FOLDING AND ASSEMBLY OF ANTIBODIES

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I. A short introduction into protein folding

I-1 General considerations

Hardly any field of biological research has undergone such considerable changes and advancements in recent years as has the field of protein folding. Almost five decades ago the observation that protein folding in the test tube is reversible (Anfinsen and Haber, 1961) laid the basis of the field. Countless studies on protein folding have been published since then, and especially in the last few years, the advent of new techniques has pushed barriers and protein folding emerged as one of the first truly interdisciplinary fields in biological sciences. This work is supposed to contribute to this development, in particular by bridging the gap between in silico, in vitro and in vivo studies on protein folding for a well known model system: antibodies.

I-2 A brief history of protein folding

As soon as it had been appreciated that proteins are irregularly but well defined structured molecular objects, the question of how proteins could be able to self-organize began to bother scientists. The adverse effect of this motivation was that the structural perspective of the protein folding problem shaped its perception. In other words, early protein folding studies aimed at a well-defined description of the protein folding phenomenon with only few states involved, unfolded, native and intermediate, often applying the conceptual framework of organic chemistry developed for small molecules. The new view of protein folding, despite several shortcomings, has highlighted this conceptual dilemma of protein folding (Dill and Chan, 1997; Leopold et al., 1992; Onuchic et al., 1995). Some of its basic ideas had already been developed earlier and applied to the native state of proteins (Austin et al., 1973; Frauenfelder et al., 1979; Hartmann et al., 1982). Together they shifted the focus, and nowadays proteins are regarded as highly dynamical objects. This dynamical character should not only influence the native state of proteins but in particular be a dominant factor in folding (Hartmann et al., 1982; Leopold et al., 1992; Henzler-Wildman et al., 2007a; Henzler-Wildman et al., 2007b). Dynamics implies
heterogeneity and heterogeneity suggests parallel rather than strictly sequential pathways for protein folding. This was a main idea of the new view which arose from a merge of polymer physics and protein science (Dill and Chan, 1997; Leopold et al., 1992; Onuchic et al., 1995). It put forward that polypeptide chains navigate on a free energy hypersurface which is generally funnelled towards the native state (Figure I-1). Accordingly, already few interactions can tip the energetic balance of a polypeptide chain, predisposing it towards further movement down the funnel and finally the native state(s) (Dill and Chan, 1997; Onuchic et al., 1995). Even though the funnel concept alone does not necessarily explain different protein folding phenomena and is very often applied quite generously, it facilitated novel perspectives on protein folding as will be outlined in the following.

Figure I-1: A schematic protein folding funnel. Shown are possible folding funnels for a two-state (A) and a multi-state folder (B). The smoothness and the number and characteristics of the available energy minima on the free energy hypersurface will shape the folding mechanism. Figure adapted from (Bartlett and Radford, 2009).

I-3 The unfolded state

Originally, the unfolded state of proteins was termed random coil. This implies the absence of defined interactions or conformations of an unfolded protein. It is accordingly expected to behave like a random polymer in a good solvent. This view is out-dated. It was mainly founded on techniques with too low resolution or too low sensitivity which could hence only provide global properties for the investigated
proteins. In contrast, several recent studies on chemically denatured proteins, even under harsh conditions, have been able to detect significant residual structure, local and non-local, native and non-native (Crowhurst et al., 2002; Mok et al., 2007; Sanchez and Kiefhaber, 2003a; Le Duff et al., 2006; Smith et al., 1996). This residual structure can be expected to have a pronounced effect on the folding free energy landscape of a protein. If native, it can entropically destabilize the unfolded state thereby rendering folding more favourable. The classical examples are disulfide bridges (Pace et al., 1988; Clarke et al., 1995a; Clarke et al., 1995b; Abkevich and Shakhnovich, 2000; Eyles et al., 1994), which historically also played a significant role as probes to study protein folding (Creighton, 1974; Creighton, 1975). Non-covalent interactions, though, seem to be far from absent in unfolded states of proteins. If non-native, enthalpical stabilization of the unfolded state might be a consequence. It should be kept in mind, however, that most studies on the unfolded state of proteins dealt with rather harsh conditions of elevated temperature or high denaturant concentrations. Under physiological conditions, the amount of structure in unfolded states of proteins can be expected to be significantly higher as have revealed the few accessible cases (Religa et al., 2005; Marsh et al., 2007; Chugha et al., 2006) and a compaction of chemically unfolded proteins with lowering denaturant concentrations suggest (Sherman and Haran, 2006; Merchant et al., 2007). Even more important, in vivo, proteins encounter quite a different scenario than dilution from a chemically denatured state. They begin their folding process often during translation or translocation; processes, which only make parts of a protein at a certain time amenable for folding, and in particular to form non-local interactions (Kosolapov and Deutsch, 2009; Zhang et al., 2009; Kimchi-Sarfaty et al., 2007; Tsai et al., 2008; Clarke and Clark, 2008; Hsu et al., 2007; Hsu et al., 2009). Furthermore, if folding occurs, it takes place in a confined space like the maximum 20 Å wide ribosome exit tunnel (Ban et al., 2000) which is chemically distinct from an aqueous solution (Lu et al., 2007).
I-3 Movements on the free energy hypersurface

Proteins are far more complex than the small molecules transition state theory has been developed for. Furthermore, in protein folding reactions a large number of weak non-covalent interactions are formed and broken and shaped by entropy, in contrast to the small molecules of organic chemistry. It is thus very unlikely that the dynamics of polypeptide chains can be adequately described by a system developed for a completely different class of molecules. Significant recent progress has been obtained in this field. Using fast spectroscopic techniques (Lapidus et al., 2000; Fierz et al., 2007), probes which are straightforward in the interpretation of the data (Krieger et al., 2003) and molecular dynamics simulations (Yeh and Hummer, 2002) has been a fruitful approach which provided significant new insights. Polypeptide chains are now believed to explore the free energy surface limited by local geometry around certain amino acids (Krieger et al., 2005) as well as solvent parameters and intrinsic hydrogen bonding (Moglich et al., 2006). Additionally, local chain motions have been found to be strongly coupled to motions of other chain segments (Fierz and Kiefhaber, 2007). Overall, polypeptides seem to move on a rough free energy surface within timescales of picoseconds to microseconds (Fierz et al., 2007) with some generalizable but also sequence-specific dynamic parameters (Krieger et al., 2003) not only in the unfolded state (Moglich et al., 2006), but also in partially folded or native states (Fierz et al., 2009).

I-4 The transition state

As the transition state of any chemical reaction represents a maximum on the free energy surface, it never becomes significantly populated in equilibrium. This makes indirect techniques necessary to explore its structural and dynamical features. Ablating specific interactions in a protein’s native state and monitoring the impact on the folding kinetics, i.e. the stabilizing effect of this specific interaction on the transition state, has been coined *Phi-value* analysis for protein folding reactions (Matouschek et al., 1989). Being an energetics-based approach, it has been considerably debated in its structural interpretation, the adequate way of mutating a protein and the impact on other states than the native and the transition state.
Undoubtedly, though, it has provided insights into an otherwise elusive state of protein folding (Fersht et al., 1992; Matouschek et al., 1992a; Matouschek et al., 1992b; Matouschek et al., 1989; Serrano et al., 1992a; Serrano et al., 1992b; Serrano et al., 1992c; Cota et al., 2001; Hamill et al., 2000; Lappalainen et al., 2008). In particular, the combination with molecular dynamics simulations proved to be a rewarding approach as Phi-value analyses can only provide local information which, when coupled to molecular dynamics simulations, can be extended to provide global conformational features of protein folding transition states (Mayor et al., 2003; Geierhaas et al., 2004; Paci et al., 2003; Vendruscolo et al., 2001). Based on these analyses, protein folding transition states are now mostly believed to roughly resemble expanded counterparts of a proteins native state – with a native-like topology yet only very few tertiary interactions (Vendruscolo et al., 2001; Lindorff-Larsen et al., 2004).

I-5 Protein folding intermediates

Originally believed to be the key to circumvent the famous Levinthal protein folding paradoxon, intermediates in protein folding have lived through a quite diverse career. They are considered to be partially folded states of proteins, which can either accelerate, decelerate or even misguide protein folding (Wagner and Kiefhaber, 1999; Mucke and Schmid, 1994; Liu et al., 2000; Jahn et al., 2006; Jahn and Radford, 2008). As such, they were hard to position in classical protein folding descriptions. Yet, being considered as local minima on the protein folding energy hypersurface, their role and properties come quite natural (Brockwell and Radford, 2007). If separated by high free energy barriers, intermediates can be readily observable which directly implies that non-observability does not mean absence. Indeed, for a lot of proteins previously believed to be two-state folders partially folded states could be detected with recent techniques of higher sensitivity and time resolution (Sanchez and Kiefhaber, 2003b; Bai, 2006; Lipman et al., 2003; Korzhnev et al., 2004). Accordingly, the population of partially folded states is rather a gradual than an all-or-none process with the possible and highly debated extreme case of barrierless downhill folding (Onuchic and Wolynes, 2004; Garcia-Mira et al., 2002; Sadqi et al., 2006; Ma and Gruebele, 2005; Ferguson et al., 2007). Intermediates can be
stabilized by native or non-native secondary and tertiary structure interactions (Bai, 2006; Brockwell and Radford, 2007; Creighton et al., 1996; Feng et al., 2005; Kameda et al., 2005; Kiefhaber et al., 1995; Korzhnev et al., 2004; Liu et al., 2000; Religa et al., 2005; Udgaonkar and Baldwin, 1988). As the underlying interactions are *per se* unspecific, it is not surprising that a whole plethora of different intermediate characteristics is observed. First high resolution structural information on folding intermediates (Hughson et al., 1990; Kiefhaber et al., 1995; Udgaonkar and Baldwin, 1988; Kameda et al., 2005; Korzhnev et al., 2004; Mizuguchi et al., 2003; Religa et al., 2005) or complete folding pathways (Feng et al., 2005; Mayor et al., 2003) put them in a structural line with the transition states: well defined topology, sometimes ill-defined native interactions or non-native interactions underlining the hierarchical character of protein folding and the role of topology for protein folding mechanisms in general (Plaxco et al., 1998; Zarrine-Afsar et al., 2005). In contrast to the transition state, however, intermediates can be populated for significant amounts of time and therefore might be bifurcation points of a protein folding reaction towards the native or misfolded potentially harmful states (Liu et al., 2000; Kameda et al., 2005; Jahn et al., 2006; Jahn and Radford, 2008; Simpson et al., 2009).

**I-6 The native state**

Quite unexpectedly, the native state of proteins, formerly considered as the most “reliable” of all states, has undergone a drastic change in its perception, too. It has been considered dynamic for a long time (Austin et al., 1973; Frauenfelder et al., 1979; Hartmann et al., 1982) and recent technical advances have tightened the boundaries between dynamics and functions of proteins (Eisenmesser et al., 2005; Henzler-Wildman et al., 2007a; Henzler-Wildman et al., 2007b). In particular, studies on single protein molecules emphasized the ensemble character not only of unfolded, partially folded but also the native state of proteins (Lipman et al., 2003; Schuler et al., 2002). The most surprising discovery, however, was that a significant amount of naturally occurring proteins do not or hardly possess any defined native three-dimensional structure. This class of proteins has been named *intrinsically disordered* and is believed to make up almost 40% of the human proteome (Dunker et al., 2008; Uversky et al., 2005; Yang et al., 2005). Historically, this class of proteins has been
overlooked for a long time as most early biochemical studies focussed on enzymes where well defined structures are a prerequisite for well defined functions. Intrinsically disordered proteins are consequently underrepresented in this functional class. In contrast, a significant amount of proteins involved in signalling and regulation are thought to be unstructured (Dunker et al., 2008; Uversky et al., 2005; Yang et al., 2005). In general unstructured until assembled with their partners, which can either be small molecule ligands or, in most cases, other proteins (Dyson and Wright, 2002; Fink, 2005). Action through interaction thus in particular applies to these proteins which are able to couple high specificity with low affinity by enthalpy-entropy compensation. Sometimes promiscuous and often highly dynamic regulatory networks can thus be realized by intrinsically disordered proteins.

**I-7 Multidomain and multimeric proteins**

Being experimentally and theoretically much easier to address than oligomeric proteins, isolated one-domain proteins have dominated protein folding studies. Yet, in the organism, almost all proteins are found to be multidomain oligomers (Han et al., 2007) or at least involved in transient protein-protein interactions to fulfil their function. The evolutionary reasons to form oligomeric structures are diverse. Functionally, heterooligomers possess the advantage of being modular and therefore evolutionary more adaptable. Furthermore, oligomers or multidomain proteins can in principle work more efficiently on substrates by e.g. substrate channelling. In general, the stability of a protein is increasing with the hydrophobic surface buried upon folding (Miller et al., 1987). Therefore, based on simple surface to volume considerations, larger proteins have the possibility to reach higher stabilities than their smaller counterparts. This opens up the possibility for more non-ideal local geometries which might functionally be necessary. In particular, the symmetric or pseudo-symmetric association of smaller building blocks is energetically favourable (Ponstingl et al., 2005) and concomitantly reduces the problems of transcription/translation errors which are more likely to occur in monocistronically encoded large proteins. On the other hand, oligomeric proteins have to be kinetically stabilized to avoid dissociation in the crowded cellular milieu with often very low concentrations of the cognate association partners. Furthermore, in particular for multidomain proteins, individual
domains have to be evolved for robust folding in the local proximity of other domains being in the course of folding (Han et al., 2007).

Of course, the same principles deduced for the folding of small proteins still hold but often they have to be extended. One example is the stability of oligomeric structures. Up to 50% of the observed free energy change upon folding can be attributed to interactions of the association surfaces which often has a significant effect on the observed folding behavior (Jaenicke and Lilie, 2000). In many cases, partially structured intermediates or very unstable monomers form which only upon association are able to attain their native structure and/or stability. The association-coupled structure formation processes and the necessary recognition specificity of partially folded species, conceptually intimately linked to intrinsically disordered proteins, are still largely open questions.

I-8 The evolution of foldable polypeptides and folding mechanisms

Proteins are not primarily evolved for folding but for function. This becomes obvious for intrinsically disordered proteins. Though, as function is in most cases linked to structure, folding comes as a necessary additional selection criterion for proteins and dominant negative effects or protein misfolding have to be avoided. Indeed, folding mechanisms and protein structures seem to be more conserved than functions or sequences in protein families (Chothia and Lesk, 1986; Zarrine-Afsar et al., 2005). It should be kept in mind that at a first glance quite different folding reactions, e.g. two-state or multi-state, might be dictated by the same folding mechanisms (Friel et al., 2004; Spudich et al., 2004). The smoothness of the underlying free energy surface will govern the population of partially folded species; topology and few key interactions are believed to make up the general mechanism (Bartlett and Radford, 2009). On the one hand, this argues for a high robustness of folding pathways against mutational changes. On the other hand, it might imply that efficient folding to a defined structure is far from being trivial to achieve evolutionary and therefore conserved in the cause of evolution. Indeed, even though random polypeptide sequences can sometimes fold into a defined structure, they do so in an uncooperative manner (Davidson and Sauer, 1994) and structure-based design of proteins led to highly complex and uncooperative folding pathways (Dantas et al.,
2003; Scalley-Kim and Baker, 2004; Watters et al., 2007). This shows that simple and cooperative folding pathways are not physical prerequisites for a defined protein structure but evolutionary selected traits. Cooperativity is assumed to be achieved by avoidance of an over-stabilization of the native state, selective destabilization of partially folded states, favorization of non-local stabilizing interactions and overlapping folding nuclei – non-trivial features of a protein (Chan et al., 2004; Watters et al., 2007; Haglund et al., 2008). They are upheld even if in conflict with fast folding (Capraro et al., 2008; Jager et al., 2006) or functional needs (Jager et al., 2006; Nordlund et al., 2009; Friel et al., 2009) pointing towards their importance for the homeostasis of the organism.

I-9 A synopsis?

Our understanding of protein folding has undergone considerable progress in recent years. Even structure-prediction and de novo protein design can now be carried out with a low but remarkable success rate (Dahiyat and Mayo, 1997; Kuhlman et al., 2003), often still relying on semi-empirical approaches, though. A common understanding of protein folding, even though a lot of details are still highly debatable, entirely missing and most likely crucial, might be the following: Proteins start from a non-native state, which can already possess local and/or non-local fluctuating structural elements and thereby are limited in their conformational freedom. The polypeptide chain explores the ensemble of conformations accessible in a time scale of picoseconds for local conformational changes to nanoseconds or even microseconds for more global conformational changes. Once crucial residues have come into contact, folding can proceed via a transition state with a roughly native-like topology and different degrees of structural polarization. From this state of high free energy, folding can proceed to partially folded states which will be more native-like but might possess non-native elements and interactions. Dependent on the successive free energy barriers to complete folding, these states can be absent, populated very transiently or quite stably before complete folding to the native state occurs via much more native-like transition states. Dependent on the cooperativity of the process, the number of folding nuclei (“foldons” (England et al., 2007)), the nature and kind of rate limiting-steps etc. a variable number parallel pathways, mostly
differing in local properties, can be taken finally resulting in a dynamic native state (Figure I-2). Even though all these considerations should also hold in vivo, a considerable modulation of the outlined principles can be expected due to the nature of the environment and the ubiquitous presence of folding helpers and catalysts (Walter and Buchner, 2002; Hartl and Hayer-Hartl, 2009; Kramer et al., 2009).

**Figure I-2: A generalizing view on protein folding.** From the unfolded ensemble, where the protein can sample multiple states within ps to µs, folding proceeds via a first transition state. This is assumed to be roughly native like in topology with only very few key interactions established. In a subsequent intermediate state, first structural elements rigidify before complete folding to the native state can occur. Typical timescales of the processes are indicated. As folding takes place on a funnel-shaped multidimensional energy landscape, the indicated pathway A is just one of the multiple pathways possible (e.g. an intermediate-free pathway B) and bifurcation might occur at different stages.
II. Antibody folding and assembly – classical themes & novel concepts

II-1 Protein folding in the endoplasmic reticulum

Naturally, proteins do not fold in the well controlled environment of the test tube but in the crowded cellular environment. Posttranslational modifications and chaperone interactions can and will occur in most cases – features which will heavily influence most folding reactions. Approximately one third of all human proteins is destined for secretion or transport to cellular membranes and therefore has to traverse the endoplasmic reticulum (ER). Polypeptides generally enter this compartment cotranslationally. Accordingly, the ER represents the major folding compartment of the cell and sets the quality control standards all transported proteins have to pass. It is thus compulsory for the ER to be equipped with a large variety of folding helpers and catalysts (Meunier et al., 2002). The most prominent ER chaperones are Hsp70 family members like BiP (Haas and Wabl, 1983) and the accessory J-proteins, ERdj’s (Dudek et al., 2009), the Hsp90 homologue Grp94 (Melnick et al., 1994; Ostrovsky et al., 2009) and the lectins calnexin and calreticulin (Helenius and Aeby, 2004; Trombetta and Helenius, 1998). Folding catalysts like protein disulfide isomerases (PDIs) (Freedman, 1989; Jessop et al., 2004; Ellgaard and Ruddock, 2005) and peptidyl-prolyl isomerases (PPIases) (Fischer et al., 1989; Lang et al., 1987) are also abundant in the ER.

Generally, folding and posttranslational modification of proteins in the endoplasmic reticulum occur with high efficiency and fidelity. Nevertheless, the ER folding machinery can become acutely or chronically overwhelmed. Therefore, the ER has evolved backup mechanisms to cope with ER-folding stress: the unfolded protein response (UPR) (Mori et al., 1992) and the ER-associated degradation (ERAD) (Lippincott-Schwartz et al., 1988; McCracken and Brodsky, 1996). ER-load is sensed by at least three sensors in the ER membrane of higher eukaryotes, IRE1, PERK and ATF6, which can, if necessary, induce an upregulation of the ER folding machinery via a signalling cascade to the nucleus (Ron and Walter, 2007) and additionally shut down the synthesis of new proteins via phosphorylation of translation initiation factors (Harding et al., 2000). If stress persists, the UPR can even lead to the induction of apoptosis (Lin et al., 2007). Terminally misfolded proteins can be retro-translocated
from the ER to the cytosol, ubiquitylated and degraded by the proteasome in the ERAD pathway (Vembar and Brodsky, 2008). Key players of the folding machinery in the ER, as well as the major ER stress response mechanisms, UPR and ERAD, have been deciphered in recent years. What is still missing though is a detailed insight into the molecular mechanisms of folding and quality control in the ER. Only a combined biophysical and cell biological approach can address this complex issue and has been applied to the folding of antibodies within this work.

II-2 A short overview over antibody biology

For most genes, great care is taken to ensure that mutations are rapidly detected and repaired in order to maintain primary amino acid sequences that fold efficiently into biologically active proteins. In contrast, highly active mutational mechanisms are essential to generate antibody diversity and fast adaptability of the immune response, which clearly increases the likelihood of producing immunoglobulin (Ig) proteins that are incapable of folding, assembling or being transported to the cell surface. Failure to closely monitor and control maturation of antibodies would wreak havoc on the proper functioning of the immune system, which can only be avoided by the action of robust folding and quality control systems.

Ig proteins serve as cell surface antigen receptors on B cells, and upon antigen stimulation and plasma cell differentiation they are secreted as soluble effector molecules (antibodies) that provide protection against infections and foreign antigens. In their simplest form, the IgG antibodies, each molecule is composed of two identical heavy chains (HC) and two identical light chains (LC) that are linked by disulfide bonds. Both chains are composed of multiple domains; each composed of ~100 amino acids (Figure II-1A). The N-terminal domain of each chain is unique and is therefore called the variable domain (V_H respectively V_L). Together both variable domains form the antigen recognition region (Figure II-1A). Within the variable domains are stretches of particularly diverse amino acids (hypervariable regions) that provide the exquisite binding specificity of the antibody molecule. The remainder of the polypeptides is conserved within antibody classes (constant domains) and is...
important for effector functions like complement activation or recruitment of macrophages and natural killer cells. Five different classes or isotypes of antibodies are made in most higher vertebrates: IgM, IgG, IgA, IgE, and IgD that differ in the HC constant regions used. Only two different types of LC (κ or λ) generally exist, which can assemble with all HC classes. However, in a given cell only one heavy chain and one light chain allele are expressed, so that antibodies with a single specificity are produced.

II-3 Biosynthesis of antibodies in the cell

The development of progenitor cells committed to the B cell lineage is characterized by the sequential expression of the HC and LC subunits. In preB cells, variable (V_H), diversity (D_H) and joining (J_H) gene segments are rearranged at the DNA level of a single allele to create a variable region (Maki et al., 1980; Early et al., 1980), which is then spliced to the constant region at the mRNA level (Calame et al., 1980). Once a functional HC is made, gene rearrangements begin for the LC, to form the light chain variable region (Bernard et al., 1978). The B cell is characterized by surface expression of Ig molecules, which are composed of two identical HC that possess a hydrophobic membrane anchor sequence and two identical LC.

DNA rearrangements that give rise to Ig variable regions are characterized by imprecise joining of the three gene segments, the addition of non-templated bases at the site of joining of these gene segments, and finally, during later stages of differentiation, the directed hypermutation of the variable region exons of HC and LC genes (Alt et al., 1987). These mechanisms are essential to generate antibody diversity and allow affinity maturation of the immune response, yet they clearly increase the likelihood of producing a protein that is incapable of folding and assembling properly, being transported to the cell surface or secreted, or engaging the appropriate signaling molecules thus compromising the functioning of the immune system. Accordingly, B lineage cells are particularly dependent on the ER quality control system to ensure that only correctly assembled Ig molecules are transported to the cell surface. In light of this, it is not surprising that many of the major components of the mammalian ER quality control machinery (e.g. BiP (Haas and
Wabl, 1983; Bole et al., 1986), calnexin (Hochstenbach et al., 1992), GRP170 (Lilie et al., 1993), ERdj3 (Meunier et al., 2002), and pERp1 (Meunier et al., 2002)) were first identified by virtue of their association with HC, and that Ig molecules were some of the earliest identified substrates of ER folding enzymes (e.g. protein disulfide isomerase (Roth and Pierce, 1987) and prolyl isomerases (Lang et al., 1987; Lilie et al., 1993).

Like all cell surface and secreted proteins, HC and LC are co-translationally translocated into the endoplasmic reticulum (ER) and folding begins even before the polypeptide chains are completely translated, starting with the V domain and proceeding though the C domains (Bergman and Kuehl, 1979). IgGs predominantly assemble first as HC dimers to which LC are added one at a time (Baumal et al., 1971). The CH3 domain is important in this assembly, as IgG HC mutants with a deleted CH3 domain do not form HC dimers readily and are often secreted as HCLC ‘hemimers’ (Zack et al., 1981). N-linked glycans are also added to nascent HC co-translationally (Bergman and Kuehl, 1978) and can restrict the potential folding pathways available to the nascent chain due to their highly polar nature. The heavily glycosylated heavy chains of IgM require the glycans for assembly and transport, while monoglycosylated heavy chains of IgG mature properly even without them (Hickman and Kornfeld, 1978). Although in vitro folding studies have not addressed the role of glycans in Ig domain folding due to the fact that the isolated domains in most cases have been produced in bacteria, they have provided a molecular explanation for most of the folding and assembly processes previously identified in cells.

**II-4 Antibody structure and the evolution of the immunoglobulin fold**

In the case of IgG, the complete molecule is composed of two four-domain heavy chains and two two-domain light chains (Figure II-1A), whose overall shape resembles a “Y”. It should be noted, however, that the orientation of the two arms of the Y is flexible because of an unstructured hinge region between the first (CH1) and the second (CH2) constant domain of the heavy chain. The protease papain cleaves the IgG molecule in the hinge region and divides it into three functional segments, each of which is a dimer (Kalmanson and Bronfenbrenner, 1942; Porter, 1950). The
two N-terminal fragments are termed Fab fragments (for fragment antigen binding). The remaining fragment, the Fc fragment (for fragment crystallisable), is important for connecting antigen binding to antibody effector functions. Dimerization of the Fc fragment is largely mediated by interactions between the C\textsubscript{H}3 domains and stabilized by disulfide bonds in the hinge region. The C\textsubscript{H}2 domains only interact via the attached sugar residues (Figure II-1A). They determine the C\textsubscript{H}2-C\textsubscript{H}2 orientation and spacing and are therefore crucial for the binding of downstream effectors (Huber et al., 1976; Krapp et al., 2003; Feige et al., 2009b).

Each of the Ig domains forms a highly similar beta sandwich structure, known as the Ig fold. The Ig fold is characterized by a greek-key β-barrel topology in which the barrel is not continuously hydrogen-bonded but composed of two sheets forming a sandwich-like structure. The variable domains (Figure II-1B) are comprised of nine strands (abcc’c”defg) and the constant domains (Figure II-1C) of seven strands (abcdefg) (Bork et al., 1994). In most antibody domains, a buried disulfide bridge connects strands b and f, which spans ~60-70 residues (Huber et al., 1976; Bork et al., 1994). It is orientated roughly perpendicular to the individual sheets and significantly stabilizes the folded domain (Goto and Hamaguchi, 1979). Another characteristic feature antibody domains share is a conserved tryptophan residue that is located in proximity to the internal disulfide bridge. Since its fluorescence is specifically quenched only in the native state by the adjacent disulfide bond, it can be used as a reporter group for the conformational state of antibody domains (Goto and Hamaguchi, 1979). Additionally, (cis-) proline residues are unusually abundant in antibody domains, contributing up to 10% of the amino acids. Particularly characteristic is a conserved cis-proline residue in the loop connecting strand b and c of the constant domains which is often preceded by an aromatic amino acid (Figure II-1C).
Figure II-1: Overall antibody structure and domain architecture. (A) Domain arrangement of an IgG antibody molecule. The light chains are shown in green, the heavy chains in blue. The oligosaccharide between the C_H2 domains is depicted as grey hexagon. Interchain disulfide bridges and important functional elements of the antibody (antigen binding paratope, Fab fragment, Fc fragment) are indicated. Domain architecture of the light chain variable domain (V_L) (B) and the light chain constant domain (C_L) (C). The strand nomenclature is indicated. The intrachain disulfide bridge (yellow) and the proximal conserved tryptophan residue (blue) are shown. The proline residues of the two domains are shown in green with the highly conserved cis-proline residue between strands b and c of C_L highlighted in a CPK representation. Small helices (red) connect strands a and b and strands e and f of the C_L domain.

Even though the Ig domain or “fold” is characteristic of antibodies, it is in fact one of the most widely used protein topologies in nature, giving rise to what is referred to as the Ig superfamily (IgSF). The origin of the Ig fold dates back to ~750 million years of evolutionary history, with the identification of IgSF members already in sponges (Du et al., 2004; Hsu et al., 2006). In contrast, the ability to produce antibodies is a more recent occurrence (~500 million years), dating back to cartilaginous fish, such as sharks, skates and rays (Hsu et al., 2006; Dooley and Flajnik, 2006). In vertebrates, the Ig fold is the major building block of extracellular recognition systems (Williams and Barclay, 1988; Barclay, 2003), while in invertebrates, IgSF members are more
limited to the neural system (Rougon and Hobert, 2003). The Ig fold has also been detected in prokaryotic and viral proteins, albeit it much less frequently, suggesting that it might have been acquired in these instances by horizontal gene transfer (Halaby and Mornon, 1998). The evolutionary success of the Ig superfamily can most likely be attributed to its robust fold, which provides stability against proteases and harsh environments, and the ability to build on this common core structure by incorporating highly diverse binding loops or edge strands.

