Mechanistic Plasticity and Molecular Recognition: The Structural Biology of the MAP Kinase Interacting Kinases 1 and 2, the NAD Synthetase and the Zinc Finger Associated Domain

PhD Thesis

In partial fulfillment of the requirements for the degree of

Doctor of Philosophy (Ph.D.) / Doctor rerum naturalium (Dr. rer. nat.)

in the Molecular Biology Program

at the Georg-August University Göttingen,

Faculty of Biology

Submitted by

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Born on the 28.06.1977 in Weida (Thüringen)

Göttingen, 2005
Affidavit

Here I declare that my Ph.D. thesis entitled “Mechanistic Plasticity and Molecular Recognition: The Structural Biology of the MAP Kinase Interacting Kinases 1 and 2, the NAD Synthetase and the Zinc Finger Associated Domain” has been written independently with no other sources and aids than quoted.

Ralf Jauch

Göttingen, 27.09.2005
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Abstract

In this study a set of four different proteins has been examined structurally and functionally. The core results embody the structural analysis of the catalytic domains of the mitogen activated protein kinase (MAPK) interacting kinases (Mnk) 1 and 2. These proteins are Ser/Thr protein kinases which target the translation initiation machinery in response to activation by MAPK cascades. Despite a highly similar primary sequence the crystal structures of Mnk1 and Mnk2 exhibit pronounced differences in structural modules which are required for enzymatic activity. Both proteins were found to adopt a conformation that is incompatible with productive binding to either ATP or the peptide substrate. The structural snapshots obtained from Mnk1, Mnk2 and Mnk2\textsuperscript{D228G}, a Mnk2 active site mutant, in combination with activity assays enable to derive a model for the functional dynamics common to Mnk subfamily members. The model predicts that activation of these protein kinases is guided by pronounced rearrangements of the activation segment. The results indicate that the regulation of Mnks involves autoinhibition mediated by the structurally reorganized activation segment and suggests strategies for inhibitor development.

A second example highlighting the structural rearrangements which accompany the activity cycle of an enzyme is the \textit{Escherischia coli} nicotinamide adenosine dinucleotide synthetase (\textit{ecoNADS}). An ensemble of five crystal structures of this enzyme was analyzed. They show that the structural rearrangements of \textit{ecoNADS} are dependent on the presence of its substrates and products. The results allow defining a catalytic state and a substrate/product exchange state of the enzyme.

Finally, the insect-specific zinc-finger associated domain (ZAD) was investigated. The ZAD constitutes a highly expanded sequence motif in \textit{Drosophila}. Biochemical and structural data revealed that the ZAD of the \textit{Drosophila} transcription factor Grauzone adopts a novel zinc coordinating fold. This fold bears a fingerprint of hydrophobic residues that facilitate homodimerisation in an antiparallel fashion. The results show that the ZAD constitutes a protein-protein recognition module which participates in the transcriptional regulation in specific phylogenetic lineages.
Introduction

Scope and Motivation

Richard Feynman, who received the Nobel Prize in physics in 1965, once pointed out that: “...everything that living things do can be understood in terms of jiggings and wigglings of atoms” (Feynman et al., 1963). Although this reductionist view may not be common sense, there is no doubt that structural biology is concerned with one of the most fundamental aspect of biology – how the physical shape of biological macromolecules determines their function. The present work aimed at the elucidation of dynamical features of biological catalysts and addresses the questions how those features guide the enzymatic reaction and regulate the activity. Furthermore, it was attempted to study the molecular recognition on the protein-protein as well as on the protein-ligand level. To this end, the structure and function of three nucleotide utilizing enzymes, two closely related protein kinases MAP kinase interacting kinases (Mnk) 1 and 2 (Chapters I and II) and the *Escherichia coli* nicotinamide adenosine dinucleotide synthetase (ecoNADS) (Supplementary Chapter III) were investigated. In addition, the zinc-finger associated domain (ZAD) of the *Drosophila* transcription factor Grauzone was studied (Supplementary Chapter IV).

