification of muscle infiltrating B-cells, plasma cells and autoantibodies suggests that these diseases may be at least partly driven by a loss of B-cell tolerance and, in the case of PM and IBM patients, not solely by the oligoclonal expansion of T-cells. We therefore investigated whether there is clonal expansion of infiltrating, autoantibody producing B-cells in situ in IM.

4.2 The muscle infiltrating B-cell response in myositis
4.2.1 The cellular composition of infiltrating lymphoid cells in myositis
To determine whether specific, antigen-driven, B-cell adaptive immune responses were occurring in situ, we used the methods described in section 2 to study the cellular composition of muscle infiltrating cells in twelve different muscle samples (2 DM, 9 PM, 1 IBM); we also examined their Ig V-gene repertoire and the processes of somatic hypermutation and clonal diversification of the rearranged V-genes. In contrast with other autoimmune diseases (see above), no classical ectopic germinal centre structures were observed within the inflamed muscle; instead, muscle-infiltrating cells were present in cellular aggregations which varied from loose to dense in the appropriate perivascular/perimysial or endomysial locations, as in previous studies. B-cells were a significant component of the inflammatory infiltrate in all samples examined for all three myositis subsets, either as CD20+ B-cells or differentiated plasma cells (Figure 7A-D), although the most significant infiltration of CD20+ B-cells was observed within the muscles of the two DM patients. FDCs were rare, and were seen only in one IBM and three PM samples, and not at all in DM. In addition to these cell phenotypes, CD3+, CD4+, CD8+, CD68+ and FoxP3+ cells were also present. Double immunofluorescence staining of cell phenotypes with the proliferating cell marker Ki67 identified proliferating cells within the infiltrating population. In addition to CD20+ B-cells (Figure 7E & F), proliferating CD3+, CD4+, CD8+ and CD68+ cells were observed, as well as FoxP3+ cells in one DM patient.
4.2.2 The Ig V-gene repertoire and clonally proliferating, muscle-infiltrating B-cells

Analysis of the repertoire of rearranged Ig V-genes expressed by infiltrating B-cells and plasma cells revealed significant biases for and against individual gene segments and families relative to the normal peripheral blood B-cell and the germline gene repertoires. V-gene usage varied between patients and myositis subsets and, in a few instances, differed significantly between the DM and PM subsets. Interestingly, naïve or unmutated B-cells (0-2 mutations per V_H gene) constituted almost 50% of the B-cells in DM, but <10% in PM, where a large proportion of sequences was highly mutated (c.30% >20 mutations). As expected, mutations were prevalent within CDRs 1 & 2. A total of nine clonally related sequences was found in five of the IM patients studied; 2 DM and 3 PM patients, each with up to four different clones comprising between two and ten clonal variants (Figure 8). These clonally related sequences provide evidence for specific, antigen-driven B-cell immune responses within the inflamed muscle. However, using the method of Hershberg et al. (2008), we found no evidence of positive selection in the CDRs of clonally related sequences, nor in any sequences isolated from the DM patients, and only in a small percentage from the PM patients. Finally, using biotinylated recombinant antigens, we identified antigen-specific B and plasma cells in infiltrates from the five out of twelve patients whose autoantibody specificities were known, including Jo-1 (Figure 9).

Parallel studies (Bradshaw et al., 2007) also demonstrated B-cell responses in muscle of 3 DM, 2 PM and 7 IBM patients but very few CD19^+ or CD20^+ cells were observed, the
The majority of B-lineage cells being CD138+ plasma cells that had class switched to either IgG or IgA. Clonally related sequences were isolated from whole muscle sections from ten of the twelve myositis patients, with up to four different clonal sets isolated from each muscle sample. Further studies also support the absence of classical ectopic germinal centre structures and the clonal expansion and maturation of B-cells within inflamed muscle (Salajegheh et al., 2010). Collectively this and our work strongly suggest the participation of antigen-specific B-cell immune responses within the muscle.