II-5 From the folding of antibody domains to complete molecules

Antibodies are highly complex molecules. Dissection of the overall molecule into individual domains or fragments was necessary to determine folding differences between the structurally similar antibody domains. The pioneering studies on antibody folding were performed on secreted LCs that were denatured and allowed to refold in vitro (Goto and Hamaguchi, 1979). Further studies examined the constant domain of the light chain (CL) (Goto and Hamaguchi, 1982a) and IgG heavy chain CH3 domain (Isenman et al., 1979) and established that individual Ig domains could fold autonomously. A key characteristic in the folding of antibody domains, is that proline-isomerization reactions, which are intrinsically slow due to their high activation energy, strongly influence antibody folding (Goto and Hamaguchi, 1982a; Ramm et al., 1999; Feige et al., 2004), often as the rate limiting step (Thies et al., 1999; Feige et al., 2008a). Recent in vitro studies have addressed the folding of Ig domains in more detail with a view to include all the IgG constant region domains (Freund et al., 1996; Ramm et al., 1999; Feige et al., 2004; Feige et al., 2007; Feige et al., 2008a; Feige et al., 2009a; Lilie et al., 1995; Simpson et al., 2009; Thies et al., 1999; Thies et al., 2002). These studies provided the first high resolution data on antibody folding and revealed that, although all Ig domains appear very similar in terms of the final structure, they can be grouped into three different folding categories (Figure II-2).

In the first category, exemplified by the well-studied CL protein, domains are able to fold autonomously to a monomeric state. In the denatured protein, no residual structure was detected, regardless of whether or not its internal disulfide bridge was present (Feige et al., 2007; Feige et al., 2008a). However, once folding was initiated, the presence of an internal disulfide bridge exerted an important guiding impact on
the folding pathway (Goto and Hamaguchi, 1982b; Feige et al., 2007; Ramm et al., 1999). The reason for this became clear when studies revealed that the folding nucleus for Ig domains involves the clustering of hydrophobic residues in strands b, c, e and f (Freund et al., 1996; Geierhaas et al., 2004; Lappalainen et al., 2008). The term folding nucleus implies that these residues cluster together in an early reaction when the overall topology of the polypeptide chain starts to arrange. The covalent linkage of residues in strands b and f via a disulfide bond facilitates the establishment of this folding nucleus, creating a structured, on-pathway intermediate and preventing unproductive off-pathway interactions. Indeed, a population of misfolded, aggregation-prone off-pathway folding intermediates for Cl was detected in the absence of the internal disulfide bridge (Feige et al., 2007).

While folding intermediates are usually quite transient and elusive, they can be populated for longer times if a slow reaction limits the subsequent folding step. This is the case for Cl, where the major on-pathway folding intermediate is relatively long-lived due to the non-native trans isomerization state of a proline residue between strands b and c, which must isomerize to its cis state before folding can proceed (Figure II-2) (Feige et al., 2008a). Using NMR combined with molecular dynamics simulations, it was possible to follow the changes in the chemical environment of most amino acids and to obtain an atomic resolution view of the intermediate structure. It revealed that the core β-strands of the Ig topology were almost completely formed, while higher flexibility was observed for flanking strands, particularly strand d (Figure II-2) (Feige et al., 2008a), in keeping with data on other IgSF members (Mizuguchi et al., 2003; Kameda et al., 2005; Jahn et al., 2006). Importantly, two small helices linking strands a and b and strands e and f (Figure II-1C, Figure II-2) are natively structured very early (Feige et al., 2008a). They can be assumed to stabilize the orientation and spacing of the beta-strands of the Ig fold and furthermore correctly position a bulky hydrophobic residue in the core of the protein (Feige et al., 2008a). Thus, if present, these helices should render the folding of Ig domains more robust as they stabilize a highly-structured on-pathway intermediate in a conformation which is poised for subsequent productive folding. These small helices are highly conserved in most constant region domains and other members of the Ig superfamily (Bork et al., 1994; Barclay, 2003; Williams and Barclay, 1988) but are absent in several members of the IgSF, in particular those few, where high resolution data is available on folding intermediates (Mizuguchi et al., 2003; Kameda
et al., 2005; Jahn et al., 2006). Importantly, no such helices are found in variable antibody domains (Figure II-1B) or in several IgSF members that are prone to misfolding and amyloid formation (Kameda et al., 2005; Jahn et al., 2006; Qin et al., 2007), suggesting that differences between partially structured folding intermediates, e.g. the presence or absence of small, highly structured helices, might hold the clue for the tendency of IgSF members to either reliably fold or tend to misfold as could be shown for β2-microglobulin (Feige et al., 2008a). The structural insights gained for the major C_L folding intermediate seem to be transferable to other HC constant domains, in particular C_H2.

The second category of antibody domain folding pathways (Figure II-2) is represented by the C_H3 domain (Isenman et al., 1979; Thies et al., 1999). In addition to folding slower than the domains in the first category, this domain forms an obligatory homodimer. As with the C_L domain, a partially folded species was observed, which was trapped by a non-native prolyl isomerization state (Thies et al., 1999). Interestingly, the formation of the correct prolyl isomers was required for dimerization to occur, whereas the internal disulfide bridge was dispensable for folding and self-association (Thies et al., 2002). Thus, proline switches can influence not only folding but also dimerization.

The third, and most unexpected, category of antibody domain folding is the recently discovered, template-assisted folding of the C_H1 domain, which interacts with the C_L domain in the intact antibody (Figure II-1A). The isolated C_H1 domain is intrinsically disordered as determined by various spectroscopic techniques (Feige et al., 2009a). This is in marked contrast to all antibody domains previously studied and unexpected from the crystal structure of antibodies (Huber et al., 1976). To form a hydrophobic folding nucleus in the C_H1 domain, the interaction with a few key residues in the dimerization interface of the folded C_L domain is required. Unlike other domains where disulfide bond formation only enhances the folding process, it was absolutely required for C_H1. Another unique twist is that the rate-limiting proline isomerization between strand b and c observed in the folding of other Ig domains can only occur after association with the C_L domain. These observations are in agreement with previous studies on the complete antibody Fab fragment where C_H1 folding was proposed to be the slowest step occurring after association of the HC and the LC (Lilie et al., 1995), and underline the special role of the C_H1 domain in B cell biology.
Figure II-2: Three pathways of antibody domain folding. C_L folds via a highly structured on-pathway intermediate that is trapped by the trans state of a proline residue in the loop connecting strands b and c (highlighted in yellow). In the intermediate, the core β-sheet structure and the two short helices connecting strands a and b and strands e and f are fully formed (shown in red). The obligate dimer C_H3 folds via two intermediates, both most likely similar in structure to the one of the C_L domain. In a first, rapidly formed intermediate, a critical proline residue (highlighted in yellow) must isomerize to its native cis state, leading to a dimerization competent second intermediate. C_H1 is intrinsically disordered in isolation. Upon association with C_L it forms a loosely folded intermediate. In this intermediate, isomerization of the conserved proline residue between strands b and c (highlighted in yellow) limits the complete folding to the native state and formation of the interchain disulfide bridge between C_H1 and C_L. Accordingly, for most constant antibody domains, isomerization of a proline residue between strands b and c, which is usually preceded by an aromatic amino acid, is the rate limiting step for the folding of the major part of the molecule.

Together, the data reveal that the same final fold is reached for all antibody domains by overall conserved mechanisms, yet with strikingly different details. The biological significance of these differences becomes clear when one considers the biosynthesis of antibodies, where the C_H1 domain is indeed a built-in assembly sensor.
The mechanisms for generating between $10^7$ and $10^9$ different antibody specificities from a limited amount of genetic information are equivalent to extensive mutagenesis. Therefore, it is not surprising that a number of quality control checkpoints are required during B cell development and differentiation that monitor the integrity of the antibody (Figure II-3). The newly created HC produced by a preB cell is retained in the ER unless it associates with a "surrogate LC". This LC-mimetic is assembled from the $V_{\text{preB}}$ protein (contributing the variable domain) (Kudo and Melchers, 1987) and the $\lambda_5$ protein (supplying the constant domain) (Sakaguchi and Melchers, 1986). The preB-specific surrogate LC "tests" the ability of HC to assemble properly with a LC-like protein and to pass ER quality control measures. If properly associated, the HC will be transported to the cell surface and engage the B cell receptor signaling components (Karasuyama et al., 1990). The surrogate LC focuses in large part on the $C_{\text{H1}}$ domain, as unassembled $C_{\text{H1}}$-deleted HC can traffic to the cell surface of the preB cell and transmit a "functional" signal (Shaffer and Schlissel, 1997). Once the heavy chain is judged functional by these criteria, conventional light chain gene rearrangements commence. The $C_{\text{H1}}$ domain remains a critical focus of Ig quality control efforts throughout B cell development and plasma cell differentiation. Unlike LC, which can be secreted alone, HCs are retained in the ER and eventually degraded unless they assemble with LC (Figure II-3) (Mains and Sibley, 1983). LC loss variants of plasmacytomas were very rarely observed, whereas HC loss variants occurred much more frequently (Coffino and Scharff, 1971). This was argued to be due to the "toxicity" of free HC, which were neutralized by LC (Kohler, 1980). Exceptions to this rule occur in the rare B cell lymphoproliferative disorder known as Heavy Chain Disease, where truncated Ig HCs are secreted from cells without LCs. Notably, although these short HC have been identified for a number of different isotypes (i.e., IgA, IgG, and IgM), the deletions nearly always involve portions of the $V_{\text{H}}$ and $C_{\text{H1}}$ domains (Seligmann et al., 1979). Similarly, mouse plasmacytoma lines expressing HC with deletions of the $C_{\text{H1}}$ domain can secrete free HC (Morrison, 1978), whereas deletion of any of the other constant region domains does not permit this. Finally, the serum of Camelidae (i.e., single humped dromedary and llamas) contains a significant fraction of antibodies that are naturally devoid of LC. These HC only antibodies do not possess a $C_{\text{H1}}$ domain (Hamers-Casterman et al., 1993; Vu et
al., 1997), further underscoring the evolutionary conserved importance of this domain in regulating Ig transport and quality control.

Figure II-3: Immunoglobulin quality control checkpoints at various stages in B cell development. After HC gene rearrangements, preB cells produce BiP-bound \( \mu \) HC. If their association with the surrogate LC, which is assembled from the \( V_{\text{preB}} \) and \( \lambda_5 \) proteins, induces BiP release and folding of the \( C_{\text{H1}} \) domain and if the rest of the Ig domains fold properly, the HC can traffic to the plasma membrane and engage BCR signalling molecules. If there is a failure in any of these steps, the \( \mu \) HCs are retrotranslocated to the cytosol for degradation by the 26S proteasome. Once conventional LC are produced in the B cell, they similarly assemble with \( \mu \) HC, displace BiP from the \( C_{\text{H1}} \) domain and induce its folding. As the rest of the constant region domains were tested at the preB cell stage, only the pairing and folding of the V domains must pass the quality control scrutiny. Plasma cell differentiation leads to the synthesis of extremely high levels of antibodies. The ability of the specific HC and LC combination to assemble and fold properly was verified at earlier stages of development, thus quality control at this point involves monitoring the completeness of Ig assembly, focusing on the LC-induced release of BiP from the \( C_{\text{H1}} \) domain and its concomitant folding. In addition, there is a shift to the secretory form of \( \mu \) HC in plasma cells, which possess a terminal cysteine that is involved in assembly with J chain and pentamer formation (IgM). Thiol-mediated retention mechanisms monitor this assembly and prevent IgM monomers from being secreted (Anelli et al., 2003).
The term “ER quality control” refers to the process of monitoring the maturation of nascent secretory proteins and allowing only properly folded and assembled proteins to transit further along the secretory pathway. Proteins that fail to mature properly are retained and eventually retrotranslocated to the cytosol for degradation by the 26S proteasome in a process known as ER associated degradation (ERAD). Immunoglobulin heavy chain binding protein (BiP), the first component of the eukaryotic ER quality control apparatus to be identified, was found by virtue of its association with the unassembled, non-transported HCs produced in pre-B cell lines (Haas and Wabl, 1983). It interacts transiently with Ig assembly intermediates but not with completely assembled H2L2 molecules in plasmacytomas (Bole et al., 1986). BiP is the ER orthologue of the Hsp70 family of chaperones (Haas and Meo, 1988) and is retained in the ER, along with any protein associated with it, by virtue of a C-terminal KDEL tetrapeptide (Munro and Pelham, 1987). Similar to the differences detected by in vitro folding experiments, the folding requirements and dependence on BiP for the various Ig domains are quite different in cells. BiP binds transiently to some Ig domains (i.e., VL, VH, and some CH domains) (Kaloff and Haas, 1995), while other domains appear to fold rapidly and stably without ever interacting with BiP (i.e., CL) (Hellman et al., 1999; Lee et al., 1999). Only the CH1 domain interacts stably with BiP in the absence of LC (Hendershot et al., 1987) (Figure II-3). The association of this domain with BiP is crucial for controlling Ig assembly and transport, because its deletion, and the resulting ablation of BiP binding, leads to the secretion of incompletely assembled Ig intermediates. Studies to determine the folding status of proteins or domains in the ER of cells rely on the analysis of the oxidation status of cysteine residues, which often form disulfide bonds upon folding making the protein more compact and faster migrating on non-reducing SDS polyacrylamide gels (Braakman et al., 1992). Surprisingly, unlike other Ig domains, the CH1 domain remains reduced in the absence of LC (Lee et al., 1999). In contrast to the in vitro studies described above (Feige et al., 2009a), there was no evidence of BiP associating with an oxidized CH1 domain. This may be due to the fact that BiP does not appear to continuously cycle from HC in the absence of LC in cells, which is required for oxidation (Vanhove et al., 2001), or that association with light chain, oxidation, and folding of CH1 are more tightly coupled in vivo. In keeping with in vitro studies, although the ATP-mediated release of BiP from isolated HC resulted in CH1 domain oxidation, this was not sufficient for proper folding (Vanhove et al., 2001).
This argues that LC association is also required for folding of this domain in vivo. Only LC in which both domains are folded led to C_{H1} domain oxidation and secretion from cells. In vivo experiments also confirmed the requirement for proline isomerization in promoting C_{H1} oxidation, assembly with LC and secretion (Feige et al., 2009a). Thus, the evolution of a unique C_{H1} domain that absolutely requires assembly with a C_{L} domain for folding, allows the cell to ensure that newly produced HCs in B cells will be retained unless they are able to pass that first important test: their ability to combine with a LC and be transported only after assembly. It also ensures that plasma cells, which have been estimated to synthesize up to 10^3 antibody molecules a second, are not releasing partially assembled subunits that cannot properly bind to antigen or perform effector functions.

Many LC fold readily and can be secreted by themselves as either monomers (Dul et al., 1996) or dimers (Leitzgen et al., 1997), suggesting that the V_{L} domains of these LCs are likely to belong to categories 1 and 2 described above. However, LC exist that cannot be secreted alone due to a failure of the V_{L} domain to fold properly by itself (Skowronek et al., 1998). These LC fold perfectly well and are secreted when they assemble with a HC (Horibata and Harris, 1970), suggesting that their V_{L} domains may belong to category 3. This requirement for assembly assisted folding of some V_{L}, and presumably V_{H} domains, could explain the restriction of possible HC and LC pairings (Wiens et al., 1998) and might play a role in antibody misfolding diseases.

**II- 7 A comprehensive view of antibody folding, assembly and quality control – the current status and challenges ahead**

In summary, detailed biophysical and in vivo studies on individual Ig domains, antibody fragments and complete antibodies have identified common themes that provide a detailed picture of IgG folding (Figure II-4). Once the internal disulfide bridge is formed, most domains will autonomously fold in a three state reaction via on-pathway folding intermediates whose lifetime is increased by incorrect peptidyl-prolyl isomerization states. Peptidyl-prolyl isomerization reactions control folding, assembly and formation of disulfide bridges and might also play a role in inhibiting
aggregation. The very first steps in domain folding involve hydrophobic interactions in the inner face of strands b, c, e and f. Once folded, C_{H3} induces Fc dimerization and the covalent linkage of the heavy chains in the hinge region (Figure II-4). At this stage, all the constant region domains except C_{H1} are folded in a heavy chain dimer. Association of a folded light chain will induce folding of the C_{H1} domain, and the assembly of heavy and light chains will be stabilized by an interchain disulfide bond (Figure II-4). This picture shows nicely how the combination of biophysical properties of the individual domains revealed through in vitro studies and the delineation of cellular components controlling the synthesis of antibodies provide a comprehensive view of the basic mechanisms that govern antibody synthesis. Future work must focus on further integrating the role of the complex ER folding machinery in modulating, synchronizing and controlling assembly and folding of antibodies and other IgSF members.

**Figure II-4: An overall view over antibody folding and assembly.** Folding, formation of disulfide bridges and glycosylation of the heavy chain (blue) and light chain (green) begins co-translationally in the ER. The molecular chaperone BiP (red) interacts with most of the domains before folding is completed. All constant domains and most variable domains fold autonomously, populating an on-pathway intermediate on the way to the native state. C_{L} is known to fold particularly fast in the cell. Once C_{H3} is folded it induces HC dimerization which will be solidified by disulfide bridges in the hinge region. C_{H1} remains unfolded, unoxidized and stably bound to BiP until the LC displaces BiP and C_{L} induces folding of the C_{H1} domain. Once all C_{H1} prolines are in the correct isomerization state and C_{H1} is folded, a disulfide bridge between the LC and the HC forms. Most of the steps are likely to hold for other Ig classes. Chaperones and folding catalysts, like Grp94, protein disulfide isomerase PDI and the peptidyl-prolyl isomerase CyclophilinB are likely to contribute to the individual steps in immunoglobulin biogenesis.
III. A short summary of the work

Within this work, novel insights into the dynamics, folding and association of polypeptide chains in silico, in vitro and in vivo could be obtained. For simple oligopeptides, we were able to show that rugged energy landscapes govern their behaviour (Feige et al., 2008b) and laid the basis for future studies on biologically relevant molecules and molecular processes. Our structural work on the Fc fragment helped to answer a long standing functional question about IgG: why deglycosylation goes hand in hand with the complete loss of effector functions (Feige et al., 2009b).

The major efforts of our in vitro folding studies focussed on the constant domain of the antibody light chain, CL. There, we were able to show that the evolutionary conserved intrinsic disulfide bridge in antibody domains not only confers stability, but, as it is located in the folding nucleus, also guides the folding pathway and inhibits misfolding (Feige et al., 2007). Follow-up studies on the CL domain focussed on its major folding intermediate which we could trap in equilibrium and characterize at atomic resolution by a combined experimental and molecular dynamics approach.

Two small helices, previously unrecognized as important in immunoglobulin folding, were found to be completely structured in the intermediate and to decisively influence immunoglobulin folding for another member of the superfamily (Feige et al., 2008a).

Finally, we were able by a combined in vitro and in vivo study to provide a molecular description of the quality control mechanism antibodies have to pass prior to secretion from the endoplasmic reticulum. The key is an unfolded domain within the heavy chain, which strongly binds to the molecular chaperone BiP. Thereby it retains the heavy chain in the ER until paired with the light chain which induces folding (Feige et al., 2009a). Taken together, our work has provided significant new insights into the folding and assembly of antibodies from the test tube to the cell and paved the path for rational optimization of antibodies as well as further detailed studies on their folding, assembly and evolution.
IV. References


V. Published scientific papers
Rate of Loop Formation in Peptides: A Simulation Study

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Experimental techniques with high temporal and spatial resolution extend our knowledge of how biological macromolecules self-organise and function. Here, we provide an illustration of the convergence between simulation and experiment made possible by techniques such as triplet-triplet energy transfer and fluorescence quenching with long-lifetime and fast-quenching fluorescent probes. These techniques have recently been used to determine the average time needed for two residues in a peptide or protein segment to form a contact. The timescale of this process is accessible to computer simulation, providing a microscopic interpretation of the data and yielding new insight into the disordered state of proteins. Conversely, such experimental data also provide a test of the validity of alternative choices for the molecular models used in simulations, indicating their possible deficiencies. We carried out simulations of peptides of various composition and length using several models. End-to-end contact formation rates and their dependence on peptide length agree with experimental estimates for some sequences and some force fields but not for others. The deviations are due to artefactual structuring of some peptides, which is not observed when an atomistic model for the solvation water is used. Simulations show that the observed experimental rates are compatible with considerably different distributions of the end-to-end distance; for realistic models, these are never Gaussian but indicative of a rugged energy landscape.

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Introduction

Molecular dynamics (MD) simulations in principle provide information on proteins and other biological macromolecules that is unrivalled in spatial and temporal resolution. However, only short timescales are accessible due to limitations on the speed of computers. This prevents, for example, the direct simulation of given amino acid sequences folding to their stable biologically active conformation and also limits the extent to which the different force fields generally used in simulations can be evaluated by direct comparison to experiments. Proteins usually adopt their native state within milliseconds to seconds under ambient conditions; this is orders of magnitude away from the nanosecond regime accessible to MD simulations of “large” systems such as even a single protein in its aqueous environment.¹ An essential step in a protein folding reaction is the formation of a contact between two residues in the folding chain, which determines the closure of loops or formation of turns; these events occur on a submicrosecond timescale.²,³ The formation of contacts, whether native or not, dramatically reduces the fraction of conformations that have to be explored by a folding polypeptidic chain and is of fundamental importance in shaping the free-energy landscape of a foldable polypeptide sequence.⁴–⁶ The formation of contacts in an unfolded chain (depending on the length of the chain and the separation in sequence between the pair of residues) takes place on a timescale directly accessible to MD simulations.

A variety of experimental approaches have been developed to characterise these basic folding reactions.³ A particularly promising one is based...
on triplet–triplet energy transfer (TTET).\textsuperscript{2} In TTET experiments, the donor is excited by a laser pulse and is able to transfer its triplet state to a suitable acceptor within a few picoseconds, but only when the donor and acceptor are in van der Waals contact,\textsuperscript{7} so that donor quenching is limited by contact formation rather than by reaction time. TTET has been successfully applied to end-to-end contact formation in unstructured peptides from short to intermediate length and from picoseconds to nanoseconds under a variety of conditions.\textsuperscript{2,7–10} Another approach to measuring contact formation close to the diffusion limit has recently been proposed,\textsuperscript{11,12} it makes use of the rapid quenching of a long-lifetime fluorophore [2,3-diazabicyclo[2.2.2]oct-2-ene (DBO)] by tryptophan. A similar technique, in which a tryptophan at one end of the peptide is optically excited into a triplet state that is quenched upon contact formation with a cysteine at the other end, has also been applied for the same purpose,\textsuperscript{13} however, as molecular simulation clearly demonstrated,\textsuperscript{14} triplet quenching by cysteine is mainly reaction-limited and, therefore, does not allow direct measurement of contact formation kinetics.

Such experimental data can provide direct insight into the dynamics of an unfolded polypeptide chain. The conformational heterogeneity and the dynamics by which the conformation space can be explored play an essential role in the folding of a polypeptide chain. To date, most MD studies of peptides have focused on structured peptides, aiming at the reproduction of the experimentally observed structure (see, e.g., Ref. \textsuperscript{15} for a review) and neglecting to a certain extent the role the unfolded state plays in a protein folding reaction. However, neglect of the denatured state limits the utility of such simulations for predicting the stability of the native state and the rates of folding and unfolding, because the thermodynamics of the system as a whole depends fundamentally on the properties of both the native and the denatured states. Therefore, it remains essential to learn how best to simulate the dynamics of unfolded peptides and to validate these simulations against appropriate experimental data.

We carried out equilibrium simulations of peptides of various composition and length using several force fields, including implicit (i.e., a continuum model) and explicit (i.e., an atomistic model) solvation; beside such “realistic” models, we also used a model in which only the covalent bonds and the repulsive part of the van der Waals interactions have been retained. Polysine, \(S_x\), and poly(glycine–serine), (GS), which have been systematically characterised with TTET, have been chosen as models in our studies. Furthermore, we studied Y(GS), FSG peptides of different lengths in which the aromatic residues have been incorporated to mimic the donor–acceptor pair used in the TTET experiments. Our simulations provide a microscopic interpretation of the experimental data, yielding new insight into the disordered state of proteins. Conversely, the experimental data provide a new test of the validity and performance of the alternative molecular models used in the simulations, indicating deficiencies and strengths of each of these models.

Theory and Models

Implicit solvent simulations

Implicit solvent simulations have been carried out using different force fields. In one [SASA (solvent-accessible surface area)],\textsuperscript{16} the polar and nonpolar contributions of each atom to the free energy of solvation are assumed to be proportional to their solvent-accessible surface areas approximated with a simple analytical function; the electrostatic screening between solute charges is accounted for by a dielectric constant \(\varepsilon = 2r\), where \(r\) is the distance between charges, and by neutralising the formal charges (Asp, Glu, Arg, Lys, and the termini) as in the EEF1 (effective energy function 1) model.\textsuperscript{17} The EEF1 model\textsuperscript{17} is similar to SASA but uses excluded volumes instead of surfaces and a dielectric constant \(\varepsilon = r\) (i.e., charge screening is weaker than that in SASA). In ACE (analytical continuum electrostatics),\textsuperscript{18} the electrostatic solvation free-energy and the nonpolar (i.e., nonelectrostatic) solvation free-energy contributions are calculated separately. The electrostatic contribution is an analytical approximation to the solution of the Poisson equation; the nonpolar solvation free energy is approximated by a pairwise potential that yields results similar to the surface area approximation solvation energy; the version used here (ACE2) takes account of the overestimation of the desolvation of charges by the pairwise descreening potential. GBSW (generalised Born with simple switching)\textsuperscript{19} uses a generalised Born method for the electrostatic part and the solvent-exposed surface area for the nonpolar part with a phenomenological surface tension coefficient; the self-electrostatic solvation energy is calculated using a simple smoothing function at the dielectric boundary.

While SASA, EEF1, and ACE use a protein model in which nonpolar hydrogens are not explicitly included, all hydrogens are present with GBSW. Additional simulations have been performed with an all-atom model in which all the attractive interactions have been removed [ORI (only repulsive interactions)]. Langevin dynamics simulations were performed with the programs CHARMM\textsuperscript{20} at 300 K for 50 ps and using the Brooks–Brusinger–Karplus algorithm.\textsuperscript{21} Simulations with an alternative integrator\textsuperscript{22} have also been performed in a few cases and gave identical results. Simulations have been performed with a friction coefficient of 1 ps\textsuperscript{−1}. This is lower than the friction coefficient corresponding to the viscosity of water by a factor of \(\sim 60\) at 300 K.\textsuperscript{22} Since the Kramers theory\textsuperscript{23,24} predicts linear dependence of rates on

† [www.iapws.org/relguide/visc.pdf](http://www.iapws.org/relguide/visc.pdf)
friction coefficient in the limit of large friction, meaningful rates could be determined in this way while economising a factor of $\sim 60$ of the simulation time. For a few peptides, a broad range of frictions was explored to precisely extract the rate of contact formation in a medium of viscosity comparable to that of water and, thus, test the validity of Kramers’ prediction in our system. Simulation time ranged from 0.25 to 2 μs, depending on the peptide length. Full convergence of the average contact formation time and the distribution of the end-to-end distance was assessed by comparing results from the first and second half of each trajectory.

**Explicit solvent simulations**

Explicit solvent simulations were performed using GROMACS\textsuperscript{25} and the OPLS-AA force field\textsuperscript{26} in TIP4P water\textsuperscript{27} at 300 K. A cubic water box with a minimum distance of 13 Å to the ends of the extended initial peptide conformation was used.\textsuperscript{28} Electrostatic interactions were calculated using particle mesh Ewald. Bond lengths were constrained using the LINCS algorithm.\textsuperscript{29} Both the temperature and pressure were maintained close to 300 K and 1 atm, respectively, using the weak coupling algorithm\textsuperscript{30} ($\tau_T=0.1$ ps and $\tau_P=1$ ps). Explicit solvent simulations converge much slower than implicit solvent ones. All explicit solvent simulations were sufficiently long to observe relaxation from the initial extended conformation and continued for a time ranging between 160 ns for S4, GS2, and GS4 and 2.5 ns for GS10. Contact formation times have only been estimated for peptides up to 12 residues long, where simulations at least 10 times longer than the average contact time could be performed. To compare contact formation rates with experimental ones, we estimated the diffusion coefficient of the TIP4P water at 300 K temperature and 1 atm pressure by simulating 1000 molecules for 20 ns; this turns out to be $D_{\text{TIP4P}}=(3.93\pm 0.04) \times 10^{-9} \text{ m}^2/\text{s}$ as compared to $D_{\text{exp}}=(2.3\pm 0.04) \times 10^{-9} \text{ m}^2/\text{s}$; the reciprocal of the ratio ($\sim 1.7$) provides an estimate for the corresponding ratio of viscosities (based on the Stokes–Einstein relation).

**End-to-end distance and contact formation rates**

In the TTET experiment\textsuperscript{7} xanthone was used as donor and 1-naphthyl alanine was used as acceptor. In the simulations, contact formation rates were determined between the C- and N-terminal amino acid for the (GS)\textsubscript{x} and S\textsubscript{x} peptides, whereas in the case of the Y(GS)\textsubscript{x}FSG peptide, the Tyr and the Phe moieties were defined as donor and acceptor, respectively. The contact distance was calculated as the minimum distance between any non-hydrogen side-chain atom of the defined acceptor and donor. In the case of glycine residues, the C\textsuperscript{α} atom was used instead of the side-chain atoms. Contact formation was assumed when the distance between donor and acceptor is less than 4 Å. This takes the side-chain hydrogens and the necessary van der Waals contact for TTET into account. Contact formation rates were calculated as

$$k^{-1} = \int_0^\infty S(t)dt$$

where $S(t)$ is the survival probability of the triplet state (i.e., the probability that the two ends are not in contact). If the decay is single exponential, $S(t)=e^{-t/\tau}$, $k^{-1}=\tau$. The original TTET experiments\textsuperscript{7} used a nanosecond laser flash to excite the triplet state; the relaxation appeared to be single exponential for any time above the dead time of the experiment. A more recent femtosecond laser experiment\textsuperscript{10} showed that, for short times, the relaxation is not single exponential, as predicted by a theory for a simple polymer model.\textsuperscript{32} The distribution of survival times of the triplet state displays a long time exponential decay and a nonexponential short-time decay, which fits well to a power law as the theory predicts. Despite a nonexponential distribution, the average contact formation time is still very close to the time constant extracted from the exponential long-time part of the distribution.