The studies on Mnk1 and Mnk2 were initiated to define the structural basis for Mnk regulation and to propose novel strategies for protein kinase inhibitor development. The investigation of various structures of the ecoNADS was considered to provide an instructive complementation of studies on the mechanistic dynamics of nucleotide utilizing enzymes. In addition, the bacterial NADS has recently been highlighted as promising target to develop anti-anthrax drugs (Sutherland, 2003). The phylogenetic comparison to the *Bacillus subtilis* NADS (Rizzi et al., 1996;Rizzi et al., 1998), which is highly similar to the enzyme of the pathogen *Bacillus anthracis*, was therefore expected to reveal intricate details within the substrate recognition site which are relevant for designing antimicrobial agents.

The investigation of the ZAD extended the *in silico* characterization of this insect specific protein module (Chung et al., 2002). The ZAD constitutes and abundant sequence motif that is expanded in insect genomes and shares no sequence homology to previously
determined structures. By characterizing ZAD it was attempted to clarify the functional roles of this protein module and to provide a structural view on a potential insecticide target.

By the time of completion of this PhD thesis, three manuscripts presented herein have been published (Supplementary Chapters III and IV) or are in press (Chapter I; scheduled for publication October 2005) in peer-review journals. The manuscript presented in Chapter II will be submitted for publication in due course. Copies of the manuscripts can be found in the corresponding chapters. The potential application of the investigated structures for drug design efforts has been protected by patent application in the case of the ZAD (European Patent 03017076) and Mnk1/2 (European Patent 050198899). The study on the Mnk subfamily as a prototype variation of eukaryotic protein kinases (PKs) has been the main focus of this work and has therefore received the largest overall attention. Hence, the Chapters I and II containing the studies on Mnk1 and Mnk2 constitute the core of this PhD thesis, whereas the work on NADS and ZAD is considered as a supplementation that frames the core project. To highlight this weightening, the chapters discussing NADS and ZAD are denoted as Supplementary Chapters III and IV. Priorities given within the introduction and the discussion were adjusted accordingly.

**Tuning Cells with Phosphoryl Switches**

Reversible phosphorylation acts as a molecular switch which regulates a multitude of biological activities (Krebs, 1983; Pawson and Scott, 2005). Pioneering work that demonstrated this profound biological insight has been carried out by Edwin G. Krebs, Edmond H. Fischer and Earl Sutherland in the 1950s by studying the rabbit muscle phosphorylase (Fischer and Krebs, 1955; Sutherland and Wosilait, 1955; Krebs and Fischer, 1956). Subsequent studies lead to the discovery of PKs as the molecular players behind those events (Krebs et al., 1958). PKs constitute a class of enzymes which catalyze the transfer of the terminal phosphat group of ATP or related nucleotides onto the hydroxyl group of the ami-
no acid residues serine, threonine or tyrosine within substrate proteins. This covalent modification may lead to an alteration of the activity, stability or subcellular localization of the affected proteins (Krebs, 1983; Pawson and Scott, 2005).

Protein kinases are often activated by extracellular stimuli. Thus, they enable cells to adapt their physiology in response to external signals, e.g. during cell-cell communication. Genome sequencing projects underlined the significance of the PKs by identifying more than 500 members of this gene family encoded by the human genome (Manning et al., 2002). This large number groups PKs among the most abundant gene families only topped in number by genes encoding the C2H2 Zinc-finger and the immunoglobulins (Lander et al., 2001).

**The Making of a Protein Kinase Follows a Conserved Construction Plan**

The investigation of an increasing number of PKs led to the recognition of common features with a putative role in the catalytic reaction. The functional domain, which spans 250 to 300 amino acids, has been defined and was further divided into 11 conserved subdomains (I to XI form N-to C-terminus; Figure 1; Hanks et al., 1988).
Certain amino acid motifs have been found in the vast majority of protein kinases such as the Gly-X-Gly-X-Gly (X corresponds to any amino acid) motif of subdomain I, the Lys of subdomain II, an Asp-X-X-X-Asn motif of subdomain VI and an Asp-Phe-Gly (DFG) motif of subdomain VII (Hanks et al., 1988). The region between the DFG motif and the Asp-Pro-Glu (APE) motif, the latter denotes a conserved tripeptide of subdomain VIII, often bears residues which are targeted for activating phosphorylation by upstream kinases and has thus been termed the ‘activation segment’ (Johnson et al., 1996; Nolen et al., 2004).