![Fig. 8. Oligoclonal expansion of B-cells and plasma cells in inflammatory myopathies](image)

Representative examples of clonal genealogical trees constructed from sequences isolated from muscle-infiltrating B-cells and plasma cells in individual patients, representing the minimum number of cell divisions required to generate each daughter cell. Clone A from a DM patient; clone B from a PM patient. The letters in the circles refer to individual sequences isolated from each B-cell clone. Genealogical trees were constructed and mutations numbered as described in the legend to Figure 5. Bracketed figures representing additional silent mutations. Dashed circles represent hypothetical intermediates whose sequences were not isolated from the muscle-infiltrating population.

![CD20 Jo1 and Plasma Cell Jo1](image)

CD20+ B-cells and antigen-specific cells were visualised by Fluorescein-Avidin D (green) and Texas Red-Avidin D (red) respectively. Original magnification was 630X. Arrows identify antigen-specific cells within the muscle-infiltrating population of polymyositis patients.
4.3 Conclusions
As previously described in other autoimmune diseases, a role for B-cells in IM is implied by
the clinical improvement in patients administered Rituximab® therapy, including
improvements in muscle strength. Some patients relapsed as their B-cell pools repopulated
and depletion of autoantibody titres was variable (Chung, Genovese, & Fiorentino, 2007;
Cooper et al., 2007; Levine, 2005). The potential of B-cells as therapeutic targets is further
supported by the elevations in serum levels and gene expression of B-cell activating factor
(BAFF) in IM patients, a cytokine crucial for B-cell maturation and survival, which is also
thought to play a role in autoantibody production (Krystufkova et al., 2008; Salajegheh et al.,
2010).
Despite all the evidence described here implicating B-cells and loss of B-cell tolerance in the
IM, numerous questions still remain to be resolved, including identification of the
stimulating antigens and epitopes, sequence characteristics and pathogenicity of high-
affinity, antigen-specific antibodies produced in situ, and the factors regulating and
controlling these autoimmune reactions. The resolution of these questions will enhance our
understanding of the immune pathology of IM and facilitate the diagnosis, treatment and
management of these diseases.

5. The ectopic germinal centre response in breast cancer
In autoimmune diseases there is a failure of immunological tolerance resulting in an
immune response to self-antigens, causing pathological damage to the target organ and
tissues, the nature of the target tissue depending on the specificity of the response. In
malignancy the immune response, if it occurs, is similar in that it is essentially directed
against self, i.e. tumour-associated, antigens. These antigens may be mutated, altered by
metabolic processes, or merely aberrantly or over-expressed on the tumour. The problem,
however, is converse to autoimmune disease in that, whereas in autoimmune disease the
aim is to suppress the immune response, preferably specifically against the target organ, in
cancer the hope is that it will be possible to boost the immune response which is often too
weak to overcome the rapidly growing tumour.
Several autoimmune disorders sometimes associate with certain tumours, most often small
cell lung cancer, breast or ovarian carcinomas, ovarian teratomas, neuroblastomas and
lymphomas (with Sjögren’s syndrome), reviewed by Lang & Vincent and Rosenfeld et al.
(Lang & Vincent, 2009; Rosenfeld & Dalmau, 2010). In the examples studied, the target
autoantigen(s) are expressed on the tumour which seems to autoimmunise against them.
Indeed, if the tumour is removed, autoantibody levels often decline (Chalk et al., 1990). In
many syndromes, the autoimmune disorder serves as a valuable early warning of the
associated tumour, and may even slow its growth (Maddison & Lang, 2008).