**Results**

Most of the simulations in this work have been performed with a simple, computationally efficient implicit solvent model where the solvation effect is based on the solvent-accessible surface (SASA, see Theory and Models). This model allowed a broad and rigorous exploration of different peptide lengths [from 4 to 52 amino acids for each of the (GS)\textsubscript{x}, S\textsubscript{x}, and Y(GS)\textsubscript{x}FSG sequences]. We used uncapped peptides; simulations with acetylated N-terminus showed that this does not significantly influence the rates or the distributions. For some peptides only, simulations were carried out with three alternative implicit solvent models (GBSW, ACE, and EE1F, see Theory and Models) as well as an explicit solvent representation (OPLS/AA + TIP4P).

**The dynamics of polypeptides**

Figure 1 shows the dependence of the average time (or inverse rate) of contact formation as a function of the number of peptide bonds for the three different sequences. For both an implicit (SASA) and explicit solvent, qualitative agreement between experimentally determined and computed values is observed for (GS)\textsubscript{x} and Y(GS)\textsubscript{x}FSG. Since implicit solvent simulations were carried out at a low friction compared to water, absolute rates differ. At the friction of water (\sim 60 ps\textsuperscript{-1}, empty black circles in Fig. 1), the agreement with the experiment is excellent (the whole friction dependence of the rates for some peptides is shown in Fig. 2). Experimental TTET\textsuperscript{7} contact times fit the equation $\tau=\tau_0+\tau_1N^m$ (solid magenta lines in Fig. 1), with $m=1.72$ for (GS)\textsubscript{x}. The exponent $m$ characterises the asymptotic behaviour for long chains and is 1.5 for
an ideal freely jointed chain, larger values are expected when excluded volume effects are present. From SASA simulations, we obtain $m=2.0$ for $(GS)_x$ and $m=1.7$ for $Y(GS)_x FSG$; these values are close to the experimental estimation although both are potentially subject to error because much longer peptides cannot be studied either in vitro due to limitations in peptide synthesis or in silico due to the required computation time, which is exponential in the peptide length.

The value $\tau_0$ represents the limiting contact formation time for short peptides. For $(GS)_x$ and $Y (GS)_x FSG$, the fits give 0.07 and 0.1 ns, respectively; corrected by a factor of 60, this gives a value of $\tau_0 \sim 4.2-6$ ns as compared with the experimental value of $\tau_0 = 5.6$ ns. For a few peptides, we performed simulations at water-like friction and direct agreement for $\tau_0$ is found between simulation and experiment; for $Y(GS)_x FSG$, we obtain a contact formation time of $7 \pm 1$ ns (see Fig. 2) in agreement with the experiment for $(GS)_x$. Overall, for peptides containing the GS repeat, agreement between contact formation times estimated by implicit solvent simulation and experiment is excellent. In explicit solvent, while fewer peptides were studied, the agreement with the experiment is good when
corrected for the lower viscosity of the TIP4P water model relative to real water (by a factor of $\sim 1.7$, see Theory and Models).

The case of $S_x$ is different. From TTET experiments, a very similar behaviour was found to that of the (GS)$_{x}$ series, that is, a similar functional dependence with $m=2.1$ and $\tau_0=12$ ns, reflecting a slight increase in rigidity. However, in simulations of $S_x$ in implicit solvent (at least with the SASA model), we obtain $m=4.8$, which reveals an unusually high rigidity, due to artefactual formation of persistent structure in the simulations (discussed further below). This behaviour is not observed for $S_x$ in explicit solvent where the computed values are always close to the experimental ones, at least for peptides of lengths up to 12 residues (the maximum we simulated with explicit solvent).

Figure 1 also shows the result for a simpler model of the peptides in which only covalent bonds and steric interactions are retained (ORI). For large $N$, we observe the expected power-law behaviour with exponents $m=1.7$ for (GS)$_x$ and Y(GS)$_{FSG}$ and $m=1.9$ for S$_x$; for small $N$, no plateau is evident for short peptides, suggesting that the rigidity of short peptides depends, in part, on attractive interactions; this is in line with the experimental observation that in 8 M GdmCl, the onset of length-independent rate occurs at shorter distances, indicating decreased chain stiffness in 8 M GdmCl compared to water.

Thus, both the overall experimental kinetics of contact formation and the small increased rigidity of $S_x$ relative to (GS)$_x$ can be reproduced well by a simple steric model; indeed, the latter, for $S_x$, performs much better than an implicit solvent model such as SASA.

Figure 1 also shows, as green crosses, fluorescence lifetimes for DBO/Trp-containing peptides determined by Nau et al.,$^{11,12}$ these are about 2.5 times longer than both those measured by TTET and calculated (at water-like friction) in the present simulations. Interestingly, Nau et al. observed an increase in contact formation time for the shortest peptide studied$^{11,12}$ as predicted by the theory.$^{34}$ Ignoring the shortest peptide, the looping time scales as $\tau = \tau_0 + \tau_1 N^m$, where $m=1$; such value may reflect an important role of hydrodynamic interactions, which are neglected in our implicit solvent simulations and which have been shown by theory and simulation to reduce the exponent relative to the ideal case.$^{35}$ However, given the similarity of such times with those determined by TTET and simulation (see Fig. 1), it seems more likely that the problem is that small errors in the times for short peptides affect enormously the asymptotic behaviour for long ones.

**End-to-end distance distributions**

Unlike in experiments, end-to-end distance distributions are readily available from the simulations. This provides information on the conformational properties of the different peptides under simulation, including the differences between force fields, and helps to clarify the discrepancy between the experimentally derived and computed contact formation rates for $S_x$ observed with an implicit solvent model, as mentioned above. The distributions of the end-to-end distances for (GS)$_{x}$, Y(GS)$_{FSG}$, and $S_x$ for various different lengths are shown in Fig. 3 for three different implicit solvents, explicit solvent, and ORI. Some general tendencies emerge. For the pure steric model (ORI), a skewed Gaussian distribution, $p(r) = c \exp(-b(r-a)^2)$ with $a=AN^m$ where $A=5.3$ and $\alpha=0.44$ and with $b=BN^{1.5}$ where $B=0.41$ and $\beta=1.5$ [N is the number of peptide bonds and $c= (a,b)$ a normalisation constant], is observed for all peptide sequences and length. This type of end-to-end distribution has often been assumed to underlie experimental data.$^{16-38}$ However, a clear deviation from the skewed Gaussian distribution is observed for all more realistic force fields. In shorter peptides, the distributions show several peaks and are rather similar for different models due to the intrinsic stiffness of the peptidic chain as the determining factor. For longer peptides, all implicit solvent representations favour more compact conformations than the explicit solvent model. A significant peak at the donor–acceptor contact distance is found for the Y(GS)$_{FSG}$ peptide with all force fields. This reflects hydrophobic interactions of the aromatic donor–acceptor pair. The area under the first peak, that is, the fraction of peptides with donor and acceptor in contact, decreases with the peptide length, and for (GS)$_{16}$ it is about 15%, in agreement with that found experimentally, $\sim 10%$.$^{7}$

In some cases, we observed a pronounced difference between random-coil behaviour and computed distance distributions. In the case of the GBSW force field and for all sequences, the difference can likely be attributed to an overestimation of electrostatic interactions; for example, for $S_x$, the peak at $\sim 5.4$ Å is due to structures where N-terminal NH$_3^+$ and the C-terminal COO$^-$ groups are in contact; the peak at $\sim 10.4$ Å is due to structures where COO$^-$ is in contact to backbone NH amide groups or NH$_3^+$ in contact with CO groups or Ser oxygen atoms. The peculiar behaviour observed for $S_x$ with the SASA implicit solvent is due to a strong helical propensity, as revealed by a peak at end-to-end distances $\sim 3 \times 1.4$ Å.

**Structure formation**

To rationalise the differences in the distance distribution for different force fields, solvent representation, and different peptides on a structural basis, we also analysed hydrogen bonds and secondary-structure formation.

In all cases, the average number of intramolecular hydrogen bonds present depends linearly on the peptide length (Fig. 4). Nevertheless, the slope differs for different models. For all implicit solvation models (SASA, GBSW, ACE, and EEF1), the number of hydrogen bonds increases by $\sim 0.35$ with each residue. In Fig. 4, the result is shown for (GS)$_{x}$. For $S_x$ (not shown), ACE and GBSW display the
same dependence as for \((\text{GS})_x\), while with SASA, the proportionality factor turns out to be \(\sim 0.8\), consistent with the overestimation of the helical propensity mentioned above. In the case of explicit solvent, the number of intrapeptide hydrogen bonds increases much more weakly, at a rate of 0.18 hydrogen bonds per residue. In summary, all implicit solvent models used here seem to overestimate the intramolecular hydrogen-bonding propensity relative to explicit solvent simulations.

The secondary-structure analysis provides additional insight into the differences between force fields. With the explicit solvent model, the residual secondary structure is low but not absent [e.g., for \((\text{GS})_{11}\), about 4% of the residues are in either a helical or an extended conformation, according to DSSP]. Implicit solvent models suggest instead that there is more secondary structure: GBSW and ACE give predictions only slightly larger than the explicit solvent; SASA gives, also taking \((\text{GS})_{11}\) as example, a helical and extended propensity of 17% and 15%, respectively; while the helical propensity decreases with the length, that of extended structure increases. For \(S_x\), the implicit solvents GBSW and ACE and the explicit solvent provide secondary structural propensity very similar to \((\text{GS})_x\), while SASA predicts a
helical propensity of 87% for S_{1,3}, increasing for longer peptides and again indicating that artefactual helical structure is the source of the poor agreement with experimental data for this force field and peptide sequence.

**Radius of gyration**

The most unambiguous measurable characteristic of the state of a disordered polypeptide chain is its radius of gyration. This is shown for (GS) \(_x\) peptides with different force fields in Fig. 5. The radius of gyration has a power-law dependence on the number of residues for all models, while the exponent depends on the molecular model.

This situation mirrors the experimental estimation of compactness of disordered peptides and proteins. Kohn et al.\(^{40}\) recently analysed small-angle x-ray scattering data for 19 proteins spanning a broad range of size and found that under strongly denaturing conditions, the radius of gyration scales as \(R_g = R_g^0 N^\nu\), where \(\nu = 0.598\), in good agreement with the analytical result for a random coil with excluded volume effect (\(\nu = 0.588\)).\(^{41}\) In our simulations, the purely steric (ORI) model leads to \(\nu = 0.68\), slightly larger than the recent experimental estimations but identical with the value reported by Tanford et al. using intrinsic viscosity measurement for 12 proteins.\(^{42}\) All implicit solvents give a considerably lower exponent (\(\nu \sim 0.31-0.37\)). This value is not far from the experimental value found for native proteins\(^{43}\) and highlights overcompaction of unstructured peptides in implicit solvent. For explicit solvent simulations, we found an exponent \(\nu \sim 0.5\), which is in agreement with that experimentally observed for natively unfolded proteins under native conditions.\(^{43}\)

While small-angle x-ray scattering data on the peptides studied here are not available, the root-mean-square end-to-end distance for (GS)\(_{16}\) has been recently determined by FRET\(^{38}\) to be 18.9 Å in water and 39.2 Å under highly denaturing conditions (8 M GdmCl). Simulations with all implicit solvents provide values that are lower than the experimental value in native conditions (4.2 Å for GBSW, 12.1 Å for SASA, 14.5 Å for ACE, and 12.6 Å for EEF1). Simulation with a purely steric potential (ORI) gives a value (39.1 Å) that is close to the high denaturant one. In explicit solvent, we found 21.9 ± 3.4 Å (averaged over the last 2.5 ns of a 6.5-ns simulation). Thus, all implicit solvents estimate the two ends of the peptide to be closer on average than for the peptide in pure water. Conversely, a pure steric model gives a value close to that observed at high denaturant concentration. Explicit solvent gives a value intermediate between the two.

**Discussion**

The unprecedented resolution of recent measurements of the rate of loop formation in peptides has added important elements to our knowledge of the dynamical properties of disordered polypeptide systems. Here, we have shown how simulation is instrumental in interpreting and provoking such experiments. Implicit solvent simulations reproduce well the experimental rates of contact formation for some sequences. Nevertheless, comparison with explicit solvent simulations shows that a general trend of implicit solvent models is to overestimate the compactness, the degree of secondary structure, and the number of hydrogen bonds present. For some peptides, despite these artefacts, we were still able to obtain good estimates of the rates of contact formation with implicit solvents, explicit solvent, and even with the purely steric model. In a pioneering simulation study, Yeh and Hummer also observed that differences in distribution of the end-to-end distance for the pentapeptide CAGQW given by two different explicit solvents (AMBER and CHARMM) were also compatible with identical rates of contact formation.\(^{14}\) One possible reason is that contact formation is an activated process and
that the height of the main free-energy barrier is similar for the different force fields. However, this is certainly not the case for the steric model whose end-to-end distribution, a skewed Gaussian, shows no barriers between contact and noncontact and which itself provides a surprisingly accurate prediction of experimental rates of contact formation. This suggests that excluded volume and chain flexibility may be the main determinant of the kinetics of contact formation. Attractive or long-range interactions only have an effect on the rates of contact formation when they induce a state (e.g., the helical state for 5a seen above) segregated by a sizable free-energy barrier from all the other conformations.

Similarly, the effect of the hydrophobic residues Tyr and Phe (included here to model the triplet donor and acceptor) is negligible on the rates of contact formation but considerable on the distribution of end-to-end distance independent of the underlying force field. While a peak corresponding to conformations where the two ends are in contact is always observed, the peak is considerably larger when the hydrophobic groups are present. The power-law relationship between the radius of gyration and the peptide length, with an exponent close to that of a random coil with excluded volume and close to that determined experimentally for proteins in strong denaturing conditions,40 is reproduced only by a purely sterically model. This suggests that ignoring all interactions except steric ones may represent proteins at highly denaturing conditions better than more sophisticated models of unstructured peptides.

Polypeptides that do not fold in native conditions are particularly interesting because they share properties with natively unfolded proteins. Natively unfolded proteins have an important biological role in phenomena such as transcription, signalling, and regulation;43,44,45 they constitute an important class of proteins, whose conformational properties can help elucidate much of what has not yet been understood about how structured proteins fold. The analysis of the hydrodynamic radius of natively unfolded proteins43 showed that the radius of gyration has a power-law dependence on the number of residues; interestingly, proteins divide into two groups: one, identified as "coil-like," is characterised by an exponent ~0.5, identical with our findings for explicit water. Intriguingly, for a second group, identified as "pre-molten-globule-like," the exponent is 0.41, closer to our results for all implicit solvents. A difference in hydrogen bond formation properties observed in the simulations with either implicit or explicit solvents may explain the origin of different groups of natively disordered proteins.

In conclusion, we found that the properties of naturally unstructured peptides calculated in computer simulations are sensitive to variations in the chosen force field; with implicit solvation models, the peptides are very compact, contain a considerable number of (generally short lived) hydrogen bonds, and are mostly unstructured. Interestingly, for a number of implicit solvents, such "odd" features do not seem to be an obstacle to the simulation of the folding of peptides or miniproteins to their experimental structure.15,46,47 The scaling behaviour of the radius of gyration observed experimentally for structured proteins at high denaturant concentrations is reproduced only by a model in which all attractive interactions are switched off, while all more realistic models are far off. On the other hand, while scaling of the radius of gyration with implicit and explicit solvents is quite different, both behaviours have been observed for natively unfolded proteins in native conditions. Thus, while it is difficult to decide which model is most appropriate, it is certainly interesting to observe that all the models provide results that are within the spectrum of the experimental observations.

Apart from one exception, we observe highly disordered conformations with little or no persistent structure; however, this is not necessarily due to a Gaussian end-to-end distance distribution or to an absence of secondary structure and hydrogen bonds.48

We note that all force fields utilised here have been optimised to accurately predict experimental structures for naturally folded proteins, yet they give markedly divergent results for unfolded peptides except in the case of end-to-end contact formation rate. This suggests that the ability of peptides under simulation to explore the accessible conformation space and fold to a specific structure may be relatively independent of their global behaviour in the denatured state (which we have shown, in this study, to be model dependent in some important aspects) but strongly dependent on rates of contact formation, in keeping with a model of the folding process dominated by diffusion and bifurcations in the folding pathway induced by closure of loops with resulting constraints on accessible conformation space. Finally, we observe that the different representation of the denatured state between models may correspond to different conditions in which proteins happen to fold, either in vitro or in vivo.

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References


Influence of the Internal Disulfide Bridge on the Folding Pathway of the $C_L$ Antibody Domain

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Disulfide bridges are one of the most important factors stabilizing the native structure of a protein. Whereas the basis for their stabilizing effect is well understood, their role in a protein folding reaction still seems to require further attention. We used the constant domain of the antibody light chain ($C_L$), a representative of the ubiquitous immunoglobulin (Ig)-superfamily, to delineate the kinetic role of its single buried disulfide bridge. Independent of its redox state, the monomeric $C_L$ domain adopts a typical Ig-fold under native conditions and does not retain significant structural elements when unfolded. Interestingly, its folding pathway is strongly influenced by the disulfide bridge. The more stable oxidized protein folds via a highly structured on-pathway intermediate, whereas the destabilized reduced protein populates a misfolded off-pathway species on its way to the native state. In both cases, the formation of the intermediate species is shown to be independent of the isomerization state of the Tyr141-Pro142 bond. Our results demonstrate that the internal disulfide bridge in an antibody domain restricts the folding pathway by bringing residues of the folding nucleus into proximity thus facilitating the way to the native state.

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Keywords: antibody folding; disulfide bridges; folding intermediates; $C_L$; NMR

Introduction

To maintain the integrity of secreted proteins, internal stabilizing factors are of outstanding importance. The most prominent example is the disulfide bridge. Disulfide bonds are able to covalently link amino acids far apart in the primary sequence. The resulting conformational restriction of the unfolded state destabilizes it and thus increases the free energy difference to the native state.1,2 In practice, however, this desirable effect is sometimes foiled by strain or reduced dynamics imposed on the native state, especially in the case of artificially introduced disulfide bonds.3-5 Even though disulfide bridges are one of the best understood factors stabilizing the native structure of a protein,6-8 their kinetic role in protein folding is not comprehensively understood. Disulfide bonds can lead to the persistence of partially structured or at least conformationally restricted regions in otherwise unfolded polypeptides.9,10 These pre-formed tertiary interactions may have a profound influence on the kinetics of a protein folding reaction.11-14 Accordingly, genetically engineered disulfide bonds have been used successfully as a tool to probe nucleation sites of the protein folding process.5,15-17 Additionally, a number of studies dealt with the formation of disulfide bridges within the folding process.18-22 However, the influence of naturally occurring disulfide bridges on the mechanism of a protein folding reaction is still not completely understood even though theoretical considerations suggest it to be of pronounced importance.23 To address this issue experimentally, the constant domain of the antibody light chain ($C_L$) of the murine monoclonal antibody MAK33 of the subtype $\kappa/IgG1$ was chosen.24 Antibody domains represent a well suited model system, since their Greek-key

Abbreviations used: ANS, 8-anilino-1-naphtalene sulfonic acid; $C_L$, constant domain of the antibody light chain; CD, circular dichroism; DJ, double jump; EDTA, ethylene diamine tetra-acetic acid; GdmCl, guanidinium chloride; Ig, immunoglobulin; MAK33, monoclonal antibody from mouse of the subtype $\kappa/IgG1$; NOESY, nuclear Overhauser enhancement spectroscopy.

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β-barrel topology is stabilized by a single internal disulfide bond located in the hydrophobic core bridging the two β-sheets. The folding of this domain structure has already been investigated, in most cases with intact disulfide bridges or in naturally disulfide-deficient model systems. A unifying feature for all the different immunoglobulin (Ig) proteins analyzed in this respect so far appears to be the formation of a folding nucleus which is mainly established by hydrophobic residues located in the inner part of the β-sheets. These residues are part of the hydrophobic core in the native state. Details of the folding process vary for the different Ig proteins studied. In some cases, two-state folding behavior has been reported; in others, folding via a highly structured intermediate or a misfolded species was assumed. These differences do of course not rule out a conserved folding mechanism. They might rather represent variations of a motif due to sequence differences which influence the stabilities of different populated species. For the C1 domain, it was shown that folding in the presence and in the absence of its disulfide bridge leads to an identical structure. Thus, the C1 domain is a well suited model system to delineate the kinetic role of the disulfide bridge in the structure formation process of the Ig-topology in antibodies.

Results

Structural characterization of the oxidized and reduced C1 domain in its native and unfolded state

All members of the immunoglobulin superfamily are characterized by the Greek-key β-barrel topology. The recombinant C1 domain shows the typical far-UV circular dichroism (CD) spectrum of an all-β protein characterized by a minimum at 218 nm and a low overall intensity (Figure 1(a)). Importantly, the far-UV CD spectrum, and thus the secondary structure of the C1 domain, does not change significantly upon reduction of its internal disulfide bridge (Figure 1(a)). Furthermore, near-UV CD and 1D 1H NMR/2D 1H-1H NOESY spectra of the reduced and oxidized C1 domain show that the structure of the domain remains essentially unchanged upon reduction of its disulfide bridge (Figure 1(b) and (c) and data not shown).

The presence of a covalent cross-link bridging 60 residues influences the conformational freedom of the unfolded chain. To investigate whether significant structural elements persist in the unfolded state
of the oxidized CL domain, compared to its reduced counterpart, the spectroscopic techniques used to characterize the native state were also applied to the denatured state of the oxidized and reduced protein. The far-UV CD spectra of both forms are very similar and typical for random coil polypeptides (Figure 1(a)). Near-UV CD spectra clearly show the absence of significant asymmetric environment around the aromatic residues for the reduced and the oxidized CL domain (Figure 1(b)). The small and broad signal in the latter case can likely be attributed to the intrinsically asymmetric disulfide bridge. NMR spectra also argue against significant differences between the unfolded states. In the presence of 2 M guanidinium chloride (GdmCl), both forms show no significant signal dispersion in the 1D 1H NMR spectrum (data not shown). Additionally, iodide fluorescence quenching experiments were carried out to investigate the solvent accessibility of the two tryptophanes in the unfolded states of the oxidized and reduced CL domain. Solvent accessibility of tryptophane residues was observed to be slightly higher in the reduced protein (data not shown). Upon reduction, the Stern-Volmer constant $K_{SV}$ increases from 4.6 M$^{-1}$ to 4.9 M$^{-1}$. In the native state, no quenching difference for the oxidized and the reduced CL domain is observed (data not shown).

The hydrodynamic properties of the CL domain are influenced by the disulfide bridge as expected. Pulsed field gradient diffusion NMR experiments of the native proteins show a decrease in the diffusion coefficient of $\sim$10% when the disulfide bridge is reduced, i.e. the reduced form is more extended. For the native state, the calculated diffusion constant using the pdb structure file (pdb code 1FH5) containing oxidized CL is $1.083 \times 10^{-10}$ m$^2$ s$^{-1}$ which is in good agreement with the measured value of $1.255 \times 10^{-10}$ m$^2$ s$^{-1}$. In the presence of 2 M GdmCl, the diffusion constant decreases $\sim$tenfold as compared to the native state (data not shown). Using the Debye-Einstein equation, the hydrodynamic radius of the different CL species can be calculated assuming a spherical shape for all proteins. Upon reduction, the hydrodynamic radius increases from 1.46 nm to 1.60 nm. These data emphasize the role of the disulfide bridge for the compact structure of the folded CL domain. In the presence of 2 M GdmCl, the increase in the hydrodynamic radius is around 40% for both the oxidized and the reduced form showing that the presence of the disulfide bond does not significantly restrict the unfolded state of the CL domain.

**Stabilization of the CL domain by its internal disulfide bridge**

Equilibrium unfolding experiments were carried out to determine how the internal disulfide bridge contributes to the stability of the CL domain. Unfolding was monitored either by changes in the far-UV CD signal or by changes in the intrinsic tryptophane fluorescence. Two tryptophanes are found in the CL domain, one is located in strand $C$ and is part of the hydrophobic core of the Ig-fold. The other one is located in strand D and is rather solvent-exposed. As can be seen in Figure 2(a), the disulfide bridge is a major stabilizing factor of the CL domain. In its absence, the midpoint of the GdmCl transition is shifted from 0.8 M GdmCl to 0.3 M GdmCl. This corresponds to a reduction in stability of 8.2 kJ mol$^{-1}$ from 14.6 kJ mol$^{-1}$ in the oxidized form to 6.4 kJ mol$^{-1}$ in the reduced form. The cooperativity parameter $m$ of the transition is found to be 17.4 kJ M$^{-1}$ mol$^{-1}$ in the oxidized and 20.9 kJ M$^{-1}$ mol$^{-1}$ in the reduced state. Since the respective far-UV CD and fluorescence transitions of the reduced and oxidized domain coincide, significantly populated equilibrium unfolding intermediates are very unlikely. Independent of the redox state, the temperature-induced unfolding of the CL domain was not completely reversible. Nevertheless, the major stabilizing effect of the disulfide bond is obvious from the decrease in the melting point by 20 °C, from 51 °C to 31 °C upon reduction (Figure 2(b)).

![Figure 2](image.png)
Folding mechanism of the C₅ domain

To investigate the role of the disulfide bridge in the folding process of the C₅ domain, the kinetics of folding and unfolding of the reduced and the oxidized protein were monitored at different GdmCl concentrations by fluorescence spectroscopy. The experiments were performed in standard buffer at 20 °C. In all cases, a small signal loss was observed during the dead time of the stopped-flow experiments which can likely be attributed to the reorganisation of the polypeptide chain upon change of the solvent conditions. From the chevron plots of the respective rate constants for the oxidized and the reduced protein, it is evident that the folding reaction consists of three observable, kinetically well separated phases (Figure 3 and Table 1). The disulfide bridge has a pronounced effect on the

Table 1. Microscopic rate constants and kinetic m-values derived from the chevron plots for the oxidized and the reduced C₅ domain

<table>
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<tr>
<th>Parameter</th>
<th>Value for oxidized C₅</th>
<th>Value for reduced C₅</th>
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<tr>
<td>kO cis−→ N [s⁻¹]</td>
<td>0.00068</td>
<td></td>
</tr>
<tr>
<td>(mO cis−→ N [M⁻¹])</td>
<td>(1.48)</td>
<td></td>
</tr>
<tr>
<td>kI cis.trans−→ cis/trans [s⁻¹]</td>
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<td></td>
</tr>
<tr>
<td>(mI cis/trans−→ cis/trans [M⁻¹])</td>
<td>(0.0)</td>
<td></td>
</tr>
<tr>
<td>kI cis−→ trans [s⁻¹]</td>
<td>0.011</td>
<td>(0.0)</td>
</tr>
<tr>
<td>(mI cis−→ trans [M⁻¹])</td>
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<td></td>
</tr>
<tr>
<td>kU cis−→ trans [s⁻¹]</td>
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<td></td>
</tr>
<tr>
<td>(mU cis−→ trans [M⁻¹])</td>
<td>(−1.80)</td>
<td></td>
</tr>
<tr>
<td>kO cis−→ trans [s⁻¹]</td>
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<td></td>
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<tr>
<td>(mO cis−→ trans [M⁻¹])</td>
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</tr>
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<td></td>
</tr>
<tr>
<td>(mI trans−→ cis [M⁻¹])</td>
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</tr>
<tr>
<td>kO trans−→ cis [s⁻¹]</td>
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<td></td>
</tr>
<tr>
<td>(mO trans−→ cis [M⁻¹])</td>
<td>(0.0)</td>
<td></td>
</tr>
</tbody>
</table>

Data were obtained from a fit of the chevron plot to a three-state on-pathway model for the oxidized C₅ domain or a three-state off-pathway model for the reduced C₅ domain (see Figure 6). Since the intermediate states build up independent of the Tyr₄₄−Pro₄₂ bond isomerization state, the kinetic parameters for their formation were regarded as independent of this reaction.