The first crystal structure of a PK had been published in 1991 and revolutionized the understanding of PK function (Knighton et al., 1991a). The cAMP dependent PK (PKA) in complex with Mg^{2+}-ATP and an inhibitory peptide unraveled the overall fold as well as the modes of nucleotide and peptide recognition of a prototype PK (Knighton et al., 1991a; Knighton et al., 1991b; Figure 2). Subsequent studies demonstrated that the overall architectural makeup of PKs is conserved (Taylor and Radzio-Andzelm, 1994; Johnson et al., 1996; Huse and Kuriyan, 2002). In short, PK domains (PKDs) fold into two structural modules which are termed the ‘N-terminal lobe’ (subdomain I to V) and the ‘C-terminal lobe’

Figure 2: The topology of a prototype protein kinase and the nomenclature of the activation segment. (A) The cAMP dependent protein kinase (PKA) in complex with an ATP molecule (ball-and-stick-representation) and a peptide inhibitor (PDB-id: 1atp). The peptide inhibitor mimics a substrate peptide (red). The N-terminal lobe is depicted in black and the C-terminal lobe in gray. The secondary structure is labeled according to standard conventions and used henceforth in the text accordingly. (B) Enlarged depiction of the activation segment and αF of PKA in the same orientation as in (A). The activation segment in (B) represents a prototype activation segment in its active configuration. Structural elements of the activation segment discussed in the text are indicated.
(subdomain V to XI), respectively. This makeup leads to a characteristic bilobal architecture of the PKD. The ATP molecule is sandwiched in a deep groove between both lobes termed the interlobal cleft (Figure 2A). The N-terminal lobe is composed of a twisted β-sheet comprising five antiparallel strands (β1-β5) and the helix αC. Certain PKs such as PKA contain more α-helices within the N-terminal lobe of which only αC is conserved throughout the PK superfamily. The linker between the two lobes is referred to as the ‘hinge’ region. The hinge region participates in the interaction with the ATP molecule through main-chain H-bonds with the N1 and 6-amino group of the adenine. The C-terminal lobe consists largely of a α-helical bundle (αD-αH) and bears the structural elements required for peptide binding and phosphoryl transfer. One of these elements is the activation segment (Figure 2B). The activation segment is a highly dynamic entity and its conformation is often associated with the catalytic activity. The activation segment can be further subdivided into structurally and functionally distinct components. It harbors (in N- to C-terminal direction) the magnesium binding DFG/D motif, the activation loop which is often phosphorylated by upstream PKs, the P+1 loop and the short helix αEF (Nolen et al., 2004). The αEF/αF loop constitutes the C-terminal ‘anchor’ of the activation segment and connects it to helix αF (Figure 2B). The structural elements downstream of αF lack a functionally copious annotation. However, recent reports suggest that helices αG and αH at the C-terminal end of the PKD provide an important scaffold required for the interaction with substrate and/or regulatory proteins (Kannan and Neuwald, 2005; Kim et al., 2005).
Figure 3: Active site residues and reaction mechanism of a protein kinase.
An ATP molecule and the side chains of amino acids with immediate relevance for the phosphoryl transfer are shown. The reaction mechanism depicted here assumes a general base catalysis (as indicated by the arrows). However, the actual biochemical events are still under debate (Adams, 2001; Hart et al., 1998). Only the non-hydrogen and non-carbon atoms are shown. The subdomains from which the amino acid residues originate are shown in brackets.