5.1 Pathology of breast cancer
Breast cancer is the second most common malignancy in women, accounting for 31% of all
types of cancer, with a lifetime incidence in the U.K. of 1/8 in women and c.1/1000 in men.
Despite advances in screening, diagnosis and therapy, 12,000 women die of breast cancer
each year in the U.K. and the global incidence in females is 23%, but there are marked
variations between different regions, it being the highest in Western Europe, Australia, New
Zealand and North America. The incidence is relatively low in Asian and African countries
(figures from Cancer Research UK). There are several different histopathological types of
breast cancer, of which the major types are the ductal and lobular carcinomas, either of which can be \textit{in situ} or invasive, the \textit{in situ} type being considered a possible precursor of invasive carcinoma. Ductal and lobular carcinomas \textit{in situ} are confined to the mammary ducts and lobules and have a very high cure rate, approaching 100%. Invasive carcinomas account for the majority of breast cancers and have a much poorer prognosis. Malignant cell growth appears to start in the ducts and lobules and then invades the surrounding tissues and ultimately metastasises to other tissues and organs. A less common type is medullary carcinoma, comprising only $c.1 - 5\%$ of breast cancers; this typically has heavy infiltrates of B-lymphocytes and a significantly better prognosis than the invasive ductal and lobular types. Length of disease free survival in breast cancer is unpredictable, with relapse occurring up to ten years post treatment and even beyond; it has been postulated that this may be due to host factors, including the nature and extent of the immune response.

5.2 The immune response to breast cancer

Most breast cancers contain infiltrates of lymphoid cells with large numbers of T-cells, including CD4$^{+}$ and CD8$^{+}$ T-cells, and variable numbers of B-cells, natural killer cells and macrophages. The degree of infiltration varies between different types of breast cancer with extensive lymphoid cell infiltrates in ductal carcinoma \textit{in situ} and some invasive ductal and lobular carcinomas (Ben Hur \textit{et al.}, 2002). Most studies have focused on the role of cytotoxic T-cells in tumour immunity, with variable success in attempting to suppress tumour growth by boosting the T-cell response to tumour-associated antigens. Relatively few studies have addressed the role of B-cells and humoral immunity in response to cancers, including breast cancer, despite the observation that $c.40\%$ of ductal breast carcinomas have significant B-cell infiltration.

There is increasing evidence that B-cells play important dual opposing roles in the immune response to tumours; on the one hand as antigen presenting cells and producers of cytotoxic antibodies effective at killing tumour cells by antibody-dependent cell-mediated cytotoxicity (ADCC) and complement-dependent cell lysis, and as tumour antigen-presenting cells capable of very efficient T-cell activation; on the other hand as promoters of inflammation aiding tumour progression (de Visser, Korets, & Coussens, 2005; de Visser, Eichten, & Coussens, 2006). These seemingly contradictory effects may be due to the difference between a specific, high affinity immune response to antigen versus a low affinity, polyclonal response, or even suppression of the cytotoxic immune response via regulatory B-cells (Mauri, 2010). The importance of antibodies in eliminating tumours is clearly demonstrated by the results of treatment of breast cancer patients with humanised monoclonal antibodies (MAbs) specific for the epidermal growth factor receptor HER-2 (trastuzumab/herceptin and pertuzumab). Not only is herceptin effective in slowing down the progression of established metastatic disease, it has also recently been demonstrated to prevent the emergence of metastases when given as an adjuvant treatment (Hortobagyi, 2005). Pertuzumab has also yielded promising results in clinical trials (Bianco, 2004). Synergistic effects between herceptin and pertuzumab suggest promising new approaches to therapy using cocktails of antibodies (Nahta, Hung, & Esteva, 2004) and elucidation of the molecular structure of the herceptin Fab/HER-2 complex (Cho \textit{et al.}, 2003) allows rational design of therapeutic anti-HER-2 antibodies. MAbs specific for other tumour-associated antigens (TAAs) are needed to work synergistically with trastuzumab and to treat patients who do not overexpress HER-2.
Several molecules have been identified that are either over-expressed, mutated, or structurally modified on tumour cells and are therefore potential targets for immunotherapy, including HER-1, HER-2, MUC-1 and p53 (Taylor-Papadimitriou et al., 2002). Some TAAs appear to overcome tolerance and induce a natural immune response as a result of mutation or altered expression; humoral immune responses to these antigens in breast cancer patients are associated with better early disease stage-specific survival (Angelopoulou et al., 2000; Visco et al., 2000; von Mensdorff-Pouilly et al., 2000) and anti-MUC-1 antibodies are cytotoxic to tumour cells (Snijdewint et al., 2001). TAA-specific tumour infiltrating (TIL) B-lymphocytes and recombinant antibodies have been isolated from both tumour (Kotlan et al., 2000; Kotlan et al., 2005) and lymph nodes (Petrarca et al., 1999; Rothe et al., 2004) of medullary and ductal carcinoma patients, showing that they are responding specifically to the tumour. Evasion of the immune response by the tumour can be overcome by passive immunotherapy or active immunisation regimes. B-cells actively responding in the draining lymph node and tumour are therefore ideal sources to study the immune response to the tumour and provide the most relevant source of potentially therapeutic antibodies.