Figure 3. Chevron plots of the oxidized and reduced C₅ domain. Unfolding and refolding of the oxidized (a) or reduced (b) C₅ domain at different GdmCl concentrations were measured. The fast phase (open circles) was monitored by stopped-flow fluorescence spectroscopy and the slowest phase (filled circles) was monitored by both techniques. Additionally, rate constants obtained from DJ experiments are shown (crosses). The data obtained for the oxidized C₅ domain were fit to a three-state on-pathway model, whereas the data for the reduced protein were best described by a three-state off-pathway model (continuous lines). The slowest phase was fit to a Pro-isomerization reaction in both cases. The rate constants and kinetic m-values for the formation of the intermediate species were assumed to be independent of the isomerization state of the Tyr₄₄−Pro₄₂ bond.

fast and intermediate phase observed. If reduced, the refolding limb of the intermediate phase not only shows a rollover but passes through a maximum (Figure 3(b)). This shape of the chevron plot makes the formation of a misfolded species within the fast phase likely which has to undergo unfolding before productive refolding reactions can occur. If unfolding of this species becomes rate limiting, as observed for the reduced C₅ domain at low GdmCl concentrations, an apparent deceleration of the productive folding reactions is expected which explains the shape of the intermediate phase in the chevron plot. For the oxidized domain, on the other hand, a linear GdmCl-dependency of the fast and intermediate phase is observed (Figure 3(a)). This implies that the reactions occurring in the fast phase are significantly different for the oxidized and the reduced C₅ domain. To further analyze the underlying processes, interrupted refolding experiments and double jump (DJ) experiments were carried out. In interrupted refolding experiments, the protein is refolded for different times before changing to unfolding conditions again. The observed amplitudes of the different unfolding phases directly correspond to the amount of different species present. In DJ experiments, slow isomerization reactions
in the unfolded state are indirectly monitored by the amount of molecules following different pathways after varying denaturation times. These experiments allowed us to monitor the formation of intermediates and native molecules as well as slow isomerization reactions giving rise to the different observable phases. As can be seen in Figure 4(a) and Table 2, two exponential phases are necessary to describe the formation of native molecules in the case of the oxidized C_L domain in interrupted refolding experiments. Hence, refolding of the oxidized protein occurs on two parallel pathways. The slower one accounts for ~90% of the molecules after extended denaturation times (Figure 4(a) and (c) and Table 2) indicative of a proline-isomerization reaction. Only one cis peptide bond, between Tyr141 and Pro142, is found in the native state of the C_L domain. After equilibration, ~80 to 90% of the molecules are expected to populate the non-native trans conformation in the unfolded state which corresponds very well to the experimentally observed amount of slow folding species (Table 2). Taking the rate constant for refolding and the amount of slow folding molecules into account, one of the parallel pathways can likely be attributed to the isomerization of the Tyr141-Pro142 bond to the native cis state. In the case of the oxidized C_L domain, an obligatory on-pathway folding intermediate seems to be populated in the fast phase as inferred from the lag time for the formation of native molecules (Figure 4(a), inset). The large extent of secondary structure, native fluorescence, and NMR signals which are already observed after the fast refolding

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value for oxidized C_L</th>
<th>Value for reduced C_L</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k_{I \text{cis} \rightarrow N}$ [s$^{-1}$]</td>
<td>0.069 (8%)</td>
<td>–</td>
</tr>
<tr>
<td>$k_{I \text{trans} \rightarrow N}$ [s$^{-1}$]</td>
<td>0.0014 (92%)</td>
<td>–</td>
</tr>
<tr>
<td>$k_{I \text{cis} \rightarrow I \text{trans}}$ [s$^{-1}$]</td>
<td>–</td>
<td>0.051 (14%)</td>
</tr>
<tr>
<td>$k_{I \text{trans} \rightarrow I \text{cis}}$ [s$^{-1}$]</td>
<td>–</td>
<td>0.0013 (86%)</td>
</tr>
</tbody>
</table>

The unfolding amplitudes obtained in the interrupted refolding assay were either fit to an irreversible three-state on-pathway model (oxidized C_L, equation (7)) or an irreversible two-state model (reduced C_L, equation (8)) with two parallel pathways in each case. The amount of molecules following a certain pathway is denoted in brackets. Values for the oxidized domain were determined in the presence of 140 mM residual GdmCl, values for the reduced domain in the presence of 90 mM residual GdmCl.
phase also implies an on-pathway folding intermediate (Figures 4(c) and 5(a) and (d); far-UV CD data not shown). The persistence of the fast refolding phase after short- or long-term denaturation in the DJ experiments (Figure 4(c)) shows that this intermediate species is formed independent of the isomerization state of the proline residues. This finding is in agreement with the results from the interrupted refolding assay, where the decay of the intermediate species occurs in a slow and a fast reaction. The rate constants for the decay of the intermediate species are in good agreement with the ones for the formation of native molecules. In contrast to these results, an intermediate trapped by a trans Tyr141-Pro142 bond would only be populated after prolonged denaturation times, i.e. the amplitude of the fast phase would increase in DJ experiments, and would only decay with a single, isomerization limited reaction in the interrupted refolding assay.

For the reduced C1 domain, the interrupted refolding assay also shows that the protein folds on two parallel pathways. Again, ~90% of the molecules follow the slow pathway after extended denaturation times with a similar rate constant as in the case of the oxidized domain (Figure 4(b) and Table 2). Accordingly, the isomerization of the Tyr141-Pro142 bond to the native cis state also slows down the folding of a major part of the reduced C1 domain as expected. The chevron plot for the reduced C1 domain exhibits three observable phases (Figure 3(b)), but in interrupted refolding experiments, the formation and decay of the misfolded species cannot be observed directly in the interrupted refolding assay due to its fast unfolding reaction (Figures 3(b) and 4(b)). Nevertheless, the presence of the fast refolding phase in DJ experiments, independent of the denaturation time and therefore the Pro-isomerization state, clearly demonstrates that the misfolded intermediate is populated on the slow and on the fast folding pathway (Figure 4(d)). The absence of a lag phase for the formation of native, reduced molecules strengthens the notion that the species formed is misfolded (Figure 4(b), inset). Furthermore, no

![Figure 5](image)

**Figure 5.** One-dimensional real-time NMR spectroscopy of the refolding of reduced (a) and oxidized (b) C1. Refolding of the reduced domain was carried out in the presence of 60 mM GdmCl, the oxidized C1 domain was refolded in the presence of 100 mM GdmCl. The intensity of the methyl resonances marked by an arrow is plotted against refolding time in (c) and fitted to a single exponential ($k_{\text{ox}} = 0.00062 \text{ s}^{-1}$ (filled circles, oxidized protein) and $k_{\text{red}} = 0.0015 \text{ s}^{-1}$ (open circles, reduced protein)). The first 1D spectrum of each series is shown in (d) (upper panel: reduced C1; lower panel: oxidized C1).
significant extent of native far-UV CD signal, native fluorescence, or NMR signals build up during the fast folding phase of the reduced \( \text{C}_\lambda \) domain (Figures 4(d) and 5(b) and (d); far-UV CD data not shown).

**Structural characterization of the different intermediates**

The structural properties of the intermediate states formed during the folding of the oxidized and the reduced \( \text{C}_\lambda \) domain were investigated by different spectroscopic techniques. Formation of secondary structure was monitored by far-UV CD spectroscopy. Spectra of the refolding protein were measured after 1 min incubation time within 2 min. In this period, the intermediate states have already formed but the majority of the molecules are still separated from the following folding reaction to the native state by the slow isomerization of the Tyr141-Pro142 bond. In the case of the oxidized protein, a large amount of the native secondary structure already forms in the 3 min experiment takes whereas for the reduced protein, the spectrum after 3 min is still much more random coil-like (data not shown). 8-Anilino-1-naphthalene sulfonic acid (ANS) binding to the two intermediate states was not significantly different. Only a slightly higher ANS affinity was observed in the case of the oxidized protein showing that a few more hydrophobic residues are already buried in the intermediate state of the reduced protein (data not shown). To gain further insight into the similarities and differences of the intermediate states populated, real-time 1D \(^1\text{H}\) NMR experiments were recorded (Figure 5). The reduced and oxidized \( \text{C}_\lambda \) domains refold in a similar timescale (Figure 5(a)-(c)), but the initial states are significantly different (Figure 5(d)). After the ~5 min dead time of the mixing and first data acquisition step, the fast refolding phase is finished and the folding intermediate is populated. The spectrum of the oxidized form already shows a well dispersed \(^1\text{H}\) spectrum which is indicative of a compact tertiary structure whereas the spectrum of the reduced form still shows poor dispersion. Therefore, this intermediate seems not to be compactly folded (Figure 5(d)). It should be noted that during the 5 min dead time of the experiment, ~20% of the molecules already fold to the native state due to amount of correct Pro-isomers present and the additionally occurring re-isomerization to the native conformer during that time. This likely explains the small amount of native methyl signals observed for the reduced \( \text{C}_\lambda \) domain in the first spectrum but not the pronounced resonances observed for the oxidized protein.

**Discussion**

In the present study, we have analyzed the influence of the internal disulfide bridge on the folding mechanism of the \( \text{C}_\lambda \) antibody domain. This protein folds into an almost identical structure independent of its redox state. Therefore, it is a well suited model system to delineate the kinetic role of the disulfide bond in the folding of a representative of the immunoglobulin superfamily. This advantageous feature has already been recognized by Goto et al. who carried out groundbreaking studies on the stabilizing role of the internal disulfide bridge\(^{39}\) and on its kinetic impact using a human type \( \lambda \) \( \text{C}_\lambda \) domain.\(^{40}\) The persistence of its structural integrity despite reduction is obviously a common feature for the human and the murine \( \text{C}_\lambda \) domain of the \( \kappa \) and \( \lambda \) type.\(^{39} \) The stability of the oxidized \( \text{C}_\lambda \) domain was determined by Goto et al. to be ~21 kJ mol\(^{-1}\) for a human \( \kappa \) \( \text{C}_\lambda \) domain and to be ~24 kJ mol\(^{-1}\) for a human \( \lambda \) \( \text{C}_\lambda \) domain.\(^{39,46}\) The higher values than the ~15 kJ mol\(^{-1}\) determined for the oxidized \( \kappa \) \( \text{C}_\lambda \) domain in this study might be explained in two ways. On the one hand, a higher stability of the human \( \kappa \) and \( \lambda \) \( \text{C}_\lambda \) domains compared to the murine \( \kappa \) \( \text{C}_\lambda \) domain could be assumed. On the other hand, different models have been applied for determining the free energy of unfolding. The binding model of Tanford employed by Goto et al.\(^{47} \) has been shown to overestimate protein stability, especially when GdmCl is used as a denaturant.\(^{48}\) Both considerations might explain the differences in stability observed. The stabilizing effect of the internal disulfide bridge is very pronounced for the human and the murine \( \text{C}_\lambda \) domain. In a simplistic approach, the stabilizing effect of a disulfide bond bridging \( n \) residues can be approximated to\(^{17} \):

\[
\Delta \Delta G = -T \Delta \Delta S = -T (-2.1 - [1.5 \text{Rn}(n)]) \text{ cal mol}^{-1}
\]  

In the case of the \( \text{C}_\lambda \) domain with \( n=60 \), \( \Delta \Delta G \) is calculated to ~17.5 kJ mol\(^{-1}\) which clearly exceeds the stabilizing effect of \( \Delta \Delta G=8.2 \text{ kJ mol}^{-1} \) determined experimentally and even the entire stability of the domain. The above equation only takes the entropic destabilization of the unfolded state into account but neglects possibly reduced dynamics of the native state or induced strain. These factors may lead to a deviation between the experimentally observed and the theoretically expected stabilizing effect of the disulfide bridge. Only in very few cases, the stabilizing role of a disulfide bridge could be adequately predicted by this simple model.\(^3\) In the case of the \( \text{C}_\lambda \) domain, the hydrodynamic radius increases upon reduction by ~10% as derived from pulsed field gradient diffusion NMR experiments. This points towards a certain strain the disulfide bridge imposes on the native state. Furthermore, the \( \text{C}_\lambda \) domain showed increased flexibility upon reduction of its internal disulfide bridge.\(^{49} \) Thus, more sophisticated models for analysis, e.g. including the reduced dynamics of the native state using a molecular dynamics approach, seem to be required.
Disulfide bridges have been shown to preserve a certain degree of residual structure in otherwise unfolded polypeptide chains in some cases. This might of course have a profound influence on the folding dynamics and pathway of a protein. Therefore, the denatured state of the reduced and oxidized C\textsubscript{L} domain was characterized to detect residual structure. The \( m \)-value of denaturant-induced unfolding transitions is known to correlate with the amount of hydrophobic surface exposed upon unfolding. For disulfide-bridged proteins, generally lower \( m \)-values are found than for disulfide-free proteins or their reduced counterparts. This is also the case for the C\textsubscript{L} domain. The \( m \)-value increases by \( \sim 20\% \) from 17.4 kJ M\(^{-1}\) mol\(^{-1}\) to 20.9 kJ M\(^{-1}\) mol\(^{-1}\) upon reduction of the internal disulfide bond. Furthermore, the Stern-Volmer constant for iodide-induced tryptophane quenching increases by \( \sim 7\% \) from 4.6 M\(^{-1}\) to 4.9 M\(^{-1}\) in the unfolded protein. On the other hand, far-UV and near-UV CD spectroscopy as well as 1D \(^1\)H and 2D \(^1\)H-\(^1\)H-NOESY NMR experiments rule out the persistence of significant residual structure in the case of the oxidized C\textsubscript{L} domain after unfolding. In the primary sequence, Cys\textsubscript{135} is flanked by hydrophobic residues (VVC\textsubscript{135}FL) and Cys\textsubscript{195} by residues of alternating hydrophobicity (YTC\textsubscript{195}EA). Importantly, Val\textsubscript{133} and Leu\textsubscript{137} as well as Tyr\textsubscript{193} and Ala\textsubscript{197} are part of the hydrophobic core in the native state. Interaction of these residues might induce a small amount of hydrophobic clustering around the disulfide bridge, even in the otherwise unfolded polypeptide chain. From the fluorescence quenching experiments, which report on a slightly lower tryptophane solvent accessibility for the oxidized and unfolded C\textsubscript{L} domain than for the reduced protein, the intermittent association of one or both tryptophane residues to this cluster seems possible.

Independent of the redox state, the isomerization of the Tyr\textsubscript{141}-Pro\textsubscript{142} bond to the native \( cis \) state is the rate limiting step for the folding of the major part of the C\textsubscript{L} molecules. The presence of the disulfide bridge has only a small effect on the isomerization rate of the Tyr\textsubscript{141}-Pro\textsubscript{142} bond, either from its native \( cis \) to the non-native \( trans \) state upon unfolding or on the reverse reaction upon refolding (see Table 1 and Figure 5(c)). Both the reduced and the oxidized C\textsubscript{L} domains populate an intermediate species upon refolding. This process is independent of their Pro-isomerization states as derived from DJ experiments. In the case of the oxidized C\textsubscript{L} domain, a highly structured on-pathway intermediate is formed, which leads to a lag phase for the formation of native molecules (see Figure 4(a)). It already possesses a large amount of native secondary structure and native-like dispersion of all NMR signals (Figure 5(d)). Furthermore, its stability is rather high. Assuming a pre-equilibrium between the unfolded and the intermediate state and independence of the Pro-isomerization reaction, its stability is calculated to be \( \sim 11 \) kJ mol\(^{-1}\) from the microscopic rate constants (Table 1), which is only \( \sim 25\% \) lower than the stability of the native state. Surprisingly, in this highly structured and stable intermediate, a free isomerization of the Tyr\textsubscript{141}-Pro\textsubscript{142} bond to the native \( cis \) state with a rate similar to that observed for unstructured peptides is possible. The Tyr\textsubscript{141}-Pro\textsubscript{142} bond is found in the loop connecting strands B and C of the immunoglobulin fold. Its free isomerization is in agreement with the proposed small role the loops play as folding nuclei in the structure formation process of the Ig-topology. Furthermore, the presence of the intermediate not only on the folding but also on the unfolding pathway (Figure 3(a)) shows that unfolding of the C\textsubscript{L} domain is likely to begin from the loops and expand to the hydrophobic core. If the internal disulfide bridge of the C\textsubscript{L} domain is reduced, its folding behavior changes significantly. For the intermediate phase of the chevron plot, corresponding to the formation of native molecules from the denatured state with the Tyr\textsubscript{141}-Pro\textsubscript{142} bond in the \( cis \) conformation, a pronounced rollover with the rate constant passing through a maximum is observed (see Figure 3(b)). The same behavior has been reported for the tenth fibronectin type III domain of human fibronectin (FN\textsubscript{n10}) also belonging to the Ig-superfamily. As has been argued by Wildegeger and Kiefhaber, this behavior of the observable rate constants points towards an off-pathway intermediate as part of a dead-end mechanism. This is in agreement with the absence of a lag phase for the formation of native, reduced C\textsubscript{L} molecules and the spectroscopic characteristics of the intermediate species. In contrast to the oxidized protein, in far-UV CD experiments, the intermediate is essentially random coil-like and 1D \(^1\)H NMR experiments also show random coil chemical shift dispersion after the fast refolding phase (see Figure 5). Taken together, the experimental data allow the folding model proposed for the C\textsubscript{L} domain\cite{27,40} to be refined, especially in the case of the reduced C\textsubscript{L} domain (Figure 5). Being very unstable (\( \sim 2 \) kJ mol\(^{-1}\)), the kinetic \( m \)-values for the folding and unfolding of the misfolded species are only poorly defined. Nevertheless, compared to the oxidized domain, the kinetic \( m \)-values are clearly higher and therefore the transition states for the formation and unfolding of the off-pathway intermediate are less compact (see Table 1). ANS-binding experiments on the other hand argue for a pronounced clustering of hydrophobic residues in the misfolded species, even slightly higher than for the on-pathway intermediate in the case of the oxidized C\textsubscript{L} domain. Consequently, in the folding process of reduced C\textsubscript{L}, the population of a misfolded, aggregation-prone off-pathway species can be postulated. Almost completely devoid of native secondary or tertiary structure and mainly stabilized by the collapse of hydrophobic residues, this species has to unfold before productive refolding to the native state can occur.

In summary, our results imply a new role for the disulfide bridge in antibody folding: it facilitates the...
efficient folding to the native state. In the case of the oxidized domain, irrespective of Pro-isomerization, a native-like intermediate is formed within a few hundred ms which then undergoes a second folding reaction to the native state. Importantly, the structured intermediate species shows no aggregation tendency at refolding concentrations up to a few hundred μM (data not shown). Folding of the reduced C_L domain, on the other hand, is slowed tendency at refolding concentrations up to a few hundred μM (data not shown). Folding of the oxidized CL domain runs faster, and loss in fluorescence confirmed by non-reducing SDS-PAGE, where the oxidized domain and reduced CL domain populate a folding intermediate. In the case of the oxidized protein (a), this intermediate is highly structured and on-pathway. The reduced C_L domain, on the other hand, populates a misfolded and unstable off-pathway species which has to undergo unfolding before productive folding reactions can occur (b). Folding of the major part of the molecules is limited by the isomerization of the Tyr141-Pro142 bond to the native cis state. Microscopic rate constants are denoted for the individual reactions. For the oxidized domain, isomerization of the Tyr141-Pro142 bond was assumed to occur in the unfolded state with the same rate constants as in the intermediate state.

Figure 6. Folding mechanism for the C_L domain. Independent of the Tyr141-Pro142 bond isomerization state, the oxidized and the reduced C_L domain populate a folding intermediate. In the case of the oxidized protein (a), this intermediate is highly structured and on-pathway. The reduced C_L domain, on the other hand, populates a misfolded and unstable off-pathway species which has to undergo unfolding before productive folding reactions can occur (b). Folding of the major part of the molecules is limited by the isomerization of the Tyr141-Pro142 bond to the native cis state. Microscopic rate constants are denoted for the individual reactions. For the oxidized domain, isomerization of the Tyr141-Pro142 bond was assumed to occur in the unfolded state with the same rate constants as in the intermediate state.

Materials and Methods

GdmCl (ultrapure) was purchased from MP Biomedicals (Eschwege, Germany). All other chemicals were from Merck (Darmstadt, Germany). The GdmCl concentrations were determined by the refractive indices of the solutions. Unless otherwise stated, all experiments were carried out in 20 mM sodium phosphate (pH 7.5), 100 mM NaCl, 1 mM ethylene diamine tetra-acetic acid (EDTA), 1 mM DTT (if the reduced C_L domain was investigated) (standard buffer) at 20 °C.

Cloning, expression, and purification of the C_L domain

The C_L gene was obtained by PCR amplification using the CDNA of the murine MAK33 κ light chain as a template. The PCR product was cloned into the pET28a(+) expression vector (Novagen, Darmstadt, Germany) via the Ndel and HindIII restriction sites and transformed into the E. coli strain BL21(DE3). Cells were grown at 37 °C in selective LB medium, and expression was started by addition of 1 mM isopropyl-beta-D-thiogalactopyranoside at an OD600 of 0.8. After 3 h, cells were harvested. The preparation of the C_L inclusion bodies was carried out as described previously. Inclusion bodies were solubilized in 100 mM sodium phosphate (pH 7.5), 6 M GdmCl, 20 mM ß-mercaptoethanol for 2 h at 20 °C. Insoluble components were removed by centrifugation (48,000 g, 25 min, 20 °C). The supernatant was diluted fivefold in 50 mM sodium phosphate (pH7.5), 4 M GdmCl and applied to a Ni-chelating column. After washing for five column volumes, elution was performed with the same buffer at pH 4.0. Refolding was carried out via dialysis in 250 mM Tris-HCl (pH 8.0), 5 mM EDTA, 1 mM oxidized glutathione at 4 °C overnight. Aggregates were removed by centrifugation (48,000g, 25 min, 4 °C) and 0.25 units Thrombin (restriction grade, Novagen, Darmstadt, Germany) for each mg protein were added to remove the N-terminal His-tag. The reaction was allowed to proceed for 16 h at 4 °C. After an additional centrifugation step to remove aggregates (48,000g, 25 min, 4 °C), the concentrated supernatant was applied to a Superdex 75 pg 26/60 gel filtration column (Amersham Biosciences, Upsala, Sweden) equilibrated in 20 mM sodium phosphate (pH 7.5), 100 mM NaCl, 1 mM EDTA.

The cloned C_L domain contains residues A112-E214 of the MAK33 κ light chain (labeled according to the crystal structure) and four additional N-terminal residues (GlySerHisMet) derived from the pET28a(+) vector after Thrombin cleavage of the His-tag. All vectors were sequenced and the mass of the purified proteins was controlled by matrix assisted laser desorption ionization-time of flight mass spectrometry.

Preparation of reduced C_L

The native C_L domain was unfolded and reduced in 20 mM sodium phosphate (pH 7.5), 100 mM NaCl, 1 mM EDTA, 2 M GdmCl, 50 mM DTT for 2 h at room temperature. Refolding of the reduced protein was performed overnight at 4 °C in 20 mM sodium phosphate (pH 7.5), 100 mM NaCl, 1 mM EDTA, 1 mM DTT via dialysis. Possibly present aggregates were removed by centrifugation (48,000g, 25 min, 4 °C). Successful reduction was confirmed by non-reducing SDS-PAGE, where the oxidized C_L domain runs faster, and loss in fluorescence quenching. To avoid re-oxidation, the reduced protein was always used for measurements within one day.

Materials and Methods

GdmCl (ultrapure) was purchased from MP Biomedicals (Eschwege, Germany). All other chemicals were from Merck (Darmstadt, Germany). The GdmCl concentrations were determined by the refractive indices of the solutions. Unless otherwise stated, all experiments were carried out in 20 mM sodium phosphate (pH 7.5), 100 mM NaCl, 1 mM ethylene diamine tetra-acetic acid (EDTA), 1 mM DTT (if the reduced C_L domain was investigated) (standard buffer) at 20 °C.
Influence of the Disulfide Bridge on C_{f} Folding

Fluorescence and CD measurements

Fluorescence measurements were carried out in a Spex FluoroMax II fluorimeter (Instruments SA, Edison, NJ). Equilibrium unfolding transitions (incubated for at least 12 h) as well as refolding kinetics were followed by the intrinsic tryptophane fluorescence of C_{f} excited at 280 nm (3 nm slit) and monitored at 370 nm (oxidized C_{f}) or 320 nm (reduced C_{f}), respectively. The emission slits were set to 5 nm. The protein concentration was 1 μM in 1 cm quartz cuvettes.

CD measurements were performed in a Jasco J-720 spectropolarimeter (Jasco, Grossumstadt, Germany). Far-UV spectra were recorded from 200 to 250 nm at a protein concentration of 10 μM in 1 cm quartz cuvettes, whereas for near-UV spectra a protein concentration of 50 μM in 5 mm quartz cuvettes was used. Spectra were accumulated 12 times and buffer-corrected. Equilibrium unfolding transitions and folding kinetics were monitored at 218 nm.

The method described by Santoro and Bolen was used to fit the data.55

The CD spectra of the folding intermediates were measured at a protein concentration of 10 μM in a 1 cm quartz cuvette. Seven individually measured spectra were recorded after a refolding time of 1 min within 2 min and averaged in each case. The same refolding and measuring times were used for ANS-fluorescence spectroscopy, where always four individually measured spectra were averaged. Experiments were performed at a protein concentration of 1 μM and an ANS concentration of 100 μM in stirrable cuvettes. Spectra were excited at 365 nm and recorded from 430 nm to 600 nm (slits as denoted above).

NMR measurements

NMR samples were concentrated up to 250 μM. Spectra were recorded on a Bruker DMX750 spectrometer (Bruker, Rheinstetten, Germany) equipped with a triple resonance probe head at 293 K. For 1D 1H spectra 2k data points were recorded with a spectral width of 12 ppm, zero filled to 4k data points and apodized with a squared sine window function. 2D 1H-1H-nuclear Overhauser enhancement spectra were recorded with a spectral width of 12 ppm, zero filled to 4k complex data points in the indirect dimension. Data were processed with Topspin Version 1.3. 1H real-time NMR experiments were conducted as described by Balbach et al.60 Refolding was started by a dilution of denatured protein samples (in 2 M GdmCl) to a final protein concentration of ~160 μM. Oxidized C_{f} was refolded in the presence of 100 mM GdmCl, whereas for reduced C_{f} the residual GdmCl concentration was 60 mM. Due to manual mixing and succeeding equilibration and optimization of the magnetic field homogeneity, the dead time of the experiment was approximately 5 min. During refolding, 32 experiments were recorded with 128 transients per spectrum. Processing was done with Topspin Version 1.3 (Bruker, Rheinstetten, Germany).

Iodide quenching

The presence of heavy atoms like iodide or bromide or heavy atom containing molecules like alkyl halides is known to quench the intrinsic fluorescence of aromatic or heteroaromatic fluorophores.61 The collision-based dynamic quenching process can be described by the Stern-Volmer equation:

\[
\frac{F_0}{F} = 1 + K_{SV}[Q]
\]

where \(F_0\) denotes the fluorescence intensity in the absence of the quencher and \(F\) the intensity in its presence which is linearly decreasing with the quencher concentration \([Q]\). The Stern-Volmer constant \(K_{SV}\) increases with increasing quencher accessibility of the chromophore. For quenching experiments, tryptophane fluorescence was excited at 295 nm and monitored at 350 nm. The protein concentration was 1 μM. NaI was added from a freshly prepared 5 M stock solution containing 0.1 mM of the anti-oxidant sodium thiosulfate.

Stopped-flow measurements

Rapid-mixing folding experiments were carried out in an Applied Photophysics SX18-MV stopped-flow instrument (Applied Photophysics, Leatherhead, UK). Fluorescence was excited at 280 nm and a 335 nm cutoff filter was used for detection. Dependent on the experiment, the stopped-flow or the sequential flow mode was used with mixing ratios of 1:1, 1:11, or 1:26 and averaging was performed for 1 to 7 traces. The final protein concentration in the 2 mm observation cuvette was always 1 to 2 μM. To monitor the unfolding rate of the intermediate in the case of the reduced C_{f} domain, it was first accumulated in a 15 s refolding pulse at 90 mM GdmCl before unfolding was carried out.

Double jump experiments

DJ experiments are well suited to follow isomerization reactions taking place in the unfolded state of a protein.62 To this end, the protein is denatured for different periods of time before refolding is carried out. The change in the amplitudes of different refolding phases directly corresponds to the amount of different species present in the unfolded state.

The C_{f} domain was unfolded for different times in 2 M GdmCl before refolding in the presence of 50 to 75 mM residual GdmCl was carried out. Slow refolding reactions were followed by manual mixing fluorescence experiments whereas for fast reactions, the sequential mixing mode of the stopped flow machine was used. It was confirmed that complete unfolding of the C_{f} domain is achieved within a few seconds at 2 M GdmCl, so that only isomerization processes in the unfolded C_{f} domain contribute to the observed changes in the refolding amplitudes.

Interrupted refolding experiments

Usually, the native conformation of a protein is not only thermodynamically but also kinetically the most stable state. This fact is exploited to follow the time-course of the formation of native or partially folded species in a protein folding reaction. In an interrupted refolding assay, N-test
or $t$-test, fully denatured protein is allowed to refold for different times before unfolding is carried out again. For the different times before unfolding is carried out again. Being kinetically most stable, native molecules unfold slower than all other species present. Importantly, the amplitudes of the different unfolding reactions directly mirror the amount of different species present. By selecting a final denaturant concentration where intermediates and the native state unfold with different rates, the simultaneous observation of their formation and decay becomes possible.

The $C_1$ domain was unfolded in 1.5 M GdmCl (if oxidized) or 1.0 M GdmCl (if reduced) before refolding for different times at a residual GdmCl concentration of 140 mM or 90 mM, respectively, was carried out. Independent of the redox state, the following unfolding step was performed in 1.3 M GdmCl. For all measurements, the sequential mode of the stopped flow machine was used. The final protein concentration was always 1 to 2 mM.

**Analysis of the kinetic data**

A three-state on- or off-pathway model was used to fit the data derived from $C_1$ folding experiments. For the analysis of chevron plots, the exact analytical solution was used to fit the fast and intermediate phase. For the three-state on-pathway model:

$$U \rightarrow I \rightarrow N$$

the characteristic equation is given by:

$$f(\lambda) = \lambda^2 - \lambda(k_{UI} + k_{IU} + k_{NI})$$

$$+ (k_{UI}k_{IN} + k_{IU}k_{NI} + k_{NI}k_{IN}) = 0$$

whereas for the three-state off-pathway model:

$$I \rightarrow U \rightarrow N$$

the characteristic equation is given by:

$$f(\lambda) = \lambda^2 - \lambda(k_{IU} + k_{UI} + k_{IN} + k_{NI})$$

$$+ (k_{IU}k_{IN} + k_{IU}k_{NI} + k_{NI}k_{IN}) = 0$$

The apparent rate constants $\lambda_1$ and $\lambda_2$ are the solutions of the quadratic characteristic equations. The logarithm of all microscopic rate constants was assumed to be linearly dependent on the denaturant concentration. The slowest phase, i.e. Pro-isomerization rates, were fit as described by Goto and Hamaguchi. For interrupted refolding experiments, on the other hand, irreversibility of the folding reactions was assumed to reduce the number of parameters, i.e. the data for the oxidized protein were numerically fit to the following model assuming two parallel pathways:

$$U_{P142C} \rightarrow I_{P142C} \rightarrow N$$

$$U_{P142I} \rightarrow I_{P142I} \rightarrow N$$

or in the case of the reduced protein:

$$U_{P142C} \rightarrow N$$

$$U_{P142I} \rightarrow N$$

where $c$ and $t$ stand for cis and trans. In equation (7), the rate for the formation of the intermediate was assumed to be equal on both parallel pathways. The Chemical Reactions module from the Berkeley Madonna software package was used to fit all data (Berkeley, CA). All graphs were created with Origin Pro Version 7.5 (Origin Lab, Northampton, MA, USA).