Protein Kinase Function

PKs facilitate the phosphoryl transfer through accurate positioning of the reactants by active site elements. These elements often constitute highly conserved entities (Taylor and Radzio-Andzelm, 1994; Figure 3). The phosphate groups of ATP are held in place by the Gly-rich loop (Gly-X-Gly-X-Gly motif; subdomain I) and the invariant Lys from subdomain II. This Lys is in turn positioned by an invariant Glu which emanates from helix αC and forms a Lys-Glu ion pair found in all active PKs. The terminal phosphate group of ATP is polarized and thereby activated, by a catalytic magnesium ion which is coordinated by the first Asp of the DFG/D motif (Adams, 2001). As a consequence, the γ-phosphate group becomes positioned for the nucleophilic attack by the hydroxyl group of the substrate peptide. The latter reaction is further potentiated by the catalytic Asp emanating from the catalytic loop (C-loop; corresponding to the conserved Asp-X-X-X-Asn motif of subdomain VI). The catalytic Asp has been proposed to either act as a base catalyst by accepting a proton from the substrate or as...
stabilizer of a protonated transition state of the γ-phosphate group (Adams, 2001; Hart et al., 1998; Figure 3).

In order to enable the phosphoryl transfer, the substrate peptide has to be docked to the active site with the phosphate receiving group in close proximity to the terminal phosphate of ATP (Hubbard, 1997; Brown et al., 1999; Yang et al., 2002). This positioning is achieved through contributions of residues from the C-loop as well as from the P+1 loop located within the activation segment. The P+1 loop received its designation because it binds the residue of the peptide substrate which is located downstream of the phosphorylation site (P0-site; residues upstream are designated P-1, P-2 and so forth; residues downstream P+1, P+2 etc) (Knighton et al., 1991a). Therefore, the P+1 loop plays an important role in conferring substrate specificity (Hanks, 2003). The discrimination between protein Ser/Thr and Tyr kinases, for example, is essentially achieved through a single amino acid residue from the P+1 loop. In the case of a Ser/Thr kinase, the P+1 loop bears either a Ser or a Thr which forms hydrogen bonds with the catalytic Asp (Knighton et al., 1991a; Lowe et al., 1997; Yang et al., 2002). In a Tyr kinase, however, the Ser/Thr is substituted by a Pro which engages in a stacking interaction with the phenol of the substrate tyrosine and thereby positions this amino acid residue for catalysis (Hubbard, 1997).

The Regulation of Protein Kinases is Diverse

Despite the conserved architecture of their catalytic domains, individual PKs often function in distinct physiological contexts and are regulated by a number of different mechanisms. PK activity is adjusted, for example, by signaling molecules such as cyclic nucleotides (Kim et al., 2005) and Ca\(^{2+}\)/Calmodulin (Goldberg et al., 1996), by covalent modifications such as phosphorylation (Hubbard, 1997) or by interactions with regulatory subunits and regulatory domains (Jeffrey et al., 1995; Kobe et al., 1996). In many cases, PK regulation involves a combination of these mechanisms. The biochemical events accompanying PK activation induce intricate conformational changes within the catalytic domain which leads to a positioning of active site elements. This way, recognition sites for ATP or substrate peptides are made accessible. However, there is no stereotypical mode of kinase regulation and the structural changes accompanying PK activation are highly diverse.
(see, for example, p38 (Bellon et al., 1999); cyclin dependent kinase 2 (CDK2; Jeffrey et al., 1995), Twitchin (Kobe et al., 1996); CAMKI (Goldberg et al., 1996); insulin receptor kinase (IRK; Hubbard, 1997); c-Abl (Nagar et al., 2003); FLT3 (Griffith et al., 2004)). The understanding of the regulation of individual PKs does therefore require a delineation of mechanistic events at the atomic level.

The impact of structure-function studies on PKs is underlined by the fact that deregulation of PK activities is a major cause of human diseases including cancer, inflammation and arthritis (Cohen, 2002; Vieth et al., 2005). Hence, the investigation of the structural basis for PK regulation assists in diagnosing, understanding and readjusting PK malfunction, for example, by rationalizing efforts to design specific PK inhibitors (Gill et al., 2005). The present work elucidates regulatory principles of the Mnk subfamily of PKs and proposes how these principles can be applied for therapeutic interventions.