5.3 Ductal carcinoma infiltrating lymphocytes are clustered into germinal centres
We and others found infiltrating lymphocytes in ductal carcinomas were aggregated into clusters containing T-cells, B-cells and follicular dendritic cells with plasmablasts/plasma cells in and around the aggregates (Coronella et al., 2002; Nzula, Going, & Stott, 2003a). These cell clusters appeared to be similar to those seen in the target tissues of autoimmune diseases except that there was no mantle zone (also absent in the salivary glands of patients with Sjögren’s syndrome (Stott et al., 1998)), so we examined whether they were responding as germinal centres.

5.4 The Ig V-gene repertoire and clonal proliferation of B-cells in ductal carcinoma
We cloned and sequenced 401 rearranged Ig V_{Ht}-genes from microdissected tumour-infiltrating B-cell foci of 7 patients with invasive ductal carcinoma and 271 V_{Ht}-genes from paired sentinel lymph nodes of 3 of the patients. 15 sets of V_{Ht}-genes from clonally related B-cells within individual foci were identified by their shared V_{Ht}, D, J_{Ht} and CDR3 sequences, showing that proliferating, mutating B-cell clones were present in lymphoid foci and that these foci were undergoing a germinal centre response within the tumour, similar to the ectopic germinal centres we have observed in the target tissues of autoimmune diseases (Fig. 10). There was preferential usage of certain V_{Ht}, D & J_{Ht} exons, indicating selection of B-cells expressing antigen receptors encoded by these gene combinations. V_{Ht} & V_{Lt}-genes from proliferating B-cell clones contained numerous mutations, demonstrating that the somatic hypermutation machinery was switched on within the cell cluster, again characteristic of a germinal centre response. Analysis of the pattern of mutations showed that the B-cell clones are undergoing an antigen-driven response accompanied by selection of specific mutations and affinity maturation in situ. Clone founder cells were of both naïve and memory B-cell type, showing that a secondary response involving memory B-cells was taking place, but also new B-cells that had not previously encountered antigen moved into cell clusters and were stimulated by antigen. We also cloned rearranged V_{Ht}-genes from microdissected germinal centres in the paired sentinel lymph node and identified proliferating, hypermutating B-cell clones there too. These also revealed selection for particular V_{Ht} & J_{Ht}-genes showing selection by antigen for B-cells expressing these genes during the immune response but we did not find evidence that members of the same B-cell clones had migrated.
from the sentinel node to the tumour; this could have been due to insufficient sample sizes (Nzula, Going & Stott, 2003a; Simsa et al., 2005).

The best matching germline V_{H}-gene is shown at the origin of each tree. The founder rearranged V_{H}-genes of three of these B-cell clones already appear to be mutated, implying that they originated from memory B-cells, whereas clone A1 appears to have been founded by a naïve B-cell. Genealogical trees were constructed and mutations numbered as described in the legend to Figure 5. Reproduced from Nzula et al. (2003).