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The structure of a folding intermediate provides insight into differences in immunoglobulin amyloidogenicity

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Folding intermediates play a key role in defining protein folding and assembly pathways as well as those of misfolding and aggregation. Yet, due to their transient nature, they are poorly accessible to high-resolution techniques. Here, we made use of the intrinsically slow folding reaction of an antibody domain to characterize its major folding intermediate in detail. Furthermore, by a single point mutation we were able to trap the intermediate in equilibrium and characterize it at atomic resolution. The intermediate exhibits the basic β-barrel topology, yet some strands are distorted. Surprisingly, two short strand-connecting helices conserved in constant antibody domains assume their completely native structure already in the intermediate, thus providing a scaffold for adjacent strands. By transplanting these helical elements into β2-microglobulin, a highly homologous member of the same superfamily, we drastically reduced its amyloidogenicity. Thus, minor structural differences in an intermediate can shape the folding landscape decisively to favor either folding or misfolding.

Results

The Major Kinetic Folding Intermediate of C<sub>L</sub> Is Highly Structured. The C<sub>L</sub> domain folds via an obligatory intermediate on two parallel pathways to its native state, the slower one being limited by the isomerization of the Y34–P35 bond to the native cis conformation (30, 32). This bond is predominantly trans in the unfolded state. As a consequence, only ∼10% of the molecules are able to fold to the native state within a few seconds (30, 32), and ∼90% of the molecules have to undergo the intrinsically slow isomerization reaction before complete folding to the native state (30, 32). At 2°C limited by an intrinsically slow trans-to-cis peptidyl-prolyl isomerization reaction (24, 25). Experiments in which the critical proline residue was held in a trans state confirmed that this intermediate is a major determinant in amyloid formation (24, 26). In this regard, several studies showed that the most probable amyloidogenic precursor already possesses a large part of the native β-sheet topology with only the outer strands and loop regions being distorted (24, 25, 27).

Bearing in mind that intermediates are a rather general aspect of a protein folding reaction and that most polypeptides are in principle susceptible to amyloid formation (28), the question arises of how proteins avoid aggregation in the majority of cases. To address this issue we set out to study the folding pathway of the constant domain of the antibody light chain (C<sub>L</sub>) with high structural resolution. The C<sub>L</sub> domain is a particularly instructive model system because it also belongs to the Ig superfamily and, like β2m, forms a β-sandwich composed of seven strands stabilized by a single disulfide bond between strands B and F (29, 30). The cis proline residue associated with the amyloidogenic potential of β<sub>m</sub> is conserved in the C<sub>L</sub> domain (29). Furthermore, the overall folding mechanisms of the two proteins are highly similar (24, 30), each populating an intermediate state en route to the native state. Nevertheless, the C<sub>L</sub> domain has never been directly associated with amyloidogenic diseases even if present at much higher concentrations than β<sub>m</sub> in the blood (31). By the structural characterization of its major folding intermediate, we show how the C<sub>L</sub> antibody domain might avoid such harmful misfolding reactions.


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this reaction takes several hours to complete [see supporting information (SI) Fig. S1], allowing the major kinetic intermediate to be populated for a significant amount of time. CD spectra of the intermediate argue for a partially formed β-sheet framework and the absence of defined asymmetric environment around the aromatic amino acids (see Fig. S1).

To structurally characterize the intermediate state as well as the folding process on a residue level, >70% of the Cδ domain backbone was assigned by standard NMR techniques (Fig. 1A), and real-time $^{15}$N-$^1$H HSQC spectra were measured during refolding from the chemically denatured state. The first spectrum recorded after 14 min reflects almost exclusively the kinetic intermediate and
had only to be corrected for 10% of the C$_{\text{L}}$ molecules possessing the correct Y34–P35 isomerization state (Fig. 1A; see Materials and Methods for details). Because the chemical shifts of the amide protons strongly depend on their molecular environment, overlaying the HSQC spectra of the intermediate and the native state reveals similarities and changes in their environment during the folding process (Fig. 1A). The HSQC spectra of the native C$_{\text{L}}$ domain and the folding intermediate are superimposable for some residues but non-superimposable for others where significant differences in the chemical shifts are observed (Fig. 1A). To obtain more insights into the structural properties of the intermediate, the change in the peak intensity at the native chemical shift position was followed over time for each assigned residue. In every case the change in peak intensity could be well described by a single exponential function, if not already showing a native-like intensity after the dead-time of the experiment (Fig. 1B). As can be seen in Fig. 1C, the time constants of the folding of the individual residues show stochastic behavior around a mean value of $\tau = 199$ min at 2°C without any significant systematic deviations for any part of the protein. In contrast, initial HSQC amplitudes in the folding intermediate show interesting patterns; almost native initial amplitudes are found in correspondence of the two short helices connecting strand A and B as well as E and F and adjacent $\beta$-sheet termini suggesting that they are already in a native environment in the intermediate whereas low initial amplitudes are observed for some of the $\beta$-strands, in particular strands C and D, which suggests a lack of native structure (Fig. 1D). In Fig. 1E regions of high or low initial amplitudes are mapped on the crystal structure of C$_{\text{L}}$ revealing that the two helices and their local environment are highly structured in the intermediate.

**An Intermediate Structure at Equilibrium Trapped by a Single Point Mutation.** The initial HSQC amplitudes only provide hints on the structural properties of the intermediate. Equilibrium spectroscopic data could provide information more directly related to structure. Therefore, we tried to trap the intermediate at equilibrium by exploiting the isomerization reaction separating the intermediate from the native state. We hypothesized that mutating the P35 residue against another amino acid that preferentially adopts a trans peptide bond (33), such as Ala (C$_{\text{P35A}}$), might ”trap” the kinetic intermediate making it populated at equilibrium. Indeed, far-UV and near-UV CD spectra of C$_{\text{P35A}}$ were found to be very similar to the respective spectra of the kinetic intermediate (data not shown). To determine the stability of the mutant in comparison to the wild type (C$_{\text{wt}}$), denaturant-induced unfolding transitions were performed. The unfolding of both proteins, C$_{\text{wt}}$ and C$_{\text{P35A}}$, was a two-state process because there was concurrent loss of secondary structure in both species. Crucially, the trapping of the intermediate observed in the folding of C$_{\text{L}}$ was not feasible for the transiently populated kinetic folding intermediate of C$_{\text{P35A}}$ (Fig. 2A). Formation of the final refolding species of C$_{\text{P35A}}$ was further characterized by stopped-flow fluorescence spectroscopy. It could be described by a two-state folding reaction. (G) Formation of the final refolding species of C$_{\text{P35A}}$ was followed by interrupted refolding experiments. The data were fit by a double exponential function (residuals shown as inset) with 93% of the molecules folding via the fast pathway. (D) The folding mechanism of C$_{\text{P35A}}$ can be described by a simple two-state model neglecting the 7% slow folding species. Transitions were measured at 10 $\mu$M protein concentration, and all kinetic experiments were performed at a final protein concentration of 2 $\mu$M. All measurements were carried out at 20°C in PBS.

![Fig. 2](https://example.com/fig2.png)

**Fig. 2.** Stability and folding mechanism of the C$_{\text{P35A}}$ mutant. (A) Equilibrium unfolding transitions of C$_{\text{P35A}}$(green) and C$_{\text{wt}}$(red) determined by the intrinsic tryptophan fluorescence excited at 280 nm and detected at 360 nm (circles) as well as the far-UV CD-signal at 218 nm (inverted triangles). The data were fit to a two-state unfolding model. (B) Chevron plot for C$_{\text{P35A}}$ determined by stopped-flow fluorescence spectroscopy. It could be described by a two-state folding reaction. (C) Formed by the final refolding species of C$_{\text{P35A}}$ was followed by interrupted refolding experiments. The data were fit by a double exponential function (residuals shown as inset) with 93% of the molecules folding via the fast pathway. (D) The folding mechanism of C$_{\text{P35A}}$ can be described by a simple two-state model neglecting the 7% slow folding species. Transitions were measured at 10 $\mu$M protein concentration, and all kinetic experiments were performed at a final protein concentration of 2 $\mu$M. All measurements were carried out at 20°C in PBS.
the standard backbone experiments and triple resonance experiments for aliphatic side-chain assignment, 3D-NOESY spectra. These data revealed that large parts of the proteins possess almost identical carbon chemical shifts and, in addition, corresponding NOESY strips within the same range (Fig. S2 and data not shown), indicating that the intermediate already adopts a highly ordered structure. Notably, the two helices are fully formed, as judged from chemical shifts and the NOESY pattern from CL (Fig. S3). The overall solvent-accessible surface area of P35A was found to be on average 10% larger than for the wild type, which is in good agreement with the observed 20% decrease in the cooperativity parameter, \( m_{eq} \), for unfolding. Taken together, the NMR experiments in combination with the simulations provide a detailed picture of the major CL folding intermediate. The two small helices and their local environment are completely folded, and the intermediate exhibits a native-like core structure despite the presence of flexible regions that are able to adopt a variety of conformations.

Dissection of the Amyloidogenic Properties of the CL Domain and \( \beta_2 \)-Microglobulin. The constant domain of the antibody light chain has never been reported to be directly responsible for amyloidogenic processes even though it possesses the same topology as amyloidogenic variable antibody domains (V_{1}) or \( \beta_2 \)-microglobulin (\( \beta_2 \)m). In both cases, amyloid formation is assumed to proceed from a partially folded intermediate state (27, 35). Interestingly, neither amyloidogenic protein possesses the short strand-connecting helices that we identified as highly structured elements in the CL folding intermediate. Accordingly, the sequence or structure of these helical elements might play a role in the inhibition of amyloid formation. To test this, we exchanged the unstructured loop regions connecting strands A and B as well as strands E and F in \( \beta_2 \)m against the corresponding helical elements of the CL domain (\( \beta_2 \)mNC) (see Fig. S3). The \( \beta_2 \)mNC exchange mutant folds to a well-defined structure with similar far-UV CD-spectroscopic properties as wild-type \( \beta_2 \)m (see Fig. S3 for details). The helical elements destabilize \( \beta_2 \)mNC against thermal denaturation but have only a minor effect on its pH stability as compared to \( \beta_2 \)m (see Fig. S3). According to the TANGO algorithm (36), the aggregation propensity of its primary sequence is left unaffected by the mutations (data not shown). To assess the amyloidogenicity of the different proteins, we used established reaction conditions for \( \beta_2 \)m (37) and monitored fibril formation by atomic force microscopy (AFM). As expected, CL\( _{2m} \) and the CL\( _{P35A} \) mutant were not prone to fibril formation. In only one out of seven individual experiments were fibrils observed at pH 1.5, yet no fibrils were found at pH 3.0 or under physiological conditions, whether seeded or not (see Fig. S4). Importantly, a clear difference in amyloidogenicity is observed for wild-type \( \beta_2 \)m and the \( \beta_2 \)mNC mutant. Whereas \( \beta_2 \)m readily formed fibrils under all conditions tested, \( \beta_2 \)mNC only formed short fibrils at pH 1.5 (Fig. 4). At pH 3.0, fibrils were only detected in two out of seven individual experiments for \( \beta_2 \)mNC, and no fibrils were detected under physiological conditions for this protein, even when \( \beta_2 \)m fibrils were used for cross-seeding experiments (data not shown).
Finding that the edge strands A and G on one side of the CL et al. Feige

tion for the marked difference in amyloidogenicity between CL and intermediate are highly structured provides one possible explana-

The data clearly show that transplanting the sequences correspond-

The most striking structural features of the intermediate are the two completely folded small helices. Although they are strongly conserved in constant antibody domains, their role in the folding process has not yet been recognized. These helices seem to fulfill a spacer and orienting function between strand pairs A–B and E–F and provide hydrogen bond donors and acceptors for adjacent strands and loops. In addition, the helices appear to position hydrophobic residues (e.g., Y80 in helix 2) so that they can participate in the formation of the hydrophobic core. Our data suggest that the two helices are able to fold efficiently and autonomously to their native structures in the context of the intermediate. Hence, these two helices can be regarded as a scaffold within the C1 intermediate favoring the formation of a native-like topology by correctly positioning important parts of the molecule.

An important protective role against amyloid formation has been attributed to the edge strands of β-sheet proteins (40, 41). Our finding that the edge strands A and G on one side of the C1 intermediate are highly structured provides one possible explana-

In conclusion, our data show how a high degree of local structuring in a protein folding intermediate can significantly influence the folding landscape and favor robust folding over harmful misfolding. The different characteristics of C1 and β2m can be understood in evolutionary terms. Selection of antibodies took place under harsh extracellular conditions with high concentrations of the multimeric protein present (47), whereas β2m is found at much lower concentrations and usually associated with the MHC-I complex (21). Thus, small differences, acquired over the course of evolution, between members of the same protein superfamily can lead to the avoidance of pathogenic misfolding reactions while preserving an identical protein topology.

Materials and Methods

Protein Production and Purification. Proteins were expressed and purified as described in SI Methods.

Optical Spectroscopy. CD kinetics and spectra were measured as described in SI Methods. Equilibrium unfolding transitions, stopped-flow, and interrupted refolding experiments were performed as described in ref. 30. For the interrupted refolding experiments, C1P35A was unfolded in 1.5 M GdmCl, refolded in 136 mM GdmCl for different times, and finally unfolded again in 1.5 M GdmCl.

NMR Spectroscopy. If not stated otherwise, all spectra were recorded at 25°C on Bruker DMX600, DMX750, and AVANCE900 spectrometers as described in SI Methods. For folding studies, 15N-labeled unfolded C1 in PBS containing 2 M GdmCl was diluted 10-fold by adding ice-cold PBS without GdmCl. Real-time 15N–1H HSQC spectra were recorded at 2°C every 14 min by using selective proton flip-back pulses (48). Identical processing of all of the spectra was performed by using the program TOPSPIN 1.3 (Bruker BioSpin). To obtain the 15N–1H HSQC of the intermediate, 10% of the final spectrum was subtracted from the first recorded HSQC spectrum. Peak intensities were analyzed by using the program SPARKY (www.cgl.ucsf.edu/home/sparky). For kinetic studies, the intensities of every amino acid during the folding process were corrected for 10% native molecules and normalized to the corresponding intensity in the final spectrum after 7 h. Backbone resonance assignments were transferred from 25°C to 2°C recording a temperature series of spectra referenced to the internal standard TSP.

MD Simulations. The C1P35A mutant was created from the crystal structure of the C1 domain (PDB entry 1FHS). The bond preceding A35 was set to 1.5 μM. Additionally, the proteins were incubated under physiological conditions (PBS, 37°C) either seeded with β2m or β2mSCC fibrils or not. Formation of amyloid fibrils was assessed by AFM measurements. Representative pictures of each sample are shown.

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Supporting Information

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SI Methods

Protein Production and Purification. C_{P35A}^{wt} and C_{P35A}^{wt}, the mutant generated by site-directed mutagenesis (Stratagene), were expressed, refolded, and purified as described in ref. 1. The refolding buffer was modified [250 mM Tris/HCl (pH 8.0), 100 mM L-Arg, 10 mM EDTA, 1 mM GSSG, 0.5 mM GSH]. β-m was cloned from human cDNA (RZPD), the gene for equilibrated in PBS was used as a final purification step for all described above. A Superdex 75 26/60 column (GE Healthcare) M urea, 10 mM EDTA. The protein of interest in each case did not bind to the column, and the flow-through was refolded as described above. A Superdex 75 26/60 column (GE Healthcare) equilibrated in PBS was used as a final purification step for all proteins. Isootope-labeled proteins were expressed in M minimal medium containing either 15N ammonium chloride as the only nitrogen source or additionally 13C glucose as the only carbon source. All plasmids were sequenced, and the mass of each protein was confirmed by MALDI-TOF MS. 

CD Spectroscopy. CD measurements were performed in a JASCO J-715 spectropolarimeter. CD spectra were accumulated 16 times and buffer-corrected. For far-UV CD measurements, 10 μM protein in a 1-mm quartz cuvette was used; near-UV CD measurements were carried out at 50 μM protein concentration in a 5-mm quartz cuvette. To obtain far-UV as well as near-UV CD spectra of the intermediate, three spectra were measured and averaged beginning after 2 min of refolding. At the chosen instrumental parameters this corresponds to a maximum measuring time of 10.5 min in which maximally 5% of the molecules fold to the native state at 2°C. The same experiment was repeated seven times independently for far-UV and three times for near-UV CD spectra and subsequently averaged. The individually averaged spectra were identical within ±5%. The spectra of the intermediate were corrected for the 10% of native molecules that possess the correct Pro-isomerization state and hence refold to the native state within the dead time of the experiment (1). For pH transitions, 10 μM protein was incubated o.n. at the different pH values in Theorell–Stenhagen buffer (2) at 20°C before the far-UV CD signal at 218 nm was recorded. Temperature-induced unfolding transitions were monitored by the change in the far-UV CD signal at 205 nm with a heating rate of 20°C/h. Because all temperature-induced unfolding transitions were not completely reversible, an apparent melting temperature was derived by a Boltzmann fit.

NMR Spectroscopy. For C_{P35A}^{wt} as well as C_{P35A}^{wt}, backbone sequential assignments were obtained by using standard triple resonance experiments implemented with selective proton flip-back techniques for fast pulsing (3). Aliphatic side chain assignments were completed by using a combination of CCH-TOCSY and CCH-COSY experiments. Distance information was derived from a set of 3D-NOESY spectra, including NH- and CNH-NOESY spectra (4), in addition to a 1H-HSQC-NOESY spectrum. Dihedral angle restraints were determined for backbone Φ and ψ angles based on Cα, Cβ, C', and Hα chemical shifts using the program TALOS (5).

MD Simulations. The restraints for simulations of C_{P35A}^{wt} were derived from a comparison of the NMR chemical shifts of C_{P35A}^{wt} and C_{P35A}^{wt}. (1) Cα, NH, C', and Cβ atoms of residues with chemical shifts similar to C_{P35A}^{wt} were restrained by minimizing the RMSD of the interatomic distances of the four atoms with respect to the crystal structure. (2) Cα, C', and Cβ atoms were restrained as described for restraint I for residues where the NH chemical shift could not be determined but otherwise had similar chemical shifts to the wild type. (3) Residues that were unassigned in the C_{P35A}^{wt} mutant alone were restrained toward zero native contacts. In this regard, native contacts were computed by using only Cα, NH, C', and Cβ atoms, a cut-off of 0.65 nm, and for pairs of residues separated by at least five other residues in the primary sequence. (4) Φ and ψ dihedral angles were restrained toward values determined by TALOS if available. To minimize the number of restraints, restraints I–3 were only applied to residues located in secondary structure elements. The ensembles of structures obtained with and without the dihedral restraints were largely similar. The restraint potential, U, used was \( U(p, i) = (\alpha_i/2)(p - \bar{p})^2 \), where \( i = 1, 2, 3, 4 \) was the type of restraint used and corresponded to I–4 described above, \( p \) was defined as the mean squared difference between the calculated and target quantities to be restrained, \( \bar{p} \) was the target value of the restrained quantity, and \( \alpha_i \) was the value of the bias (kcal/mol) used to drive the restrained quantities toward the target values. A simulated annealing protocol was used to enhance the sampling of the conformational space. It was composed of six stages lasting a total of 452 ps: (i) 520 K for 62 ps, (ii) 650 K for 62 ps, (iii) 500 K for 62 ps, (iv) 420 K for 82 ps, (v) 350 K for 82 ps, and (vi) 300 K for 102 ps. A total of 100 cycles were performed.

AFM Measurements. For fibrillization experiments, a 100 μM protein solution in PBS was mixed 1:1 with buffer A (25 mM sodium acetate, 25 mM sodium phosphate) at pH 1.5 or 2.5 (final pH: 1.5 or 3.0 respectively, pH-meter reading) or PBS. The solution was incubated for 7 days, under slight shaking at 37°C. Seeds were prepared from β-m protein fibrils (grown at pH 1.5 for 7 days) by sonication for 15 min in a sonication bath. Two microliters of seeds were used to seed 100 μl of MilliQ water (Millipore), and dried overnight at room temperature. AFM measurements were done in contact mode with a Digital Instruments multimode scanning probe microscope (Veeco) at a scanning speed of 1.5 μm/min. DNP-S20 tips were used for all measurements (Veeco). Each experiment was repeated seven times.

Fig. S1. Structural and kinetic characterization of the C<sub>d</sub> domain and its major folding intermediate by CD spectroscopy. The secondary structure of the native (red open circles) and the unfolded C<sub>d</sub> domain (blue dots) was investigated by far-UV CD spectroscopy (A) and its tertiary structure by near-UV CD-spectroscopy (B). During refolding, spectra of the major folding intermediate were measured (green, filled circles) (A and B). The spectra of the intermediate were corrected for 10% of the molecules that already possess the correct Y34–P35 isomerization state and therefore refold to the native state during the dead time of the experiment. Refolding kinetics were followed by far-UV CD spectroscopy at 218 nm (C) as well as by near-UV CD spectroscopy at 280 nm (D). The signal of the unfolded domain is shown in blue, refolding to the intermediate in green, and folding from the intermediate to the native state as a single exponential fit in red (C and D). (E) The overall folding mechanism on the slow C<sub>d</sub> folding pathway can be described as a three-state process. Time constants for each reaction are indicated. For simplicity, the folding mechanism does not show the parallel folding pathway of the C<sub>d</sub> domain with the Y34–P35 bond in the correct isomerization state. For far-UV CD measurements, 10 μM protein was used, for near-UV CD measurements 50 μM protein was used. All measurements were carried out at 2°C in PBS in the presence of a GdmCl concentration of 2 M for the unfolded protein and 0.2 M for the folded protein.

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E

U
P35
trans

I
P35
trans

N
P35
cis

τ ~ 1 s

τ ~ 4 h
Fig. S2. Chemical-shift deviations of the nuclei $C^\alpha$, $C^\beta$, $C^\gamma$, and $H^\alpha$ from the random coil values published by Wishart and colleagues [Wishart DS, Sykes BD, Richards FM (1992) *Biochemistry* 31:1647–1651; Wishart DS, Sykes BD (1994) *J Biomol NMR* 4:171–180] for CL$^{\text{wt}}$ shown in blue and for the mutant CL$^{\text{P35A}}$ shown in red.
Fig. S3. Secondary structure, sequence and stability of β₂m, C₄ and the β₂mtoCL exchange mutant. (A) Far-UV CD spectra were measured at 20°C in PBS at 10 μM protein concentration. β₂m is shown in black, C₄ in red, and β₂mtoCL in blue. (B) Sequence elements from β₂m (black), C₄ (red), and β₂mtoCL (blue). Exchanged elements and the corresponding wild-type sequences are depicted in bold. The temperature-induced unfolding of all three proteins is shown in C. It was followed by far-UV CD spectroscopy at 205 nm. β₂m (black) shows a transition midpoint of 63°C, for C₄ (red) the midpoint is at 51°C, and for β₂mtoCL (blue) at 43°C. (D) The pH stability of β₂m (black), C₄ (red), and β₂mtoCL (blue) at 20°C was investigated at 218 nm by far-UV CD spectroscopy.
Fig. S4. Amyloidogenic properties of CL and CL\textsuperscript{P35A}. The proteins were incubated at pH 1.5 or 3.0 at 37°C for 7 days at a concentration of 50 μM. Additionally, both proteins were incubated under physiological conditions (PBS, 37°C) either seeded with 2 μm fibrils or not. Formation of amyloid fibrils was assessed by AFM measurements, and representative pictures of each sample are shown.
Structure of the Murine Unglycosylated IgG1 Fc Fragment

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A prototypic IgG antibody can be divided into two major structural units: the antigen-binding fragment (Fab) and the Fc fragment that mediates effector functions. The IgG Fc fragment is a homodimer of the two C-terminal domains (C\textsubscript{\text{H}2} and C\textsubscript{\text{H}3}) of the heavy chains. Characteristic of the Fc part is the presence of a sugar moiety at the inner face of the C\textsubscript{\text{H}2} domains. The structure of this complex branched oligosaccharide is generally resolved in crystal structures of Fc fragments due to numerous well-defined sugar–protein interactions and a small number of sugar–sugar interactions. This suggested that sugars play an important role in the structure of the Fc fragment. To address this question directly, we determined the crystal structure of the unglycosylated Fc fragment of the murine IgG1 MAK33. The structures of the C\textsubscript{\text{H}3} domains of the unglycosylated Fc fragment superimpose perfectly with the structure of the isolated MAK33 C\textsubscript{\text{H}3} domain. The unglycosylated C\textsubscript{\text{H}2} domains, in contrast, approach each other much more closely compared to known structures of partly deglycosylated Fc fragments with rigid-body motions between 10 and 14 Å, leading to a strongly “closed” conformation of the unglycosylated Fc fragment. The glycosylation sites in the C′E loop and the BC and FG loops are well defined in the unglycosylated CH2 domain, however, with increased mobility and with a significant displacement of about 4.9 Å for the unglycosylated Asn residue compared to the glycosylated structure. Thus, glycosylation both stabilizes the C′E-loop conformation within the CH2 domain and also helps to ensure an “open” conformation, as seen upon Fc receptor binding. These structural data provide a rationale for the observation that deglycosylation of antibodies often compromises their ability to bind and activate Fcγ receptors.

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Keywords: IgG1; antibody structure; glycosylation; antibody stability; antibody effector functions

Introduction

Antibodies not only are potent mediators and effectors within the arsenal of humoral immune response but also constitute a link to cellular immunity. Among them, the IgG class is the most abundant one. This heterotetrameric glycoprotein is composed of two light chains and two heavy chains and possesses two antigen-binding...
sites that are made up of the N-terminal domains of the light chain and the heavy chain, respectively. Based on proteolytically accessible fragments, the IgG antibody can be divided into two identical antigen-binding fragments (Fab) and the so-called Fc fragment. Fabs are connected by a flexible linker (the hinge region) to the Fc fragment. The Fc part itself is composed of two C1¿ domains that form a homodimer at the C-terminal end of the antibody molecule and two glycosylated C1¿ domains with sugar moieties located between the C1¿ domains. An important role of the Fc fragment is the mediation of effector functions such as antibody-dependent cellular cytotoxicity, phagocytosis, oxidative burst, and regulation of antibody production. All effector functions mediated through Fc receptors, as well as complement factor C1q-dependent reactions, have been shown to be compromised by deglycosylation of the C1¿ domains. Since Fc¿ receptors are constitutively expressed in different leukocytes such as macrophages, B-cells, and natural killer cells, they are an essential part of the connection between specific antibody–antigen binding and cellular responses (i.e., between humoral and cellular immune defense mechanisms). Therefore, the role of Fc glycosylation has gained much attention in molecular immunology. It was shown that a complex biantennary oligosaccharide is attached to a conserved Asn residue of the C1¿ domain. The oligosaccharide possesses a defined tertiary structure, as it could be resolved in crystal structures of glycosylated antibodies. Stabilization of the sugar structure is achieved primarily via sugar–protein contacts with the C1¿ domain and, in some cases, also via some sugar–sugar contacts. The core heptasaccharide is made up of four N-acetylgalactosamine (GlcNAc) residues and three mannose (Man) residues. It is often extended by fucose at the core heptasaccharide and by galactose or galactose and N-acetyleneuraminic acid residues at the branches. The branching point of the biantennary oligosaccharide is found at the third sugar residue after the Asn attachment, which is a Man. A ternary oligosaccharide is attached to a conserved Asn residue after the Fab fragment (Fig. 1c), an equilibrium stability of $\Delta G = 15 \pm 1 \text{kJ/mol}$ was obtained. For the enzymatically deglycosylated (Fig. 1b) and recombinant Fc fragment (Fig. 1c), stability was reduced to $\Delta G = 47 \pm 1 \text{kJ/mol}$, respectively, and the cooperativity parameter was reduced to $13 \pm 1 \text{kJ/mol}$ M. Thus, the overall stability was only reduced by 4–6%, whereas the cooperativity of the overall transition was decreased by $\sim 13\%$ upon deglycosylation. The same tendency is observed in temperature-induced unfolding transition, where a similar midpoint is observed for all Fc variants (Table 1) yet the sugar-free Fc forms show a lower apparent cooperativity of the transition (Fig. 1).

Results

**Biophysical characterization of the unglycosylated Fc fragment**

Oligosaccharides attached to the antibody Fc part are special because they fulfill essential structural and functional roles in antibody molecules. It has been argued that sugars serve as a spacer between the C1¿ domains determining the shape of the Fc part, thereby influencing its affinity for Fc¿ receptors and other Fc fragment-binding proteins. To investigate the structural consequences of the complete absence of oligosaccharides, we produced the Fc fragment of the murine IgG1 MAK33 in *Escherichia coli*. The Fc fragment was refolded from inclusion bodies in an established manner. Its secondary and tertiary structures in solution, based on far-UV and near-UV CD spectra, respectively, were highly similar to that of the authentic glycosylated Fc fragment and the enzymatically deglycosylated Fc fragment produced in eukaryotic cell cultures (data not shown).

To address the influence of glycosylation on the overall stability of the MAK33 Fc fragment, we measured temperature and guanidinium chloride (GdmCl)-induced unfolding transitions. Thermal melting curves were monitored by changes in far-UV CD signals, and GdmCl-induced unfolding transitions were followed by far-UV CD and fluorescence spectroscopy. For the glycosylated Fc fragment (Fig. 1a), an equilibrium stability of $\Delta G = 30.5\%$ (Table 2). The crystals belong to the space group $P 6_1 22$, with one Fc monomer in the asymmetric unit. The structure comprises all residues from Ser241 to Ser445 of MAK33. Residues Met238-Val240 at the N-terminus and residues His445-Glu450 were disordered and not defined by electron density; this corresponds to the Fc fragment, with the connection between the C1¿ domain and the C1¿

**Structure determination**

We determined the crystal structure of the unglycosylated Fc fragment by X-ray crystallography and molecular replacement at 2.5 Å, with an R-factor of 24.2% ($R_{free} = 30.5\%$) (Table 2). The crystals belong to space group $P 6_1 22$, with one Fc monomer in the asymmetric unit. The structure comprises all residues from Ser241 to Ser445 of MAK33. Residues Met238-Val240 at the N-terminus and residues His445-Glu450 were disordered and not defined by electron density; this corresponds to the Fc fragment, with the connection between the C1¿ domain and the C1¿
domain formed by the amino acid stretch Ser340-Ala348. The electron density of all amino acid residues of the domains was well defined; however, regions with higher temperature factors and, therefore, higher mobility were observed in the CH2 domain. These include the regions around the C′E, BC, and FG loops. The quality of the refined atomic model structure was confirmed with the program PROCHECK by analyzing the binding geometry of the backbone of the Fc fragment.

Structure of the unglycosylated Fc fragment

The domains of the unglycosylated MAK33 Fc fragment show a typical immunoglobulin fold: In both domains, C1′2 and C1′3, two layers of anti-parallel β-sheets enclose the hydrophobic core of each domain, as shown in Fig. 2. Disulfide bridges are formed between amino acid residues Cys264 and Cys324 (C1′2 domain), and amino acid residues Cys370 and Cys428 (C1′3 domain), respectively. As Table 1. Stability parameters of the Fc fragment

<table>
<thead>
<tr>
<th>Sample</th>
<th>ΔG_unfolding [kJ/mol]</th>
<th>m_eq [kJ mol⁻¹ M⁻¹]</th>
<th>T_melt [°C]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fc glycosylated</td>
<td>50±2</td>
<td>15±1</td>
<td>72.8±0.2</td>
</tr>
<tr>
<td>Fc deglycosylated</td>
<td>48±1</td>
<td>13±1</td>
<td>70.8±0.2</td>
</tr>
<tr>
<td>Fc recombinant</td>
<td>47±1</td>
<td>13±1</td>
<td>70.6±0.4</td>
</tr>
</tbody>
</table>

Thermodynamic parameters derived from GdmCl-induced unfolding transitions, as well as melting points from CD (T_melt) measurements, are shown.
judged by electron density, the disulfide bridges are fully formed. These are oriented perpendicular to the β-sheets and are located in the hydrophobic core of the protein.

A common feature of many members of the immunoglobin superfamily is the presence of at least one cis-proline residue in the native structure. Peptidyl–prolyl cis/trans isomerization reactions have been found as a rate-limiting step in the folding of a number of proteins. In agreement with the already solved crystal structure of the C1s3 domain of MAK33 from mouse, Pro377, which is located in a loop structure connecting two β-strands of the C1s3 domain, shows a cis-conformation. Previous folding studies suggested the absence of any cis-proline residues in the MAK33 C1i2 domain, which is confirmed by the crystal structure.

Figure 2 shows the physiological dimer of the MAK33 Fc fragment. It is obvious that the dimer is formed by noncovalent interactions of the C1s3 domains. Both C1i2 domains show no contacts with each other as they are at least 5 Å apart. The C1i2 loop C′E, ranging from Gln293 to Phe303, contains the N-glycosylation site consisting of the recognition sequence Asn-Ser-Thr. In the unglycosylated form of the Fc fragment of MAK33, this loop is more flexible than in structures of glycosylated Fc fragments. This seems to be due to the absence of the sugar moiety, which may have a stabilizing effect on this region.

A superposition of the unglycosylated IgG Fc fragment of MAK33 from mice with a glycosylated IgG Fc fragment from humans [Protein Data Bank (PDB) ID 1H3X] is shown in Fig. 3. Due to rigid-body motions of the C1i2 domains, significant shifts between about 10 and 14 Å are observed for the FG, BC, and C′E loops, respectively (indicated by the glycosylation site Asn297 or Asn300, respectively, in Fig. 3). These rigid-body motions result in a strongly closed conformation of the Fc fragment. The C1s3 domains, including the loop structures, superimpose very well. When superimposing only the C1i2 domains (data not shown), it is observed that they retain their structural integrity in the absence of oligosaccharides. However, the location and precise

| Structure of Unglycosylated IgG Fc |

Table 2. Data collection and refinement statistics for the unglycosylated MAK33 Fc fragment

<table>
<thead>
<tr>
<th>Data collection</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Space group</td>
<td>P6122</td>
<td></td>
</tr>
<tr>
<td>Unit cell dimensions a, b, c [Å]</td>
<td>96.45, 96.45, 90.63</td>
<td></td>
</tr>
<tr>
<td>Resolution range [Å]</td>
<td>25–2.5 (2.5–2.6)</td>
<td></td>
</tr>
<tr>
<td>Unique reflections</td>
<td>9063 (966)</td>
<td></td>
</tr>
<tr>
<td>Multiplicity</td>
<td>11.4 (10.3)</td>
<td></td>
</tr>
<tr>
<td>Completeness [%]</td>
<td>99.9 (100)</td>
<td></td>
</tr>
<tr>
<td>Rmerge [%]</td>
<td>5.6 (48.3)</td>
<td></td>
</tr>
<tr>
<td>I/σ(I)</td>
<td>25.7 (2.5)</td>
<td></td>
</tr>
</tbody>
</table>

Refinement

| Number of protein atoms | 1648 |
| Number of water molecules | 62 |
| Resolution range [Å] | 10–2.5 |
| Reflections [n (%)]    | 8342 (93.6) |
| Test set               | 485 (5.4) |
| Completeness [%]       | 99.9 (100) |
| Rcryst [%]             | 24.2   |
| Rfree [%]              | 30.5   |
| rmsd from ideal geometry | 0.011 |
| Bond length [Å]        | 1.66   |
| Ramachandran plot [%]  | 80.6   |
| Residues in most favored regions | 14.4 |
| Residues in generously allowed regions | 3.9 |
| Residues in disallowed regions | 1.1 |
| (Asn387 and Lys417)    |       |
| Nonglycine and nonproline residues [n (%)] | 180 (100.0) |
| Average B-factor [Å²] | Protein | 55.3 |
|                        | Water   | 47.2 |
|                        | Ions    | 70.2 |

Values for the outer-resolution shell are given in parentheses.

\[ R_{cryst} = \sum_{hkl\not\in W} |F_{obs}| - k|F_{calc}|/\sum_{hkl\not\in W} |F_{obs}|, \]

\[ R_{free} = \sum_{hkl\in T} |F_{obs}| - k|F_{calc}|/\sum_{hkl\in T} |F_{obs}|. \]
conformations of the C'E, BC, and FG loops are changed to some extent. A major shift (4.9 Å) is observed for the C'α position of Asn300, the glycosylation site, in the C'E loop compared to the equivalent Asn297 in the human Fc. For the BC-loop position (Lys271 in murine Fc/His268 in human Fc),

**Fig. 3.** Comparison of the unglycosylated murine Fc fragment with a glycosylated human counterpart. Superposition of the unglycosylated IgG Fc fragment of MAK33 (in green) with a glycosylated IgG Fc fragment from humans (PDB ID 1H3X, in magenta). The human glycosylation site Asn297, as well as the corresponding murine residue Asn300, is shown to highlight its displacement upon deglycosylation. The sugar moiety of the human Fc fragment is shown in a ball-and-stick representation.

**Fig. 4.** Superposition of the unglycosylated IgG Fc fragment of MAK33 shown in red with a glycosylated IgG Fc fragment from humans bound to the Fcγ receptor III given in blue and yellow, respectively (PDB ID 1E4K). Oligosaccharides of the IgG Fc fragment are depicted in a ball-and-stick model. To illustrate the “super-closed” conformation of the unglycosylated murine MAK33 Fc fragment, we show the Asn residues of the glycosylation sites as ball-and-stick models in green (Asn300, MAK33) and blue (Asn297, human IgG Fc), respectively.
a shift of 2.6 Å is observed; in the FG loop, a displacement of 1.3 Å occurs. The overall motion is towards the binding site of the oligosaccharide. We conclude that especially the C′E loop, which has already been shown to show different conformations in different crystal structures,21,27,28 is supported and stabilized in the fully glycosylated antibody.

Figure 4 shows a superposition of the unglycosylated IgG Fc fragment of murine MAK33 with a glycosylated human IgG Fc fragment bound to an Fcγ receptor III (PDB ID 1E4K).29 Here as well, no significant differences in the C1γ domains are found, in contrast to the C1β domains, where huge movements are observed. The binding of the Fc fragment to the Fcγ receptor III is based on the interactions of five regions of the C1γ domain with the receptor. These interactions involve rather few side chains, including the highly conserved amino acid residue Pro329, which represents one main interaction contact with the Fcγ receptor III. It is just as well ordered in the Fc fragment of MAK33 from mouse. The regions from Asp265 to Glu269 (Val268-Ile272 in mouse) and those from Asn297 to Thr299 (Asn300-Thr302 in mouse) that interact with the Fcγ receptor III are found in loops in the unglycosylated Fc fragment of the MAK33. These residues show rather small changes in their position within each domain. Although Fcγ receptor III binding induces a structural adaptation of domain positions of C1γ domains in Fc, including a prominent deviation from 2-fold symmetry,15 the highly closed conformation observed in the present structure would require a significantly larger rearrangement that will have a prominent effect on binding affinity.

Discussion

Effect of deglycosylation on Fc overall conformation

It has been noticed that the Fc fragment assumes an “open” conformation in the complex with Fcγ receptor III15 and that stepwise deglycosylation is associated with the formation of more “closed” conformations and structural variations in the C′E loop with shifts of up to 1.5 Å.21 The distance between the Cα atom of Pro329A and the Cα atom of Pro329B has been used as reference, as they point towards the 2-fold axis of the Fc fragment and are located in a rigid part of the molecule. These distances vary between 33.7 Å for the (G2F)2 glycoform with 10 monosaccharides per chain and 21.9 Å for the (MN2F)2 glycoform with 4 monosaccharides per chain.21 In the present study, the distance between the corresponding Pro332A and Pro332B is only 11.6 Å, which indicates that the fully unglycosylated Fc fragment can adopt the most “closed” structure observed so far. It should be noted, however, that effects of the crystallization conditions cannot be ruled out completely. Additionally, the absence of hinge disulfide bonds in this work might have an impact on the overall Fc conformation. However, the first resolved residue of the C1γ2 domain is only 13.6 Å apart from its counterpart. Five flexible amino acids would separate the last hinge Cys residue from this first resolved residue and, hence, the observed structure would, in principle, be compatible with the presence of hinge disulfide bonds.

By the overall motions of the C1γ2 domains, the Cα atom of Asn300 (the sugar attachment site) moves inwards and considerably downwards to the C1γ3 platform. As a result, for assuming the Fcγ-receptor-binding-competent conformation, drastic structural rearrangements would be required from that starting conformation. For other glycoforms with known structures, at least some contacts between C1γ2 domains are maintained via oligosaccharides, which may guide the molecules towards an active conformation.

In the absence of oligosaccharides, such a rearrangement will be entropically unfavorable. In addition, there is no support for the C′E loop and, to a lesser degree, for the BC and FG loops to adopt the binding-competent conformation. It should be noted that Fcγ receptors bind in a highly nonsymmetric fashion, requiring an induced fit.15 It involves the C′E loop (with contacts between Fcγ receptor III and GlcNAc1), the BC loop, residues from the hinge region (around Leu235) on one C1γ2 domain, and residues from the hinge region and the FG loop from the other C1γ2 domain. Most significantly, the disulfide-linked (Cys229A-Cys229B) hinge region connecting the Fc and Fab parts of the antibody is pushed to the direction opposite to that of the Fcγ receptor III binding site. Nevertheless, both C1γ2 domains retain very similar loop conformations.15 In this complex, the Man4 moieties of the minimally required oligosaccharides have a minimum distance of about 13 Å. Shortening of the Pro329A and Pro329B distance below 20 Å would be prohibited by van der Waals contacts between the core units of the oligosaccharides.

Effect of deglycosylation on the C1γ2 domain

A comparison of native stepwise deglycosylated Fc structures and our completely unglycosylated Fc structure, as well as a recently solved crystal structure of a human unglycosylated C1γ2 domain,20 reveals that the C′E loop becomes more flexible with decreasing sugar content. In the latter structure, the BC and FG loops are also significantly displaced compared to the native glyco form. A shift in the C′E loop has been observed in glycoforms (M3N2F)2, which lacks the outer GlcNAc6 and GlcNAc9 of the core heptasaccharide, and (MN2F)2, which additionally lacks Man5 and Man8, the first branched sugars of the core heptasaccharide.21 It is accompanied by a concomitant movement of the entire oligosaccharide moiety and a slight reduction in affinity for Fcγ receptor IIb, which was attributed to an unfavorable change in entropy.5 It has been suggested that the higher Fcγ receptor IIb affinities of all glycoforms containing GlcNAc sugar residues on the α(1–6) and
α(1–3) arms result from a rigid C/E-loop conformation that has also been observed in Fcγ receptor complexes. A conformational change in the BC loop has been also observed upon deglycosylation using 1H NMR with His268 as reporter group.20

Large conformational differences are observed for sugars of the α(1–6) arm in glycoform (G0F)2, which only comprises the core heptasaccharide and usually fucose attached to GlcNAc1, with only GlcNAc6 remaining in an invariant position where it shields Phe243 (Phe246 in murine Fc) from the solvent. Strikingly, its absence in the glycoform (M3N2F)2 is accompanied by a major decrease in melting temperature and unfolding enthalpy.5 In addition, longer oligosaccharide chains result in lower temperature factors for oligosaccharide moieties and C/E loops.21

Taking these observations together, the oligosaccharides, which are bound by multiple weak contacts, have a significant conformational variability and, by a shift of the C/E loop bearing the covalent attachment site, can also slide along the protein surface within certain limits. The overall stabilizing effect appears to involve shielding of key residues such as Phe243 from the solvent.

**Stability of the Fc fragment**

The C12 and C13 domains in the context of the Fc fragment are mostly regarded as independent units in terms of stability and folding. The stability of the C13 domain is assumed to be unaffected by the C12 glycosylation status, whereas C12 stability itself decreases with trimming of its oligosaccharide.6,14 We find that the enzymatically deglycosylated and unglycosylated Fc fragments are slightly less stable against GdmCl-induced denaturation than the glycosylated counterpart. The observed cooperativity is reduced more significantly. This finding can most likely be attributed to two underlying transitions: one for the C12 domain and one for the C13 domain, which cannot be dissected under the applied conditions due to very similar stabilities.6,14 The observed chemical stabilities for the different Fc glycoforms correspond to the ones of the isolated C12 or C13 domain, arguing against a significant impact of the structure of the Fc fragment on the chemical stability of individual domains.6,18,19 A somehow different behavior is observed in temperature-induced unfolding experiments. Again, the midpoint of the transition is only slightly affected by an unglycosylated C12 domain, yet cooperativity seems to be significantly reduced due to the two underlying transitions.6,14 The thermal stability of the unglycosylated C12 domain itself seems to be higher in the Fc fragment than in the isolated C12 domain, where unfolding around physiological temperatures has been reported.19 Even though sequence effects at the termini of the C12 domain, which were slightly different in a previous study and this study, cannot be ruled out completely,19 stabilization by a weak interaction with C13 (the partner C12 domain) or due to excluded volume effects in the context of the dimeric Fc fragment is more likely to explain the differences.

**Conclusion**

In the absence of a short core tetrasaccharide, which comprises all sugar moieties until the branching point at Man4, the C/E loop, including Asn297 as the glycosylation site, changes its conformation. This leads to a shift of the neighboring BC loop and, to a lesser degree, also of the FG loop by about 2.2 and 1.7 Å relative to the position of the fully glycosylated Fc. In addition, this allows the Fc fragment to adopt an extremely “closed” state, indicated by a distance of only 11.6 Å between Pro332A and Pro332B (murine) compared to a distance of 30.3 Å between the corresponding residues Pro329A and Pro329B in the human Fc–Fcγ receptor III complex.15 The latter value corresponds well to that observed in native Fc.1,21 Stepwise truncation of the oligosaccharides induces progressively “closed” conformations with Pro329A-Pro329B distances of between 26.6 and 21.9 Å, resulting in a rather small loss of affinity (from 1 to 3 µM).5,6 The absence of the four core oligosaccharide moieties obviously results mainly in the destabilization of the C/E loop within a C12 domain and a “super-closed” conformation of the Fc fragment. Together, these effects explain the often observed inability of unglycosylated Fc to bind and activate Fcγ receptors.

**Materials and Methods**

**Materials**

GdmCl (ultrapure) was purchased from MP Biomedicals (Eschwege, Germany). All other chemicals were obtained from Merck (Darmstadt, Germany). GdmCl concentrations were determined by the refractive indices of the solutions. Unless otherwise stated, all experiments were carried out in phosphate-buffered saline (PBS) at 25 °C.

**Proteins**

The glycosylated MAK33 Fc part was a kind gift from Dr. Helmut Lenz (Roche, Penzberg, Germany). For complete deglycosylation, 10 mg of the native Fc fragment was incubated with 100 U of N-glycosidase F (Roche, Mannheim, Germany) for 72 h at 37 °C. The native Fc fragment and the deglycosylated Fc fragment were purified by size-exclusion chromatography on a Superdex75pg 26/60 gel-filtration column (Amersham Biosciences, Uppsala, Sweden) equilibrated in PBS. Complete deglycosylation was confirmed by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry. Recombinant Fc comprised MAK33 residues Glu239-Glu450 and was purified as described previously.22 All proteins used in this study were devoid of hinge disulfide bridges.

**Crystallization and structure determination**

The deglycosylated form of the human Fc fragment could not be crystallized in the past by dialyzing against
water or different buffers. Therefore, an extensive screening for crystallization conditions was performed. Under the final conditions using the sitting-drop vapor-diffusion method, each drop contained 2 μl of the recombinant Fc fragment with a concentration of 15 mg/ml in 50 mM potassium phosphate buffer (pH 8.0) was mixed with an equal volume of the crystallization buffer and incubated in the presence of a 300-μl crystallization buffer reservoir. The reservoir solution contained 0.05 M calcium acetate, 0.1 M imidazole/HCl (pH 8.0), and 35% 2-ethoxyethanol (Emerald Cryo I, 18; Emerald Biosystems, Bainbridge Island, WA) as crystallization buffer.

**Diffraction data collection and data reduction**

Because cryoprotection was given under crystallization conditions, unglycosylated Fc fragment crystals were directly frozen in a nitrogen stream of 100 K (Oxford Cryosystems, Oxford, UK). Data were collected at a wavelength of 1.5418 Å on an image plate system (MAR Cryosystems, Oxford, UK). Data were collected at a maximum size of approximately 300 μm × 300 μm × 100 μm at 20 °C in about 10 days, were obtained after several days.

**Model building and refinement**

Model building was performed with the program MAIN. The structure was refined with CNS. During refinement, a rigid-body routine was employed using the Fc domains as individual rigid domains. The dihedral angles of all amino acid residues were refined to allowed values in the Ramachandran plot. The model was further improved in successive cycles of model building and refinement. Individual isotropic B-factors were refined.

**Analysis and graphical representation**

Structural figures were prepared with MOLSCRIPT and BOBSCRIPT and were rendered with Raster3D.

**Stability measurements**

Fluorescence measurements were performed in a Spex FluoroMax II fluorometer (Instruments SA, Edison, NJ), and CD measurements were performed in a Jasco J-715 spectropolarimeter (Jasco, Grossumstadt, Germany). GdmCl-induced unfolding transitions were followed either by the intrinsic tryptophan fluorescence of the Fc fragment at 354 nm and excited at 280 nm or by the far-UV CD signal at 218 nm. Fluorescence measurements were carried out in a 1-cm quartz cuvette; for far-UV CD measurements, a cuvette with 1-mm pathlength was used.

At each GdmCl concentration, 50 individual data points were recorded with an integration time of 1 s at the observation wavelength and averaged. Samples were incubated for 10 days prior to measurements. GdmCl-induced unfolding transitions were completely reversible. Data were evaluated according to a two-state unfolding model for a dimeric protein:

\[
S_{\text{obs}} = y_N(1 - F_U) + y_U F_U
\]

\[
F_U = \frac{[U]}{C_P} = \frac{4c_P}{K_U + 8k_UC_P - K_U}
\]

\[
K_U = \frac{[U]^2}{[N_2]} = e^{-\frac{\Delta G_U}{RT}} = e^{-\frac{(\Delta G_{H_2O} - m(GdmCl))}{RT}}
\]

where \(S_{\text{obs}}\) denotes the observed CD or fluorescence signal at a certain GdmCl concentration, \(y_N\) is the signal of the native protein, and \(y_U\) is the signal of the unfolded protein (both signals are assumed to be linearly dependent on GdmCl concentration). The fraction of unfolded protein is termed \(F_U\), and the total protein concentration is termed \(C_P\). The protein stability \(\Delta G\) was assumed to be linearly dependent on GdmCl concentration with the cooperativity parameter \(m\).

Temperature-induced unfolding was followed by CD spectroscopy. Fc was unfolded at a heating rate of 30 °C/h in a 1-mm quartz cuvette, and the signal change at 218 nm was recorded. Melts were fitted by a Boltzmann function. In all cases, temperature melts were not completely reversible; however, in each case, three individual experiments overlaid well. Hence, no thermodynamic data could be derived from the temperature melts. All spectroscopic measurements were carried out at a concentration of 5 μM Fc monomer.

**PDB accession number**

The PDB accession number is 3HKF.

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**References**


An Unfolded C_H1 Domain Controls the Assembly and Secretion of IgG Antibodies

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SUMMARY

A prerequisite for antibody secretion and function is their assembly into a defined quaternary structure, composed of two heavy and two light chains for IgG. Unassembled heavy chains are actively retained in the endoplasmic reticulum (ER). Here, we show that the C_H1 domain of the heavy chain is intrinsically disordered in vitro, which sets it apart from other antibody domains. It folds only upon interaction with the light-chain C_L domain. Structure formation proceeds via a trapped intermediate and can be accelerated by the ER-specific peptidyl-prolyl isomerase cyclophilin B. The molecular chaperone BiP recognizes incompletely folded states of the C_H1 domain and competes for binding to the C_L domain. In vivo experiments demonstrate that requirements identified for folding the C_H1 domain in vitro, including association with a folded C_L domain and isomerization of a conserved proline residue, are essential for antibody assembly and secretion in the cell.

INTRODUCTION

In eukaryotic cells, proteins destined for secretion mature within the endoplasmic reticulum (ER) and are subject to rigorous quality control prior to their transport to the Golgi (Helenius et al., 1992). This usually involves surveillance of the folding status, the correct posttranslational modifications, and proper oligomerization (Helenius et al., 1992; Elggaard and Helenius, 2003; Christis et al., 2008). A prominent example of this is immunoglobulin G (IgG), the most abundant antibody in the blood. It is a heterotetrameric glycoprotein assembled from two light and two heavy chains that are comprised of two and four, respectively, compact Ig domains that are structurally almost identical (Huber et al., 1976). Each domain shows a β barrel topology, a two-layer sandwich structure composed of between seven and nine antiparallel β strands (Huber et al., 1976; Amzel and Poljak, 1979). The fold is stabilized by an internal disulfide bridge (Goto and Hamaguchi, 1979) that is located in the hydrophobic core and lies perpendicular to the β sheets (Huber et al., 1976). Most of these structural characteristics are shared by the ubiquitous members of the immunoglobulin (Ig) superfamily (Bork et al., 1994), which perform a broad variety of extracellular recognition functions (Williams and Barclay, 1988; Rougon and Hobert, 2003; Aricescu and Jones, 2007). The evolutionary success of the Ig superfamily has fueled a vast scope of investigations on its biophysical properties. In particular, the folding pathways of diverse members of the Ig superfamily have been studied in detail (Goto and Hamaguchi, 1982; Freund et al., 1996; Thies et al., 1999; Cota et al., 2001; Paci et al., 2003; Feige et al., 2004) and have provided insights into determinants of robust folding (Hamill et al., 2000; Feige et al., 2008) as well as potentially harmful misfolding of this class of proteins (Kameda et al., 2005; Jahn et al., 2006; Qin et al., 2007).

To become secreted from the cell and fulfill their biological functions, the individual domains of an antibody not only have to fold into their native tertiary structure, but furthermore must assemble into a defined quaternary structure (Porter, 1973; Huber et al., 1976). Whereas isolated antibody light chains can be secreted from the ER (Coffino et al., 1970; Melchers, 1971), unpaired heavy chains are actively and efficiently retained in the ER (Bole et al., 1986; Hendershot et al., 1987). Antibody heavy-chain and light-chain synthesis occur asynchronously during B cell development (Burrows et al., 1979), and only completely assembled molecules can both bind to antigen and carry out effector functions. Therefore, tight quality control of their assembly prior to secretion is vital. It is known that the first constant domain of the heavy chain, the C_H1 domain, plays an important role in this retention process (Hendershot et al., 1987; Kaloff and Haas, 1995). If deleted or replaced with another antibody domain, isolated heavy chains can be secreted, as occurs in the case of the rare heavy-chain diseases (Wolfenstein-Todel et al., 1974; Adetugbo, 1978; Hendershot et al., 1987), or naturally in camelid antibodies, which do not contain light chains (Hamers-Casterman et al., 1993). In the context of the whole IgG molecule (Figure 1A), the C_H1 domain is associated with the constant domain of the light chain (C_L) and shows the typical immunoglobulin fold (Huber et al., 1976). In vivo, C_H1 is the only antibody domain that is stably bound to the molecular chaperone BiP and remains in a reduced form even after assembly with light chain (Vanhove et al., 2001). The basis for the unusual behavior of the C_H1 domain has remained enigmatic.

Here, we set out to study the role of the C_H1/C_L association for correct antibody assembly and secretion. To our surprise, we found that C_H1 is an unfolded protein in isolation that gains...
structure only upon interaction with its cognate partner, C_L. Based on this finding, we analyzed the association-coupled C\textsubscript{1} folding pathway and its modulation by the chaperone BiP in detail and provide a comprehensive picture for the control of antibody secretion in the cell.

RESULTS

The Murine IgG1 C\textsubscript{1} Domain Is Intrinsically Disordered

Antibodies are modular structures composed of a series of structurally highly homologous domains (Figure 1A). These domains...
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can usually be produced and studied separately, as they repre-
sent independent structural units (Goto and Hamaguchi, 1982; 
Lilie et al., 1995). Surprisingly, analysis of the murine IgG1 CH1 domain revealed that, in marked contrast to all antibody domains 
studied thus far (Goto and Hamaguchi, 1982; Thies et al., 1999; 
Feige et al., 2004; Rothlisberger et al., 2005), the isolated CH1 domain is an unfolded protein, irrespective of whether its internal 
disulfide bridge is formed or not (Figure 1B; Figures S1 and S2, 
available online). To further characterize the unfolded state of 
the CH1 domain under physiological conditions, iodide fluores-
cence quenching experiments were carried out. The experiments 
indicate no significant differences in the burial of tryptophan resi-
dues between CH1 in PBS and in 3 M GdmCl (data not shown). 
Additionally, NMR experiments were recorded on a highly 
deut erated CH1 sample. Deuteration enables detection of long-
range NOEs even if they belong to only a subset of conformers. 
In these experiments, no long-range NOEs could be determined, 
and, consequently, no preferential conformation of CH1 seems to 
persist. Taken together, these data argue against the presence of 
a significant amount of stable structure in the isolated CH1 domain. However, the pattern changed completely when the CH1 
domain, the cognate association partner of CH1 in the antibody, 
was added. Only then did we observe folding of the CH1 domain 
to a well-defined β sheet structure (Figure 1B). Thus, the CH1 
domain is necessary and sufficient to induce structure formation 
in CH1. This folding process was observed only if the internal 
disulfide bridge in the CH1 domain was present (Figure S2). 
Based on these findings, a key role of this association-coupled 
folding reaction for correct antibody assembly can be antici-
pated, which we aimed to elucidate in more detail.

The Mechanism of Induced Folding of the CH1 Domain

To understand how binding to CL and folding of CH1 are coupled, 
the thermodynamic and kinetic parameters for this reaction were 
established. The dissociation constant of the two domains was 
determined to be 6.2 ± 0.4 μM (Figure 1C) by the change in 
intrinsic fluorescence emission upon CH1-induced CH1 folding 
(Figure 1C, inset). A moderate affinity is expected because CH1 
has to fold upon binding to CL. One should bear in mind that 
the observed dissociation constant is orders of magnitude lower 
than the antibody concentration in the ER of plasma cells (Cenci 
and Sitia, 2007); thus, association will readily occur in vivo. The 
analysis of the kinetics of secondary, tertiary, and quaternary 
structure formation by using far-UV CD spectroscopy (Figure 1D), 
near-UV CD spectroscopy, and analytical high-performance liquid 
chromatography (HPLC) (Figure S1) showed that all three 
processes occur with virtually identical time constants of τ = 
60 ± 10 min at 25°C. Hence, all of these processes are likely 
rate limited by the same slow reaction. This slow folding reaction 
could be accelerated by the ER-specific peptidyl-prolyl isom-
erase cyclophilin B (Figure 1D). Thus, the slow folding phase 
can be attributed to the isomerization of peptidyl-prolyl bonds 
within the CH1 domain, which possesses an unusually high 
number of three cis prolines in the native state (Augustine et al., 
2001). Prior to the slow folding to the native structure, the CH1 
domain forms an intermediate with the Cc domain in a concen-
tration-dependent reaction (Figure 1E). As this complex could be 
detected by fluorescence anisotropy measurements, but not by 
the other techniques outlined above, it is likely a dynamic species 
with an only marginally folded CH1 domain. In the complete anti-
body, a disulfide bridge covalently links the CH1 domain with the 
Cc domain (Figure 1A). If the bridge-forming Cys residues were 
included in the CH1 domain as well as the Cc domain, no change 
in the folding state of the isolated domains and the CH1-induced 
folding of the CH1 domain was observed (Figure S3), but forma-
tion of covalent dimers could be readily followed by SDS-
PAGE. As covalent dimers were formed with the same rate as 
the slow CH1 folding reaction and the reaction could be acceler-
ated by cyclophilin B (Figure 1F), it is clearly limited by proline 
isomerization and hence complete folding of the CH1 domain.

Tak en together, the CH1-induced folding of the CH1 domain can 
be dissected into three reactions: first, oxidation of the internal 
CH1 disulfide bridge has to take place. Then, a transient hetero-
dimeric intermediate is formed, and, subsequently, peptidyl-
prolyl isomerization is required to allow folding to the native state 
and covalent assembly with Cc (Figure 1G).