5.5 Cloning and characterisation of scFv antibodies
In order to identify the antigens driving the immune response within the tumour, we reconstructed the antigen receptors expressed by germinal centre B-cells as scFv antibodies.
by cloning $V_H$ and $V_L$-genes into phagemid (pHEN2), as described in section 2.3, to generate scFv-phage libraries expressing randomly assorted combinations of $V_H$ and $V_L$-genes. These “mini-libraries” were made from the B-cells in individual germinal centres within the tumour and are therefore much smaller and more restricted than the very large libraries normally made by random combination of large pools of $V_H$ and $V_L$-genes. Two scFv-phage mini-libraries were constructed from a germinal centre incorporating all the $V_H$-genes pooled from the largest proliferating B-cell clone (D4 in Fig. 10) linked randomly to either the rearranged $V_NJ$-genes or the rearranged $V_{nj}$-genes amplified from the same germinal centre.

Tumour-binding scFv were selected from the mini-libraries by three or four cycles of panning and elution on a heterologous tumour homogenate pooled from breast tumours of 5 patients. During the panning cycles we observed exponential enrichment of the $V_{nj}$ mini-library, but not the $V_NJ$ mini-library, indicating that scFv within the $V_{nj}$ mini-library bind specifically to tumour-associated antigens (Fig. 11A). This is consistent with the scFv-libraries being derived from the same B-cell clone, since a single B-cell clone uses either a $\kappa$ or $\lambda$ light chain, not both. 13 scFv-phages binding to the tumour extract were cloned and their specificity for tumour tissue confirmed by ELISA. 7 scFv-phage clones that bound to the tumour extract were identified for further characterisation. All 7 used the same combination of $V_H3-23$ with exons D1-26 & $J_H2$, expressed by B-cell clone D4, and the light chain gene $V_{nj}1c$ with $J_{nj}3b$.

We also constructed two scFv-phage libraries from DNA extracted from a whole sample of tumour tissue, as described for the mini-libraries. The 2 libraries were panned on the same heterologous tumour homogenate as used with the mini-libraries. After 4 cycles of panning we observed an enrichment of several logs for both libraries, indicating the presence of tumour-specific antibodies (Fig. 11B). The enrichment of both global libraries shows that tumour-specific B-cells derived from independent B-cell clones were present in the tumour, as expected. 19 scFv-phages were cloned from the 2 global libraries and their specificity for tumour tissue confirmed by ELISA using the same tumour homogenate as source of antigen.

### 5.6 Identification of the specificity of proliferating B-cells

Since the scFvs from the $V_{H}/V_{\lambda}$ mini-library were derived from proliferating B-cell clone D4, their sequences and antigen specificities reveal the nature of the genes and antigen receptor specificities of the original germinal centre B-cells. We therefore sequenced the scFv clones that showed the strongest binding to the tumour extract and performed a Blast search of the Genbank gene database. The $V_{H}DJ_{H}$ heavy chain gene used by all members of B-cell clone D4 exhibited 89% homology with a human anti-HER3 MAb (AF048774) and the $V_{nj}$ light chain gene, also used by the same B-cells, matched a human anti-EGFR antibody (DQ666353.1) with 96% homology. These, and scFvs from the global libraries, were tested for binding to recombinant antigens from the epidermal growth factor receptor family: HER-2, HER-3 and HER-4, kindly provided by Genentech (San Francisco, USA) and Pharmexa A/S (Hørsholm, Denmark) by ELISA. Six scFvs from B-cell clone D4 and one from the global tumour library bound to recombinant HER-2, HER-3 & HER-4, indicating that they recognised a shared epitope expressed by all three members of this EGFR family of receptors (Fig. 12). Specificity for HER-2, HER-3 & HER-4 was confirmed using soluble scFv produced in the non-suppressor strain of *E.coli*. 