An Atomic Level Description of the CH1 Folding Pathway

To resolve the specific recognition and the folding pathway of 
the intrinsically disordered CH1 domain at the level of atomic 
resolution, NMR experiments were performed. The 15N-1H 
HSQC spectrum of the isolated CH1 domain is characteristic of 
an unfolded protein (Figure 2A, red spectrum), confirming the 
results described above. In contrast, after induced folding by 
Cc, the CH1 domain shows well-dispersed spectra (Figure 2A, 
blue spectrum). The backbone assignment of the CH1 domain 
in the complex was achieved by a combination of triple-reso-
nance experiments and NH residual dipolar couplings (RDCs). 
All obtained NMR data, the carbon chemical shifts, the NH RDCs, 
and the MEXICO (Gemmecker et al., 1993) water exchange rates 
(data not shown) agree with an all-β structure for the CH1 domain 
in the presence of Cc, similar to that observed in the crystal struc-
ture of IgG antibodies (Augustine et al., 2001). Because complete 
folding of the CH1 domain is limited by proline isomerization, 
and hence associated with a high activation energy, the final 
folding step is significantly decelerated at low temperatures. 
This allowed us to characterize the trapped intermediate and 
resolve the association-coupled folding process by using real-
time 15N-1H HSQC experiments. For each assigned residue, 
changes of the amplitudes over time could be described by a 
single exponential function (Figure 2A, inset). Notably, some 
residues already exhibit significant intensities in the first spec-
trum recorded after 20 min (Figure 2B, red bars). These residues 
are likely to already adopt a native-like backbone conformation 
prior to the slow peptidyl-prolyl isomerization reaction. Alto-
together, 10 residues, which are part of the β sheets that form 
the mature structure, were found to be in a native-like environ-
ment in the intermediate. Mapping these residues on the crystal 
structure of the CH1 domain of a murine IgG1 Fab fragment revealed 
how the association-coupled folding reaction of this antibody 
domain might proceed. Residues Thr22, His49, Ser65, and Thr67 
in the CH1 domain, which form part of the Cc interface, seem to 
be already correctly positioned in the intermediate (Figure 2C, left 
panel). Importantly, His49 and Ser65 are involved in hydrogen 
bonds with the Cc domain in the native state. The interaction 
with Cc apparently initiates the formation of a hydrophobic cluster.
in the C1 domain including Val21, Val68, Trp73, and Val78 (Figure 2C, right panel). Additionally, interaction of Val48 and Val66 might also be involved in this hydrophobic cluster, although this could not be directly addressed due to peak overlap for Val66. Thus, a few key interactions between CL and CH1 establish an interface between the two domains in the intermediate, which allows for the formation of a hydrophobic folding nucleus in the CH1 domain, and subsequent prolyl isomerization paves the path to the native state.

To identify the residues responsible for the slow folding reaction of the C1 domain, each of the three proline residues (Pro32, Pro34, and Pro74) that adopt a cis conformation in the native state (Figure 2C, right panel) was individually mutated to alanine. 15N-1H HSQC spectra were recorded for each of the mutants in the absence and in the presence of C1. All three mutants showed almost indistinguishable spectra compared to the wild-type C1 domain in the absence of C1 (Figure 2D). Importantly, two of the mutants, Pro34Ala and Pro74Ala, displayed well dispersed HSQC spectra in the presence of C1 that are very similar to that of wild-type C1 in the presence of C1 (Figure 2D), arguing that isomerization of these two prolines is not essential for C1 domain folding. However, when the Pro32Ala mutant was similarly examined, identical spectra
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Figure 3. Characterization of the Interaction between BiP and C\textsubscript{H}1 In Vitro

(A) The affinity between BiP and oxidized C\textsubscript{H}1 (filled circles, straight line) as well as reduced C\textsubscript{H}1 (open circles, dashed line) was determined by analytical HPLC experiments. The data were fitted to a one-site binding model to determine the $K_D$.

(B) The association kinetics between 1 mM BiP and varying concentrations of oxidized C\textsubscript{H}1 (filled circles) and reduced C\textsubscript{H}1 (open circles) were measured to determine the rate constants of the reaction. For oxidized C\textsubscript{H}1, $k_{on} = 0.00026 \pm 0.00002$ M$^{-1}$ min$^{-1}$ and $k_{off} = 0.0050 \pm 0.0002$ min$^{-1}$ were obtained. For the reduced C\textsubscript{H}1 domain, the corresponding values were $k_{on} = 0.00041 \pm 0.00003$ M$^{-1}$ min$^{-1}$ and $k_{off} = 0.0047 \pm 0.0003$ min$^{-1}$. The left inset shows single HPLC runs of 8 mM oxidized C\textsubscript{H}1 and 1 mM BiP after 10 min (blue) and 200 min (red) of coincubation. The right inset shows the overall observed single exponential association kinetics between 1 mM BiP and 8 mM oxidized C\textsubscript{H}1.

The Antibody Domain Folding Status Controls Binding to BiP and Secretion from the ER In Vivo

It has long been appreciated that the C\textsubscript{H}1 domain is central to correct assembly and transport of IgG molecules and other immunoglobulin isotypes (Wolfenstein-Todel et al., 1974; Ade-tugbo, 1978; Hendershot et al., 1987; Shaffer and Schlissel, 1997). Deletion of this domain allows for secretion or surface expression of free heavy chains and various Ig assembly intermediates (Hendershot et al., 1987), which shows that quality control is focused on this domain. Our data put the unexpected unfolded nature of the C\textsubscript{H}1 domain at the center of the secretion control mechanism of IgG antibodies. To test this notion in a cellular context, we first expressed the MAK33 light chain, which contained the C\textsubscript{H}1 domain that was used in the in vitro experiments, in COS-1 cells and performed metabolic labeling and immunoprecipitation assays. As expected, this wild-type light chain (LC\textsubscript{wt}) was detected not only in the cells, but also in the medium, indicating that LC\textsubscript{wt} was secreted efficiently (Figure 4A, lanes 1 and 3). When we replaced the C\textsubscript{H}1 domain of the light chain with the C\textsubscript{H}1 domain, this light chain (LC\textsubscript{CH1}) now behaved like a heavy chain in terms of retention in the ER and interaction with BiP, as demonstrated by an increase in the amount of the altered light chain that coprecipitated with BiP and its absence in the medium (Figure 4A, lanes 4–6). This shows that the structural characteristics of the C\textsubscript{H}1 domain and its role in antibody retention are intrinsic, context-independent features. To more directly address the structural prerequisites for antibody retention, we exchanged the small helical elements of the C\textsubscript{H}1 domain, which have been reported to play a crucial role in the folding of this domain (Feige et al., 2008), against the corresponding elements of the C\textsubscript{H}1 domain. This exchange transformed the C\textsubscript{H}1 domain into an unfolded protein in vitro (data not shown). When a light chain containing this altered C\textsubscript{L} domain (LC\textsubscript{CLmut}) was expressed in COS-1 cells, it strongly interacted with BiP in vivo and was no longer secreted.
from the cell (Figure 4A, lanes 7–9), which argues that the folding status of an antibody domain is key for its retention.

To add support to the key role played specifically by the interactions between CH1 and CL domains, we first performed additional in vivo experiments with two different full-length heavy chains (the MAK33 g1 heavy chain and a humanized mouse IgG heavy chain [Liu et al., 1987]) in combination with both the wild-type (LCwt) and the mutated MAK33 light chain (LCCLmut). The wild-type MAK33 light chain assembled with both heavy chains and allowed their secretion from cells, whereas the mutant light chain was unable to assemble or induce secretion of the heavy chains (Figure 4B; Figure S5). This is in keeping with our in vitro data showing that the mutant CL domain was unfolded and therefore unable to induce the folding of the CH1 domain. To determine if this was also the case in vivo, we used a truncated version of the chimeric heavy chain consisting of only the VH and CH1 domains, which allows us to monitor oxidation of the CH1 domain based on an increase in its mobility (Lee et al., 1999). Indeed, we found that the wild-type MAK33 light chain induced oxidation and secretion comparable to a different light chain that was used in previous studies (Lee et al., 1999) (Figure 4C, compare lanes 4 and 6 and lanes 5 and 7). However, the mutant light chain was unable to induce either oxidation or secretion of this truncated heavy chain (Figure 4C, lanes 8 and 9). These data clearly show that the correct folding of the CL domain is absolutely required to induce the folding and oxidation of the CH1 domain as well as their covalent assembly in vivo, which is in line with our in vitro data. Furthermore, these data demonstrate that whereas both wild-type λ and κ light chains induced oxidation of the CH1 domain, assembly, and secretion, the light chain containing the CL mutant was defective in these processes.

Isomerization of a Conserved Proline Residue in the CH1 Domain Is Essential for Assembly and Secretion of IgG Molecules In Vivo

To directly assess the role of proline isomerization in the association-coupled folding reaction of the CH1 domain in vivo, each single cis proline residue in the CH1 domain was exchanged against alanine in the context of the full-length heavy chain, and metabolic labeling experiments were conducted. For the wild-type heavy chain and two of the heavy chains mutated in the CH1 domain, Pro34Ala and Pro74Ala, all Ig assembly intermediates could be detected (Figure 5A, lanes 1, 2, 4, 5, 10, and 11), and, importantly, completely assembled Ig molecules were secreted (Figure 5A, lanes 3, 6, and 12), demonstrating that isomerization of neither of these prolines was critical for
CH1 domain folding in vivo. In fact, mutation of Pro74 to alanine actually increased the assembly and secretion of the heavy chain. In contrast, when Pro32 in the CH1 domain was mutated to alanine, no significant amount of heavy-chain and light-chain assembly were detected, nor was any heavy chain secreted (Figure 5A, lanes 7–9). This is not due to poor expression of either the light chain or mutant heavy chain in this experiment (Figure 5B). These results are in excellent agreement with our in vitro data (Figure 2D) and reveal a key role of the trans to cis isomerization of Pro32 in the CH1 domain for the assembly, interchain disulfide bridge formation, and secretion of IgG molecules. It should be noted that this critical proline residue is highly conserved in the various murine Ig isotypes as well as in the immunoglobulins of different species.

DISCUSSION

In this work, we show that the C\textsubscript{H1} domain of the murine IgG1 antibody is an intrinsically disordered protein. As C\textsubscript{H1} does not possess an unusual number or distribution of charged or hydrophobic residues, it can still form a well defined globular structure once folded and does not show the typical sequence signature usually associated with intrinsically disordered proteins (Ward et al., 2004; Yang et al., 2005; Fink, 2005; Dunker et al., 2008). The large number of proline residues, also in a small helical part of the C\textsubscript{H1} domain that was recently identified as important for antibody domain folding (Feige et al., 2008), might be one determinant contributing to its unfolded nature. Our data thus suggest that the C\textsubscript{H1} domain is a representative of a distinct class of intrinsically disordered proteins.

We demonstrate that the C\textsubscript{H1} domain, which is the cognate partner of the C\textsubscript{L} domain in the complete IgG molecule, is required to fold C\textsubscript{H1} to the structure observed in IgG antibodies. The detailed analysis of the underlying pathway suggests that the reaction is initiated by the recognition of a few key interface residues between C\textsubscript{L} and C\textsubscript{H1}, which then promotes the formation of a hydrophobic core in C\textsubscript{H1}. Both reactions render each other energetically more favorable and thereby allow the entropically demanding structuring of an unfolded polypeptide chain. In vitro, this folding reaction requires the presence of the internal disulfide bridge in C\textsubscript{H1}. Therefore, one might assume that a roughly preformed topology or residual structure, which could not be detected in this study, might play a role in the folding process. Folding of the C\textsubscript{H1} domain is rate limited by proline isomerization, as observed for most isolated antibody domains (Goto and Hamaguchi, 1982; Thies et al., 1999; Feige et al., 2004). The presence of several cis proline residues in the native state and the overall very large number of proline residues, however, sets the C\textsubscript{H1} domain apart from most other antibody domains and may reflect the special role played by this domain. The key step in C\textsubscript{H1} folding, the isomerization of a single highly conserved proline residue in the loop between strands B and C, is likely to be essential for the folding and secretion of most Ig classes and potentially of other Ig superfamily members.

We show that BiP, a major ER chaperone, strongly binds to C\textsubscript{H1} in vitro, in agreement with previous in vivo studies (Hendershot et al., 1987). Oxidation of the internal C\textsubscript{H1} disulfide bridge was possible in the BiP-bound state in vitro, although this form has not been detected in vivo. After release from BiP, C\textsubscript{H1} can complete folding upon association with C\textsubscript{L} and, if successful, form an interchain disulfide bridge with C\textsubscript{L}. Because heavy chains of most isotypes that are devoid of the C\textsubscript{H1} domain can be secreted (Coffino et al., 1970; Hendershot et al., 1987), the essential steps controlling IgG assembly described here are likely to be general for all antibodies. Our data allow us to propose a possible order of events for this quality control mechanism in vivo. First,
the CH1 domain binds to BiP as it enters the ER cotranslationally in the reduced state. Then, likely triggered by association with light chain (Lee et al., 1999), the oxidation of the internal disulfide bridge between Cys25 and Cys80 takes place, which brings at least two of the residues (Val21 and Val78) that are involved in the formation of the hydrophobic folding nucleus in close proximity to each other. Only after release from BiP can the oxidized CH1 domain complete its folding in association with C. This scenario is in agreement with the fact that some residues in the CH1 domain found to be involved in the initial interaction with C were among those identified as putative BiP binding sequences in this domain in a previous study (Knarr et al., 1995). Even though most antibody domains possess BiP binding sequences (Knarr et al., 1995), they interact with BiP in the cell only transiently or not at all due to the competing, rapid folding reaction (Hellman et al., 1999). In contrast, the continued unfolded status of the CH1 domain in the absence of light-chain association allows it to permanently expose binding sites for BiP, predisposing it for a stable interaction in the ER. In the complete antibody, the C and the CH1 domain are covalently crosslinked via a disulfide bridge. We found that once this intermolecular disulfide bridge is formed, BiP no longer associates with the CH1 domain in vitro (data not shown), because formation of this interchain disulfide bridge is rate limited by proline isomerization and hence depends on the complete folding of CH1. Thus, folding-dependent, covalent assembly provides yet another checkpoint for monitoring the proper maturation of Ig molecules in the ER and allows the assembled heavy chain to escape thiol-mediated retention mechanisms in cells (Sitia et al., 1990). It is conceivable that in the ER, association, oxidation, and folding of CH1 are tightly coupled by the immunoglobulin assembly machinery, as no oxidized CH1 was found to be bound to BiP in vivo and ATP-induced release of BiP from unassembled heavy chains results in the formation of disulfide-linked heavy-chain aggregates (Vanhove et al., 2001).

Taken together, our data provide a detailed mechanism by which BiP and an intrinsically disordered antibody domain control the secretion of IgG antibodies (Figure 6). The comprehensive model incorporates previous in vitro (Lilie et al., 1995; Mayer et al., 2000) and in vivo findings (Bole et al., 1986; Hendershot et al., 1987; Lilie et al., 1995; Lee et al., 1999; Mayer et al., 2000; Vanhove et al., 2001) and suggests regulatory hubs at which additional components may come into play in vivo (Vanhove et al., 2001; Elkabetz et al., 2005) to orchestrate folding, oxidation, assembly, and secretion.

A rigorous assembly control mechanism is particularly important for antibody assembly control, which undergo a developmentally asynchronous expression of heavy- and light-chain genes. In pre-B cells, heavy-chain genes are rearranged first, and the resultant proteins are largely retained in the cell (Burrows et al., 1979), except for a limited number that assemble with the surrogate light chain (Pillai and Baltimore, 1987). The developmentally more mature B cell expresses light chains, which assemble with heavy chains and allow their transport to the cell surface. Finally, the terminally differentiated plasma cell produces enormous quantities of Ig molecules (Cenci and Sitia, 2007), which must be appropriately assembled to bind specifically to antigens and fulfill their effector functions. Accordingly, the nature of the reactions that govern CH1 folding and assembly with C, allow for the efficient and accurate assembly of antibodies prior to secretion and might hint toward the coevolution of substrates and folding factors in the ER as well as a general mechanism of quality control for oligomeric proteins.

**EXPERIMENTAL PROCEDURES**

**Protein Production**

All antibody domains were produced in a manner similar to published protocols (Feige et al., 2004, 2007). Details can be found in the Supplemental Data. Numbering of the CH1 domain begins with 1 in this work.

Hamster BiP (Wei and Hendershot, 1995) was mutated by site-directed mutagenesis to the murine sequence. Expression was carried out in HB101 cells for 3 h at 37°C. After cell disruption, Ni-NTA affinity purification was performed in 50 mM HEPES/ KOH (pH 7.5), 400 mM NaCl, 50 mM imidazole. BiP was eluted with an imidazole gradient from 0.05 M to 1 M. BiP containing fractions were applied to a Superdex 200 pg (26/60) gel-filtration column (GE Healthcare, Munich, Germany) equilibrated in HKM buffer (50 mM HEPES/KOH (pH 7.5), 150 mM KCl, 10 mM MgCl2) and finally to a Superdex 200 10/300GL HPLC column (GE Healthcare, Munich, Germany) equilibrated in the same buffer. The cyclophilin B (CypB) gene was amplified without the signal sequence from the murine cDNA (imaGenes, Berlin, Germany) and was inserted into the pET28a vector. Expression was carried out for 3 h at 37°C in BL21-DE3 cells. The cell pellet was dissolved in 50 mM HEPES/KOH (pH 7.0), 10 mM EDTA, and the cleared lysate was applied to a SP-Sepharose column equilibrated in the same buffer. The protein was eluted with a 0–1 M NaCl gradient. Subsequently, CypB-containing fractions were applied to a Superdex 75 pg (26/60) gel-filtration column equilibrated in HKM buffer. All vectors were sequenced, and protein masses were verified by mass spectrometry.

**Optical Spectroscopy**

A Jasco J-720 spectropolarimeter was used for all CD measurements (Jasco, Gross-Umstadt, Germany). Far-UV CD spectra were recorded in a 0.2 mm quartz cuvette; far-UV kinetics were recorded in a 1 mm quartz cuvette. Spectra of the isolated domains were recorded at 45 μM protein concentration; for the spectrum of the complex, 15 μM CH1 in the presence of 45 μM C, was used. Far-UV CD kinetics were recorded at 10 μM protein concentration of each domain and followed at 205 nm. Spectra of the CH1 domain in the complex were calculated by subtraction of the spectrum of the isolated CH1 domain from the spectrum of the complex, measured after a 4 hr equilibration step at 25°C. All spectra were averaged 16 times and buffer corrected. Fluorescence measurements were carried out in a Spex FluoroMaxIII spectrophotofluorometer (Jobin Yvon, Munich, Germany) in a stirred 1 cm quartz cuvette. Kinetics and titrations were measured by the change in the intrinsic tryptophan fluorescence, excited at 280 nm, and detected at 350 nm. For titrations, varying concentrations of CH1 were added to 2 μM C, and immediately as well as after a 4 hr equilibration step at 25°C, the fluorescence of the same samples was determined. The difference between initial and final fluorescence emission was normalized and analyzed according to a one-site binding model. Iodide quenching experiments were performed as published (Feige et al., 2007).

For anisotropy measurements, 1 μM lucifer yellow-labeled CH1 Ala113Cys and varying concentrations of CH1 were used. Lucifer yellow fluorescence was excited at 430 nm and detected at 525 nm. The change in quantum yield of the chromophore was less than 5% upon association of the labeled CH1 domain with CH1. Individual traces were fitted by single exponential functions. The obtained rate constants were fitted to a linear equation to derive the koff and the koff value.

**NMR Spectroscopy**

Spectra of the CH1 domain in complex with the C domain were recorded at 25°C on Bruker DMX600 and DMX750 spectrometers (Bruker, Rheinstetten, Germany), whereas spectra for the assignment of the unfolded CH1 domain were measured at 12.5°C on a Bruker AVANCE900 spectrometer (Bruker, Rheinstetten, Germany). Backbone sequential assignment of the isolated CH1 domain was obtained by standard triple-resonance experiments implemented.
with selective proton flip-back techniques for fast pulsing (Diercks et al., 2005). To gain information about any preferential conformations present in the disordered C_{H1} domain, an NNH-NOESY spectrum and a ^{15}N-HSQC-NOESY spectrum were recorded on a highly deuterated sample with a mixing time of 600 ms in order to detect long-range HN-HN NOEs (Mok et al., 1999).

For all measurements of the folded C_{H1} domain in association with C_{L}, a 2-fold excess of unlabeled C_{L} was added to ^{15}N or ^{15}N, ^{13}C-labeled C_{H1}. Prior to steady-state measurements, samples were incubated for at least 6 hr at room temperature to ensure complete folding of the C_{H1} domain. Backbone sequential assignment of the assembled C_{H1} domain was achieved with standard triple-resonance experiments with selective proton flip-back techniques for fast pulsing. The assignment of the carbon chemical shifts was limited to the C_{0} and C_{a} chemical shifts due to the relaxation properties of the whole protein complex. To verify the backbone resonance assignment, NH residual dipolar couplings (RDCs) were determined. The sample was prepared as described above and aligned with nonionic liquid crystalline media (Ruckert and Otting, 2000). NH RDC values were extracted from IPAP-HSQC spectra by using Bruker pulse sequences. The sequential information based on the C' and C'' chemical shifts as well as the NH RDC values and the crystal structure of the folded C_{H1} domain (PDB code: 1ORS) served as input for the software MARS (Jung and Zweckstetter, 2004a, 2004b).

In order to characterize the folding pathway of the intrinsically disordered C_{H1} domain, ^{15}N-HSQC spectra were recorded at 12.5 °C every 14 min immediately after adding unlabeled C_{L} to ^{15}N-labeled C_{H1} by using selective proton flip-back techniques for fast pulsing. Identical processing of all spectra measured during the folding process was performed with the software TOPSPIN 1.3 (Bruker Biospin). Peak intensities were analyzed by using the software SPARKY (www.cgl.ucsf.edu/home/sparky) and normalized to the corresponding intensities in the final spectrum after 36 hr. The backbone resonance assignment was transferred from 25°C to 12.5°C recording a HSQC temperature series.

**Analytical HPLC Experiments**

For all experiments, a Shimadzu HPLC system (Shimadzu, München, Germany) was used. Complex formation between BiP and C_{H1} was analyzed on a Superdex 200 10/300GL column in HKM buffer at a flow rate of 0.5 ml/min. For the determination of the dissociation constant respectively binding kinetics, peak intensities at the retention time of 28.4 min corrected for baseline drifts were

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**Figure 6. A Model for the Overall IgG Secretion Control Mechanism**

A schematic indicating the possible pathways for the C_{H1} domain (blue); its folding and assembly in association with C_{L} (green) and BiP (gray) is shown. C_{H1} has to form its internal disulfide bridge and to be released from BiP before it can associate with C_{L}. In vivo, these processes are tightly coupled and thus cannot be dissected kinetically. Prior to complete folding and irreversible formation of the C_{L}/C_{H1} interchain disulfide bridge, the proline residue 32 has to isomerize from trans to cis and otherwise traps a partially folded C_{H1} intermediate. The isomerization reaction can be accelerated by cyclophilin B. All rate constants were determined at 25°C.
plotted against the C\textsubscript{\text{1}} concentration or the incubation time, respectively, and were normalized. The rate constants, \(k_{\text{on}}\) and \(k_{\text{off}}\), of the binding reaction were determined from single exponential fits and evaluated with a linear equation to derive \(k_{\text{on}}\) and \(k_{\text{off}}\). Detection of all proteins was performed by the intrinsic fluorescence excited at 280 nm and monitored at 350 nm. Incubation steps were performed in HKM buffer with 1 mM ADP.

**Cell Culture Experiments**

The murine IgG1 MAK33 light-chain (LC\textsubscript{WT}) and heavy-chain (HC\textsubscript{WT}) cDNAs were obtained with an intact signal sequence for expression in mammalian ER. Two LC mutants, one in which the C\textsubscript{L} domain was substituted for the C\textsubscript{c} domain (LC\textsubscript{CLmut}) and the other in which structural features of the C\textsubscript{L} domain were substituted for the corresponding regions of the C\textsubscript{c} domain (LC\textsubscript{CCmut}) were produced, and all constructs were inserted into the pSVL vector (GE Healthcare, München, Germany). An HA epitope tag was engineered at the C terminus of the wild-type light chain and the two mutants for immunoprecipitation purposes. Heavy-chain proline exchange mutants were generated by site-directed mutagenesis. The cDNA for a chimeric humanized heavy chain was used as published (Liu et al., 1987), and a truncated version of this heavy chain containing only the VH and CH1 domains was produced previously (Lee et al., 1999), as was a mouse lambda light-chain cDNA (Hellman et al., 1999). The recombinant plasmids, along with a pMT vector encoding hamster BiP (Lee et al., 1999), were introduced into COS-1 cells (Gluzman, 1981) that were cultured as described (Lee et al., 1999) by using FuGENE 6 transfection reagent (Roche, Indianapolis, USA) following the manufacturer’s protocol. Metabolic labeling, cell lysis, immunoprecipitation, and visualization of the proteins were performed as described previously (Lee et al., 1999). Anti-rodent BiP antiserum (Hendershot et al., 1995), a monoclonal anti-HA (12CAS) antibody (kindly provided by Dr. Al Reynolds, Vanderbilt University, USA), goat anti-mouse Ig\textsubscript{g} and goat anti-mouse \(\gamma\) antibodies (Southern Biotech, Birmingham, AL, USA), and Protein A Sepharose beads were used for immunoprecipitations. For metabolic labeling experiments, cells were cultured in DMEM lacking methionine and cysteine and labeled with \(^{35}\text{S} \) Translabel (MP Biomedicals, Irvine, CA, USA) for the indicated times.

**SUPPLEMENTAL DATA**


**ACKNOWLEDGMENTS**

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Molecular Cell

Unfolded \( C_H \) Controls IgG Assembly and Secretion


Supplemental Data

An Unfolded C\textsubscript{H}1 Domain Controls the Assembly and Secretion of IgG Antibodies

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Supplemental Experimental Procedures

**Protein production.** The C\textsubscript{H}1 domain (Thr123-Arg215 of the pdb file 1FH5; numbering starts with one in this work) and an Ala113Cys mutant of the C\textsubscript{L} domain were amplified from the murine IgG1 MAK33 cDNA. Additionally, the murine IgG1 MAK33 cDNA was used as a template to amplify the gene segments encoding the C\textsubscript{L} domain with the native C-terminal cysteine residue (amino acids Ala112-Cys215 of the pdb file 1FH5) and the C\textsubscript{H}1 domain with the native C-terminal cysteine residue (Thr123-Cys217 of the pdb file 1FH5). All genes were cloned into the pET28a (Novagen, Gibbstown, NJ, USA) vector without a tag, except for the wild type C\textsubscript{L} domain which was essentially purified as published (Feige et al., 2007). Isotope labeled C\textsubscript{H}1 for NMR experiments was expressed in M9 minimal medium. For FRET measurements, the Ala113Cys mutant of the C\textsubscript{L} domain was used. In the case of C\textsubscript{H}1, the native C-terminal cysteine residue was used for labeling in FRET experiments. Proteins were expressed as inclusion bodies overnight at 37°C in BL21-DE3 cells in selective LB medium. Inclusion bodies were isolated as described (Feige et al., 2004). Inclusion bodies were solubilized in 50 mM Tris/HCl, pH 7.5, 10 mM β-mercaptoethanol, 10 mM EDTA, 8 M urea and subsequently applied to a Q-
Sepharose column equilibrated in 50 mM Tris/HCl, pH 7.5, 10 mM EDTA, 5 M urea. All proteins of interest did not bind to the column under these conditions. Refolding was carried out by dialysis as published (Feige et al., 2007). After refolding, all proteins were applied to a Superdex 75pg (26/60) gel filtration column (GE Healthcare, München, Germany) equilibrated in PBS. Labeling of C\textsubscript{L} Ala113Cys with lucifer yellow iodoacetamide (Invitrogen, Carlsbad, CA, USA) was performed according to the manufacturer's protocol. For FRET experiments between BiP, C\textsubscript{L} and C\textsubscript{H1}, C\textsubscript{L} Ala113Cys was labeled with the ATTO532 maleimide dye (ATTO-TEC, Weidenau, Germany), and murine BiP, which possesses two free cysteines, was labeled with ATTO594 maleimide. For FRET experiments between BiP and C\textsubscript{H1}, C\textsubscript{H1} was labeled with ATTO532 at its native C-terminal cysteine residue. All proteins were separated from the free dye via a Superdex200 10/300GL HPLC column equilibrated in HKM buffer. All vectors in this study were sequenced and protein masses were verified by mass spectrometry.

**Preparation of reduced C\textsubscript{H1}**. To reduce the internal disulfide bond of the C\textsubscript{H1} domain, the protein was unfolded in 2 M GdmCl in PBS at 25°C for 2 h in the presence of 10 mM TCEP. Subsequently, the protein was applied to a Superdex75 10/300GL HPLC column equilibrated in PBS supplemented with 0.1 mM TCEP.

**Optical spectroscopy**. A Jasco J-720 spectropolarimeter was used for all measurements (Jasco, Gross-Umstadt, Germany). A 2 mm quartz cuvette was used for near-UV CD spectra and kinetics (observed at 280 nm). 100 µM protein respectively 100 µM of each domain were used. Spectra of the C\textsubscript{H1} domain in the complex were calculated by subtraction of the spectrum of the isolated C\textsubscript{L} domain.
from the spectrum of the complex, measured after a 4 h equilibration step at 25°C. All spectra were averaged 16 times and buffer corrected.

**Analytical HPLC experiments.** For all experiments, a Shimadzu HPLC system (LC-20AT, SPD-20A, RF-10Axl, SIL-20AC) (Shimadzu, München, Germany) was used. To assess complex formation between C_L and C_H1, both proteins were incubated at 25°C with a domain concentration of 25 µM each. After varying times, 100 µl of the sample were applied to a Superdex75 10/300GL HPLC column (GE Healthcare, München, Germany) in PBS at a flow rate of 0.75 ml/min. The peak height corresponding to the dimer was analyzed over incubation time. To assess whether C_H1 can be oxidized if bound to BiP, C_H1 was reduced in HKM buffer with 2 mM TCEP for 1 hour. Complex formation between 30 µM reduced C_H1 and 50 µM Bip was performed at 37°C for 2 hours in HKM buffer supplemented with 1 mM ADP and 1 mM TCEP. The sample was applied at a flow rate of 0.25 ml/min to a Superdex 200 10/300GL column equilibrated in HKM buffer supplemented with 4 mM GSSG and 1 mM TCEP. Only the peak of BiP-bound C_H1 was collected and free sulfhydryls were reacted with 10 mM NEM for 15 minutes at room temperature. The samples were subsequently analyzed by MALDI-TOF/TOF mass spectrometry which showed peaks for oxidized as well as NEM-reacted C_H1. As a control, reduced, BiP-bound C_H1 was reacted under the same conditions with NEM and only showed peaks for NEM bound C_H1.

**SDS-PAGE experiments.** To assess the formation of the disulfide-bridged C_L/C_H1 dimer, both proteins were co-incubated at a concentration of 25 µM each in PBS at 25°C. If present, the concentration of CypB was 5 µM. At the indicated times, a 20 µl aliquot was withdrawn, supplemented with 10 µl 3x Laemmli sample buffer (without
β-mercaptoethanol) and boiled for 1 min at 95°C. Subsequently, 20 µl of each sample was applied to a 17.5% SDS-PAGE gel and run for 75 min at 25 mA. The bands were quantified with ImageQuant TL (GE Healthcare, München, Germany).

Supplemental References


Supplemental Figure Legends

Figure S1. Formation of tertiary and quaternary structure upon C_H1/C_L association. (A) The isolated C_L domain shows a well defined near-UV CD spectrum (cyan), whereas the isolated C_H1 domain shows the featureless spectrum of an unfolded protein (blue). The complex between C_L and C_H1 is shown in green and the spectrum of the C_H1 domain in the complex in red. The change in near-UV CD signal at 280 nm upon the co-incubation of C_L and C_H1 is shown in (B). The trace could be fitted with a single exponential function with a time constant of \( \tau = 50 \pm 2 \) min (C). Formation of stable C_H1/C_L quaternary structure was assessed by analytical HPLC-experiments using 25 \( \mu \)M of each protein. The elution profiles after 1 min of co-incubation of C_L and C_H1 (dashed line) and after 5 h of co-incubation (straight line) are shown. In (D), the normalized intensity of the peak corresponding to the C_L/C_H1 dimer (15 min) is plotted over time. It could be fitted with a single exponential function \( (\tau = 65 \pm 5 \) min). All measurements were performed in PBS at 25°C.

Figure S2. Influence of the redox status of the C_H1 domain on its folding status and its association-coupled folding process. (A) The far-UV CD spectrum of the reduced C_H1 domain is characteristic of an unfolded protein. (B) 5 \( \mu \)M oxidized (straight line) or reduced C_H1 (dashed line) were added to 1 \( \mu \)M lucifer yellow labeled C_L. The change in anisotropy over time could be well described as a single exponential function for oxidized C_H1 \( (k_{obs} = 0.19 \) min\(^{-1}\)), for reduced C_H1 no change was observed. (C) For oxidized C_H1 (straight line), folding in the presence of C_L could
be observed by far-UV CD-spectroscopy at 205 nm (τ = 44 ±4 min) whereas no structure formation was detectable for the reduced C\textsubscript{H1} domain (dashed line). A concentration of 10 µM was used for each protein. All measurements were carried out at 25°C in PBS in the presence of 30 µM TCEP.

**Figure S3.** C\textsubscript{L}/C\textsubscript{H1} association coupled to intermolecular disulfide bridge formation. (A) The isolated C\textsubscript{L} domain with the C-terminal cysteine residue shows an all-\textbeta far-UV CD spectrum (cyan), whereas the isolated C\textsubscript{H1} domain with the C-terminal cysteine residue displays a random coil spectrum (blue). The complex of both proteins is shown in green. The spectrum of the C\textsubscript{H1} domain, calculated from the individual spectra of C\textsubscript{L} and the C\textsubscript{H1}/C\textsubscript{L} complex, reflects folding of the C\textsubscript{H1} domain in the presence of C\textsubscript{L} (red). (B) Folding of the C\textsubscript{H1} domain, followed by far-UV CD spectroscopy at 205 nm, occurred with a time constant of τ = 53 ±5 min in the absence of CypB (red trace). It could be accelerated to τ = 38 ±4 min in the presence of 2 µM CypB (blue trace). The acceleration could be inhibited by 4 µM CspA (not shown). All measurements were carried out in PBS at 25°C with C\textsubscript{L} respectively C\textsubscript{H1} containing their native C-terminal cysteine residue.

**Figure S4.** Assessment of the formation of triple complexes between BiP, C\textsubscript{H1} and C\textsubscript{L}. To investigate the possibility that either C\textsubscript{L} could associate with BiP bound C\textsubscript{H1} or if the C\textsubscript{L}/C\textsubscript{H1} complex binds to BiP, 1 µM ATTO532 labeled C\textsubscript{L} was incubated with 10 µM ATTO594 labeled BiP (A). The donor, ATTO532, was excited at 500 nm and the donor fluorescence was recorded at 550 nm (green line), the fluorescence of the acceptor, ATTO594, was recorded at 625 nm (red line). After 30 min, where no binding of C\textsubscript{L} to BiP could be detected as expected (data not shown), 10 µM
unlabeled CH1 was added. Under these conditions, CH1 binds to BiP within several minutes. If either CL could bind stably to the BiP:CH1 complex or if BiP could bind to the CL/CH1 complex, a FRET signal between BiP and CL is expected. However, this was not observed arguing against the presence of BiP:CH1:CL triple complexes. As a control of the FRET system, the association between 10 µM ATTO594 labeled BiP and 1 µM ATTO532 labeled CH1 was measured (B). The association reaction occurred with a rate of $k_{obs} = 0.01 \pm 0.005 \text{ min}^{-1}$. In (C, left) a putative triple complex between BiP (grey, model based on the pdb code 3C7N, chain B), CH1 (purple) and CL (blue) is shown. The labels are schematically shown in green (ATTO532) respectively in red (ATTO594) and distances are indicated. The Förster radius of the dye pair is 6.6 nm (www.atto-tec.de). On the right, a putative complex between BiP and CH1 is shown. All measurements were performed at 25°C in HKM buffer with 1 mM ADP and donor fluorescence was linearly corrected for bleaching.

**Figure S5. A folded CL domain is required for the proper maturation and secretion of heavy chains from cells.** COS-1 cells were co-transfected with vectors encoding BiP, the HA-tagged versions of either wild type light chain (LCwt) or the mutant light chain containing an unfolded CL domain (LCCLmut) and either a humanized chimeric γ heavy chain or the MAK33 γ heavy chain. After 24 h, cells were metabolically labeled and both cell lysates (no subscript) and culture supernatants (subscript m) were immunoprecipitated with the indicated antisera. Precipitated proteins were separated by SDS-PAGE under non-reducing condition and visualized by autoradiography.
Supplemental Figures

Figure S1
Figure S2

A

\[ \Theta_{\text{MW}} \times 10^3 \text{ (deg cm}^2\text{ dmol}^{-1}) \]

wavelength (nm)

B

change in anisotropy (a.u.)

time (min)

C

CD-signal (mdeg)

time (min)
Figure S3

A

\[ \theta_{MRV} \times 10^{-3} \, (\text{deg cm}^2 \, \text{dmol}^{-1}) \]

wavelength (nm)

B

CD-signal (mdeg)

time (min)
Figure S4

A

B

c

104
Figure S5
VI. Published book chapters
CHAPTER 2

Oxidative Folding of Proteins in vitro

CHAPTER 2.1

The Role of Disulfide Bonds in Protein Folding and Stability

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2.1.1 Introduction

Disulfide bonds add covalent cross-links to the linear polypeptide chain. It is therefore intuitive to assume that this posttranslational modification has a pronounced impact on the folding and stability of proteins. These covalent linkages are found mostly in extracellular proteins where they contribute significantly to the stability of the respective protein. An extension to the intrinsic stabilization of extracellular proteins by disulfide bridges is the stabilization of quaternary structure by intermolecular cystine links. This is an effective strategy to prevent dissociation in the extracellular environment which, due to the often low concentrations, would in many cases be irreversible. However, in the context of discussing the effects on protein structure and folding, these disulfides are set apart from the intra-chain disulfides.
Interestingly, disulfide bonds in proteins are highly conserved; only Trp residues are even more conserved.¹ The natural selection for disulfide bonds in secreted proteins seems to be due to the simple and reversible redox chemistry involved in their formation and breaking and their intrinsic stability. Only at extreme conditions such as 100°C and alkaline pH values do they spontaneously break or rearrange.² Disulfide bridges come in different flavors, reflecting the different purposes they can fulfill in a protein. In addition to their effect on the global stability of a polypeptide chain, they may have important effects on the structure, stability and dynamics of local structural elements. This is especially evident for the so-called cysteine knots, in which disulfide bridges cross each other and thus determine the local topology³ and for “allosteric” disulfide bonds which serve to regulate protein function.⁴ In this review we will focus exclusively on the dissection of the effects of structural, in particular non-local, disulfide bonds on protein stability and folding.

The effect a disulfide bond will have on a given protein will of course depend on the position of the disulfide bond in the structure of the protein. The complexity of the influence of disulfide bonds is reflected by the finding that stabilizing disulfide bonds can be found either in the core of proteins as such or surface-localized. An important issue in this context is the distance between two Cys residues in the sequence of a polypeptide chain. As will be pointed out in this chapter, the stabilizing effect depends to a significant extent on the number of amino acids spanned between the two residues. In agreement with this notion the average distance of two Cys residues in proteins was found to be 15 amino acids.¹

Disulfide-bonded proteins have been studied early on in the history of protein folding. Notably, one of the first proteins analyzed extensively, RNase A, contains several disulfide bridges. In seminal work, Christian Anfinsen showed that the completely reduced and denatured protein could spontaneously regain its native structure including the correct disulfide bonds.⁵ Efforts to analyze the impact of disulfide bonds on protein folding and stability mechanistically have contributed enormously to the generation of concepts for these processes. Furthermore, engineered disulfide bonds have become valuable tools to explore them in detail. In this chapter we will discuss the basic models resulting from these studies and we will outline the thermodynamic and kinetic implications the presence of a disulfide bond has on unfolded, folding and folded proteins.

### 2.1.2 Stabilization of Proteins by Disulfide Bonds

In the early days of protein science, the native state of a protein was believed to be one, rather fixed, conformation and the unfolded state to be a random coil.⁶ An ideal random coil is devoid of any long-range interactions except excluded volume effects. It behaves as a freely joined chain of segments of defined length.⁷ In this description, the impact of a covalent cross-link between two defined residues of the polypeptide chain, such as a disulfide bridge, was thought to be exclusively on the unfolded state. The fixed geometry of the native state should be left essentially unaltered by a bond between two residues which are in proximity, yet the freedom
of the random coil polypeptide should be significantly decreased. Restricting the conformational space of the unfolded state clearly reduces its entropy. Hence, the entropy change for the reaction to the ordered native state was thought to be less negative with a net stabilization of the folded protein as a result. A quantitative description of the phenomenon was developed by Schellman, Flory, Poland and Scheraga.8–10 The decrease in entropy of the unfolded state is derived from the probability that two otherwise free elements of the chain are now found in a defined volume element (v). Mathematically and based on polymer theory, the problem is described in Equation (2.1.1) (where R is the gas constant, l the average length of a statistical segment of the chain composed of N segments; in proteins l is assumed to be 3.8 Å corresponding to one amino acid):

\[
\Delta S = -R \ln \left[ \frac{3}{(2\pi l^2 N)^{3/2}} \right] v
\]  

(2.1.1)

A major point of discussion has been the adequate choice of v. A value of 57.9 Å³ based on the closest possible approach of two thiols is still mostly in use.11 Hence, Equation (2.1.1) can be simplified to Equation (2.1.2) where n is the number of amino acids bridged by the disulfide bond:

\[
\Delta S = -2.1 - 3/2 \ R \ln(n)
\]  

(2.1.2)

Based on a study of Ribonuclease T1 with zero, one and two intact disulfide bonds, Equation (2.1.2) was developed by Pace et al.11 The authors did not only find a good correlation between n and ΔΔG upon removal of disulfide bonds in RNase T1, but additionally agreement between experimental data for lysozyme, RNaseA and the antibody C_L domain.11

The above equations have two consequences. Conceptually, the stabilization of a protein is thought to be an entirely entropy-driven process with an impact exclusively on the random coil unfolded state. Practically, the stabilization exerted by a disulfide bridge should increase with the number of amino acids between the two cysteines. This theory treats the unfolded polypeptide chain as a system devoid of any intra- or intermolecular interactions. However, the water surrounding a protein is an important factor shaping the free energy landscape of the polypeptide chain during folding and in the native state.12 Sometimes it can be regarded as being a part of the native structure.13 The impact of this scenario with respect to disulfide bonds has been addressed by Doig and Williams in a widely recognized publication in 1991.14 The authors argued that disulfide bonds might lead to a significantly decreased solvent accessible surface in the unfolded state. Hence, hydrophobic residues as well as hydrogen-bond donors and acceptors might become buried. This is supposed to lead on the one hand to a larger entropy of the solvent in disulfide-containing proteins due to buried hydrophobic residues. Consequently, the hydrophobic effect should be less pronounced for these proteins. On the other hand, hydrogen bonding to water in the more compact unfolded state will be possible to a lesser extent, and hence folding to the native state will be enthalpically more favorable. According to the authors, the enthalpic contribution has to be considered as the major stabilizing factor of
disulfide bridges. The model will therefore be called *solvent enthalpy model* in the subsequent paragraphs. As in the *chain entropy model*, eventually occurring differences in the native state such as induced strain or reduced dynamics are also neglected in this model. Both models are summarized in Figure 2.1.1 where the reduced conformational freedom of a disulfide-linked polypeptide chain is visualized as well as hydrophobic clustering in the unfolded state or reduced solvent-protein interactions which might be present for a disulfide-linked protein.

**Figure 2.1.1** Graphical representation of the chain entropy and the solvent enthalpy model. In the chain entropy model, the major stabilizing factor exerted by a disulfide bridge is believed to be the conformational restriction of the unfolded state which renders the folding reaction entropically less unfavorable (A). In the solvent enthalpy model (B), two major effects are attributed to disulfide bridges. First, hydrophobic interactions are believed to be more pronounced in unfolded, disulfide-bridged proteins decreasing the hydrophobic effect for a protein folding reaction. Second, and numerically more important, disulfide bridges are assumed to inhibit hydrogen bonding to water in the unfolded state, making the folding reaction enthalpically more favorable. In both theories, the native state is believed to be unaffected. To illustrate both models, a hypothetical 12-residue \( \beta \)-hairpin structure with a single disulfide bridge was designed.
A variety of experimental evidence argues against both theories rendering the problem much more complex yet more insightful. Site-specific mutagenesis offered the possibility to introduce artificial disulfide bonds at defined positions within a protein and allowed to explore the effects of disulfide bonds rigorously.\textsuperscript{15–18} Many results of these studies were not compatible with predictions based on the prevailing theories concerning the effects of disulfide bonds on protein stability. Most discrepancies arose from efforts to stabilize a variety of proteins by the introduction of new disulfide bridges. In contrast to expectations, destabilization was the result of a significant proportion of these attempts.\textsuperscript{19–21} High-resolution structures of the engineered proteins helped to rationalize these unexpected effects. It turned out that strain was imposed on the native state by the disulfide cross-links in several cases.\textsuperscript{22} Two particularly insightful examples are studies on barnase\textsuperscript{19,23} and staphylococcal nuclease.\textsuperscript{21} Clarke and Fersht introduced three artificial disulfide bonds into barnase. One was found to destabilize the native state, one stabilized it to an extent predicted by the chain entropy model and one to a much lesser extent.\textsuperscript{19} Furthermore, the bridge connecting fewer residues was found to be more stabilizing than the one encompassing more residues. For staphylococcal nuclease, no stabilization was found for all disulfide constructs, but in this case a peptide bond \textit{cis/trans} equilibrium was shifted in the native state and the catalytic activity of all mutants was reduced.\textsuperscript{21} The NMR H/D exchange analysis of the barnase mutants revealed an altered dynamics of the native state imposed by the presence of the covalent cross-links.\textsuperscript{23} In the study on staphylococcal nuclease, strain on the native state, as reflected by the alteration of the \textit{cis/trans} equilibrium, was evoked to explain the observed unexpected alterations.\textsuperscript{21} A more general comparison between experimental data for disulfide-bridged proteins with native or engineered disulfide bonds and the chain entropy model, which predicts that disulfide bonds stabilize proteins in a strictly loop length dependent manner, can be found in Figure 2.1.2. A clear deviation between experimental data and predictions is evident from Figure 2.1.2 for the major part of the proteins.

\textbf{Figure 2.1.2} Comparison of experimental data and theory for the chain entropy model. Experimental data on the stability of proteins in the reduced and oxidized state are summarized in the table on the left.\textsuperscript{11,20,70,75} A comparison to the predicted stabilization based on Equation (2.1.2) is shown on the right. The chain entropy model predicts a larger stabilization \textit{via} disulfide bridges at higher temperatures and longer loop lengths.
One clear conclusion can be drawn from the two mentioned and a variety of other studies: the native state is not left unaffected by a disulfide bridge. A major assumption underlying the chain entropy model as well as the solvent enthalpy model hence does clearly not hold in all cases. Strain is often found to be imposed on the native state by the presence of covalent cross-links. This can be reflected in structural changes or more subtle changes, like alterations of the dynamics of the native state. The important consequence is that the enthalpy as well as the entropy of the native state are not unlikely to be altered if two residues in the polypeptide are covalently cross-linked. Furthermore, the alterations in dynamics and structure are not always a global but sometimes a local context-specific effect. β-sheets and loops are thought to be more suited to dissipate induced strain than α-helical elements, and more dynamic parts of the structure are influenced to a greater extent. In addition, changes in the solvent-exposed hydrophobic surface which have been observed for disulfide mutants of interleukin-4 have a significant effect on the enthalpy of the native state. Changes in the dynamics, e.g. the vibrational normal modes of a protein, will also influence the entropy of the native state as has been shown in a molecular dynamics study by Karplus and co-workers. Consequently, a net destabilization due to a loss of native state entropy can be expected in some cases if disulfide bonds are introduced into a protein. In summary, novel experimental as well as theoretical insights clearly argue against the simple chain entropy model, at least concerning one side of the equation, the native state. The same holds for the solvent enthalpy model by Doig and Williams, which is additionally at odds with thermodynamic parameters derived for some disulfide-bridged proteins which showed that disulfide bridges do not necessarily stabilize the native state enthalpically. But what about the unfolded state? Do the assumptions underlying the theory hold? In other words, is it correct to assume a random coil almost devoid of any interactions except for hydrogen bonding to the solvent? Clearly not in all cases. Residual structure which is believed to be important for folding pathways has been detected in a variety of proteins. In particular, residual hydrophobic interactions or fluctuating α-helical elements seem to be a rather general than an exceptional feature of proteins, in particular under mildly denaturing conditions. Importantly, the structural features of the unfolded state like residual structure, which has been reported for the unfolded state of barnase, or the tendency towards irreversible inactivation have in some cases been shown to be influenced by the presence or absence of disulfide bridges. Hence, disulfide bonds very likely not only influence the conformational freedom of the unfolded chain but can also introduce structure which might also be protective against irreversible aggregation. Even apparently minor structural changes like clustering of some hydrophobic residues will influence the enthalpy and entropy of the unfolded state. In addition, as outlined above, dynamic phenomena, strain induced on the native state and an impact on the native structure are completely omitted in these theories. Although some proteins could be adequately described by one of the theories, it comes as no surprise that the chain entropy model as well as the
solvent enthalpy model fail to describe the major part of experimental data (Figure 2.1.2).

How can the different findings on the divergent effects disulfide bridges have on the stability of different proteins be summarized in a comprehensive way? Hardly by any theory describing the unfolded polypeptide chain as a construct composed of identical elements and devoid of structural features. The decreased entropy of an unfolded and cross-linked polypeptide chain has to be taken into account as developed in the chain entropy model. Furthermore, the solvent enthalpy model based on the decreased hydrophobic effect and hydrogen bonding in the unfolded state should be included. And to be added to these models are interactions eventually present in the unfolded state due to the disulfide bridge, which may not be localized directly around the bond but can be present as long-range residual structure. This will clearly have an effect on the enthalpy and entropy balance for the reaction to the native state. The same holds for decreased dynamics of the native state, locally or globally, as well as enthalpically unfavorable strain or enthalpically favorable induced proximity of interacting residues. In summary, the effect of a disulfide bridge on the stability of a protein can be easily assessed experimentally, yet its molecular explanation is almost as diverse as the protein under investigation. The key factors giving rise to the net effect are most likely all known but, as for the prediction of the native state of a protein, their contribution to the overall effect are blurred in their sum as well as their mutual influence on each other. It seems therefore highly rewarding to use a combined empirical and theoretical approach. Exact stability data, if possible together with structural data on the native as well as the unfolded state of the protein under investigation, are a prerequisite for the detailed understanding of the effect of a disulfide bridge. They should be complemented by theoretical approaches like molecular dynamics simulations of the native and the unfolded state to obtain a more complete picture. This might be different in detail for different proteins but in summary these analyses will most likely reveal general principles.

2.1.3 Disulfide Bonds in Protein Folding Reactions

One of the major questions in biophysical chemistry is how a linear polypeptide chain specifically adopts its intricate three-dimensional structure in a reasonable amount of time. A variety of mutational approaches, high-resolution structural techniques and ultrafast perturbation methods have recently provided deep insight into this phenomenon of biological self-organization. Nowadays, proteins are believed to fold with a certain heterogeneity via multiple individual pathways to the native state, yet distinct features of the protein dictate the general trajectories of the folding process. In particular, residues whose interactions define the overall topology of a protein are assumed to be in contact in the transition state and are believed to be key elements in a protein folding reaction in general. While there seems to be consensus concerning the general scheme of events, a variety of details are still under intense debate, like
the role of residual structure in the unfolded state, the role of folding intermediates on the way to the native state and the heterogeneity of the transition states. In all these questions, disulfide bridges have played a pronounced role as factors influencing these processes as well as tools to elucidate them.

Independent of its pathway, every protein begins its folding reaction in the unfolded state, be it at the ribosome or under denaturing conditions. Therefore, the nature of the unfolded state of proteins has gained much attention again recently, in part fuelled by the development of novel experimental techniques. In this context, deviations from a random coil are regarded as important elements of protein folding in general. They will not only have an impact on the net stability balance of a protein, as outlined above, but furthermore likely shape the energy landscape on the way to the native state. Preformed interactions in the unfolded state might therein have a rather controversial effect on a folding reaction. If native and not detrimentally influencing the ability of the remaining polypeptide chain to explore the necessary conformational space, they might lead to faster and more efficient folding to the native state. Yet if non-native or too stable, the opposite might hold. Examples for both cases have been reported for disulfide-bridged proteins, which per se possess preformed correct tertiary interactions. In the case of RNase T1 (Figure 2.1.3), deceleration of the folding kinetics was observed which has been attributed to decreased chain flexibility in the presence of a disulfide bridge. In this context it is important to note that RNase T1 possesses two disulfide bridges, one connecting a small N-terminal β-turn and one connecting the C-terminus to this N-terminal β-turn (Figure 2.1.3). Accordingly, the protein will almost be a completely looped structure in its unfolded state, which will clearly have an effect on the dynamics of the whole polypeptide chain and might interfere with the establishment of a folding nucleus, if not in proximity of the disulfide bridges, which is unlikely due to the solvent exposure of the disulfide bridges in the native state. This has been confirmed for one of the disulfide bonds connecting residues 2 and 10.

Figure 2.1.3 Structural comparison of RNase T1 and the murine C domain. RNaseT1 (A) possesses two solvent-exposed disulfide bridges. The murine immunoglobulin G C domain has one disulfide bridge buried in the hydrophobic core of the protein (B).
Its deletion led to a stability reduction of RNase T1 but had no impact on the folding mechanism of the protein.\textsuperscript{66}

For C\textsubscript{L}, the constant domain of the antibody light chain, on the other hand, its single intrinsic disulfide bridge accelerated folding up to \(\sim 100\)-fold (Figure 2.1.3).\textsuperscript{67,68} In the case of this protein, the disulfide bond is found in the hydrophobic core of the protein and part of the folding nucleus.\textsuperscript{68} Accelerated folding due to preformed interactions which facilitate the way to the native state were hence expected and experimentally observed. For many other proteins, either a deceleration or an acceleration of the respective folding reactions have been observed in the presence of their natural intrinsic disulfide bonds.\textsuperscript{20,69–74} Hence, disulfide bonds are in general far from being inert in kinetic terms. Very often, different disulfide bridges within the same protein had different effects on the folding rates. A comprehensive study in this respect has been carried out for the all-\(\beta\) protein CD2, where thirteen different artificially introduced disulfide bridges showed a markedly different impact on the folding behaviour.\textsuperscript{20} Similar results were obtained for unfolding reactions which were either left unaffected by disulfide bonds or their rates were reduced.\textsuperscript{20,75} The experimental findings are in agreement with simple lattice-based simulations, where disulfide bonds inside the folding nucleus were found to be accelerating yet decelerating if outside.\textsuperscript{76} In some simulations folding has been found to be influenced by a disulfide bond to a lesser extent than expected, revealing a larger dynamics of a real polypeptide chain in comparison to a chain moving on a lattice.\textsuperscript{76} Despite the heterogeneous effects of a disulfide bridge on protein folding/unfolding kinetics at first glance, it can be more easily rationalized than the effect on native state stability. The key lies in the transition state for folding and unfolding. Acceleration of folding is expected when residues are cross-linked which have to come into contact early in a protein folding reaction, if not weighed out by entropy/enthalpy compensation in the transition state. Analogously, deceleration of unfolding is expected when two residues, whose interactions are broken in the transition state for unfolding are covalently linked. Both effects can provide structural information about the otherwise hardly accessible transition state as exemplified for the immunoglobulin proteins C\textsubscript{L} and CD2 where the transition state for folding could be stabilized by disulfide bridges\textsuperscript{20,68} or for barnase, where the transition state for unfolding could be destabilized by a disulfide bond.\textsuperscript{75} Often, disulfide-bridged proteins are found to fold less cooperatively.\textsuperscript{70} This might be caused by the population of disulfide-stabilized intermediates in agreement with the stabilization of the transition state and subsequent partially folded states. If stabilization of partially folded structures becomes too strong, either by native or non-native interactions, this can even result in a net deceleration of a protein folding reaction as has been observed for CD2.\textsuperscript{71} Attempts to increase the folding rate by multiple disulfide bonds had the opposite effect: deceleration of the folding reaction by the over-stabilization of a partially folded state.\textsuperscript{71} This highlights the role of cooperativity for efficient protein folding which can be beneficially or detrimentally influenced by preformed tertiary interactions like disulfide bridges.
2.1.4 Conclusions

Disulfide bonds are one of the most widespread structural elements stabilizing the native state of a protein. Their stabilizing role likely arises from a variety of effects imposed on the unfolded as well as the native state. A destabilization of the unfolded state due to the restriction of the conformational freedom as well as decreased protein-solvent interaction will favor folding to the native state which, additionally, can be stabilized by induced proximity of energetically favorable interactions. In most cases, in particular for naturally selected disulfide-bond positions, the stabilizing effects of this covalent bond are more pronounced than eventually occurring destabilizing effects. These can include a stabilization of the unfolded state due to residual or non-native structure. Additionally, strain as well as reduced dynamics imposed on the native state can significantly decrease the stability of the native state. The sum of all these factors will be the net stabilizing effect of a disulfide bridge.

Having a closer look at the kinetic impact of disulfide bridges is highly insightful. By accelerated folding or decelerated unfolding, entropic, enthalpic and structural conclusions about otherwise almost inaccessible transition states can be drawn. Furthermore, how disulfide bonds are positioned in natural proteins might not only help in the design of proteins with improved characteristics but also hold a clue for the specific structural features a protein has been selected for. In this respect, it is particularly revealing to look at disulfide bridge positions in an evolutionary perspective. For many extracellular proteins which do not have to undergo a large variety of binding reactions requiring large scale dynamics, loop structures and flexible parts are often found to be disulfide-linked.\(^7\) This, on the one hand, directly reduces susceptibility to protease digestion and, on the other hand, might generally reduce unfolding rates by linking parts which are likely to come apart early in an unfolding reaction. If large flexibility is needed, as e.g. in members of the immunoglobulin superfamily, which are optimized for molecular recognition processes, another evolutionary strategy has prevailed. Here, disulfide bridges are found in the folding nucleus,\(^7\) where they seem to accelerate refolding if unfolding occurs. At the same time this strategy permits flexibility where needed and also results in a net stabilization of the native state. Positions of evolutionary selected disulfide bonds might hence not only provide insight into folding nuclei but additionally provide a basis for the rational stabilization of homologous proteins.

The elucidation of the versatile effects disulfide bonds have on protein structure, stability and folding has significantly extended our knowledge about how proteins fold and function and it paved the path to influence their properties in desirable ways.

References


Hiermit erkläre ich, dass die vorliegende Arbeit selbstständig verfasst wurde und keine anderen als die hier angegebenen Quellen und Hilfsmittel verwendet wurden. Die Arbeit wurde noch keiner Prüfungskommission vorgelegt.

Matthias J. Feige
München, September 2009


Geld bringt keine Freiheit, kann sie aber nehmen, deshalb danke ich der Studienstiftung des deutschen Volkes nicht nur für die finanzielle Freiheit während meines Studiums und der Promotion sondern auch die geistige auf all den gemeinsamen Veranstaltungen.