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Fig. 11. Exponential enrichment of scFv-phage after panning and elution on a breast tumour extract

A. $V_{H}/V_{k}$ and $V_{H}/V_{\lambda}$ scFv mini-libraries, using $V_{H}$-genes from B-cell clone D4, panned on pooled heterologous tumour extract. The eluate from each panning was then subjected to further cycles of panning and elution; B. Global $V_{H}/V_{k}$ and $V_{H}/V_{\lambda}$ scFv-phage libraries from a human breast tumour, panned on the same heterologous tumour extract as in A.
Clusters of B-cells, T-cells and follicular dendritic cells form within human ductal breast carcinomas, resembling the ectopic germinal centres observed in the target tissues of patients with autoimmune diseases. These clusters of lymphoid cells contain clones of proliferating B-cells that are undergoing somatic hypermutation of their rearranged, Ig V-(D)-J-genes and affinity selection of their B-cell receptors driven, in the case described here, by members of the epidermal growth factor receptor family Her-2, Her-3 and Her-4. The antibodies produced during this response recognise an epitope shared by these 3 cell surface receptors, which are known to be overexpressed on breast carcinoma cells and several other types of carcinoma, including ovarian cancer. It is very probable that other tumour associated antigens are also able to stimulate a local B-cell response within the tumour, e.g. antibodies specific for a ganglioside were cloned from medullary carcinoma B-cells, although it is not clear whether these are involved in attacking the tumour (Kotlan et al., 2005).

Single chain Fv antibodies cloned from tumour germinal centre B-cells can readily be converted to complete antibodies by splicing their V-genes on to Ig constant regions of any desired isotype. These fully human antibodies can be produced in large quantities in a protein expression system and are therefore potential candidates for diagnosis, monitoring and therapy of breast and other types of cancer.
6. General conclusions

It has become increasingly clear that infiltrating B- and T-lymphocytes organise themselves into ectopic lymphoid follicles and germinal centres within tissues undergoing inflammatory processes. This has been observed in several autoimmune diseases (Table 1), usually within the target organ or tissue, myasthenia gravis being the exception to the rule for reasons discussed in section 3.5. However, ectopic g.c.s are not restricted to autoimmune diseases but can also develop in other chronic inflammatory diseases, such as Crohn’s disease and ulcerative colitis; at sites of infection such as the liver during chronic hepatitis C virus infection and the skin of oncocerciasis patients (Brattig et al., 2010); and in neoplasias, including lymphoma of the mucosal-associated lymphoid tissue associated with Sjögren’s syndrome (Bombardieri et al., 2007a) and, as shown here, in breast cancer (Table 1).

How these ectopic g.c.s develop, their role in the pathology of autoimmune diseases, and in combating infections and malignancies is still unclear but evidence is beginning to emerge. Cytokines, chemokines and signalling molecules involved in lymphoid neogenesis in the secondary lymphoid organs also appear to be required for ectopic g.c. formation, including lymphotoxins-α, β and α1β2, TNFα, Grb2, the chemokine receptor CXCR5, its ligand CXCL15, and the B-cell attracting chemokines CXCL13 and CCL21, in this case due to release of these molecules within the inflammatory environment, suggesting that g.c. formation may be secondary to inflammation (Aloisi & Pujol-Borrell, 2006).

When reports first emerged of germinal centre-like structures within the target tissues of autoimmune diseases, there was some scepticism regarding whether these structures were involved in true germinal centre reactions. These doubts have now been dispelled. Identification of dark and light zones and a follicular dendritic cell network in intimate contact with B-cells was highly suggestive of a germinal centre reaction, especially when it was shown that autoantigen was trapped on the follicular dendritic cell processes, e.g. (SHIONO et al., 2003). Studies by us and other researchers have shown that the B-cells within ectopic germinal centres are activated to antigen-driven clonal proliferation, somatic hypermutation and class switching, similar to the response in orthotopic g.c.s responding to foreign antigens. That they switch on the somatic hypermutation machinery has been shown in several autoimmune diseases by sequencing studies of the expressed, rearranged Ig V-genes cloned from microdissected g.c.s. This has been confirmed in the salivary gland g.c.s of Sjögren’s patients and the synovial g.c.s of rheumatoid arthritis patients by identification of activation induced cytidine deaminase (AID), a key enzyme in somatic hypermutation and class switch recombination (Bombardieri et al., 2007b; Humby et al., 2009). Interestingly, expression of AID has recently been observed in hyperplastic fibroblasts of rheumatoid arthritis patients (Igarashi et al., 2010). Expression correlated with mutations in the p53 gene and was induced by TNFα in vitro. AID is known to induce mutations in non-Ig genes at a lower frequency and it was suggested that the mutations of this tumour suppressor gene may be the cause of the fibroblast hyperplasia. Affinity maturation of B-cell receptors during somatic hypermutation has been demonstrated by analysis of replacement mutations, although early analyses failed to take into account the bias towards replacement mutations in the CDRs resulting from targeting of AID to sequence motifs such as RGYW. Final confirmation requires direct affinity measurements of autoantibodies cloned from germinal centre B-cells and/or by 3-D molecular modelling of the antigen-binding site bound to its epitope, as we have shown for anti-hen egg lysozyme antibodies produced in an orthotopic germinal centre reaction (Adams et al., 2003).
In several cases it has been shown that the autoantibodies generated in ectopic g.c.s have similar specificities to the autoantibodies found in the blood, notably in Hashimoto’s thyroiditis, Sjögren’s syndrome and rheumatoid arthritis, suggesting that the g.c.s contribute to pathological mechanisms, although whether they are critical in the early stages of development of the disease, or only contribute to its maintenance once the initial tissue damage has commenced, has yet to be established. Production of cytokines and chemokines at sites of damage that attract lymphocytes and contribute to lymphoid neogenesis suggests that the latter may be the more likely scenario. Nevertheless, a detailed understanding of the mechanisms involved in generation of ectopic g.c. structures and maintenance of production of plasma cells and memory B-cells producing potentially pathogenic antibodies is essential for a full understanding of the pathology of autoimmune disease and holds promise for developing new methods of therapy, based on controlling this response or inducing immunological tolerance to the autoantigens.

Even more work needs to be done to determine the role of ectopic g.c.s in other diseases, including sterile and infectious chronic inflammatory diseases and cancer. What other types of cancer, in addition to breast cancer and lymphoma, induce germinal centre reactions within the tumour and the nature of their response in elimination of cancer cells have yet to be determined. The identification of intra-tumour g.c.s producing antibodies and memory B-cells with specificity for members of the epidermal growth factor receptor family holds out hope that therapeutic vaccines can be developed to boost this response for therapy of breast cancer and, potentially other neoplasias, such as ovarian cancer, in which these molecules are overexpressed. Experimental approaches using mouse models of breast cancer support this optimism (Renard et al., 2003; Renard & Leach, 2007; Mukhopadhyay, MS in preparation). Cloning of antibodies against tumour-associated antigens from intra-tumour g.c.s is also a novel way of producing fully human antibodies for passive immunotherapy.

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8. References


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The Ectopic Germinal Centre Response in Autoimmune Disease and Cancer


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Autoimmune Disorders – Current Concepts and Advances from Bedside to Mechanistic Insights


Autoimmune disorders are caused due to breakdown of the immune system, which consequently fails in its ability to differentiate "self" from "non-self" in the context of immunology. The diseases are intriguing, both clinically and immunologically, for their diversified clinical phenotypes and complex underlying immunological mechanisms. This book offers cutting-edge information on some of the specific autoimmune disease phenotypes, respective diagnostic and prognostic measures, classical and new therapeutic options currently available, pathogenesis and underlying mechanisms potentially involved, and beyond. In the form of Open Access, such information is made freely available to clinicians, basic scientists and many others who will be interested regarding current advances in the areas. Its potential readers will find many of the chapters containing in-depth analysis, interesting discussions and various thought-provoking novel ideas.

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