**Figure 4: Organization of the human Mnk proteins.**
Schematic depiction of the two human Mnk isoforms indicating the arrangement of functional domains (as labeled). The region investigated herein (Mnk1/2 kinase region, Mnk1/2-KR) is boxed with dashed lines. Alternative splicing does neither affect the N-terminus nor the kinase domain. NLS - nuclear localization signal. eIF4G – eukaryotic initiation factor 4G, the scaffolding protein of the translation initiation complex which binds Mnk1 and Mnk2 (Pyronnet et al., 1999; Scheper et al., 2001).
The Subfamily of MAPK Interacting Kinases

The analysis of the phylogenetic tree of PKs led to the definition of nine major groups termed after prominent group members (Manning et al., 2002). These groups were further subdivided into families and subfamilies (www.kinase.com). The Ca\(^{2+}\)/Calmodulin modulated protein kinase (CAMK) group comprises 80 members including the name-giving CAMK family of PKs, the phosphorylase kinase family and MAP kinase associated protein kinase (MAPKAP) family (Hanks, 2003).

The Mnks constitute a subfamily belonging to the MAPKAPs encoded by two genes within the human genome (\textit{mnk1} and \textit{mnk2}). Both Mnk isoforms are alternatively spliced leading to a total of four proteins products. The resulting proteins are termed Mnk1a, Mnk1b, Mnk2a and Mnk2b and range from 347 to 465 amino acid residues (Slentz-Kesler et al., 2000; Scheper et al., 2003; O'Loghlen et al., 2004; Figure 4). The differential splicing affects a region C-terminal of the central kinase domain which bears both a Crm-1 type nuclear export signal (Fornerod et al., 1997) and a MAPK binding motif. This way, the subcellular localization and the basal activity of the corresponding proteins is altered (O'Loghlen et al., 2004; Parra-Palau et al., 2003). In contrast, the region N-terminal of the PK domain is common to all four splice-products (Scheper and Proud, 2002). This region bears a basic sequence which functions as a nuclear localization signal and a binding region for the eukaryotic initiation factor 4G (eIF4G; Scheper and Proud, 2002). The catalytic PKDs of Mnk1 and Mnk2 locates to the center of the enzymes and spans their largest portion. The PKD of both isoforms exhibits an amino acid identity of 78% suggesting similar substrate specificities and regulatory principles.

Mnks are downstream components of intracellular signal transduction pathways (Duncan et al., 2005). They are activated in response to phosphorylation by the MAP kinases ERK1/2 and p38, which are activated upon cellular stress, cytokine or mitogen induced signaling cascades (Fukunaga and Hunter, 1997; Waskiewicz et al., 1997; Scheper et al., 2001). The hitherto most intensively studied Mnk substrate is the eukaryotic initiation factor 4E (eIF4E) (Scheper and Proud, 2002). Mnks have been shown to phosphorylate eIF4E at a conserved serine close to its substrate binding region \textit{in vitro} and \textit{in vivo} (Waskiewicz et al., 1997; Waskiewicz et al., 1999; Ueda et al., 2004).
The Mnk substrate eIF4E is a component of the trimeric eIF4F complex which is required to initiate the translation of most eukaryotic mRNAs. The eIF4F complex contains, in addition, the DEAD-box RNA helicase eIF4A and the scaffold protein eIF4G. eIF4E is required to link this complex to the mRNA’s 5’-terminus by physically binding to the m\(^7\)GpppN cap structure (m\(^7\)G represents 7-methylguanylate; p represents phosphate groups and N represents any base; Marcotrigiano et al., 1997; Preiss and Hentze, 2003). A cap-bound eIF4F complex is essential for the recruitment of the 43S pre-initiation complex including the small ribosomal subunit and the initiator tRNA to the mRNA (Sachs, 2000; Preiss and Hentze, 2003; Proud, 2005). Since the initiation of translation is tightly controlled, e.g. by posttranslational modifications of initiation factors, Mnks could potentially function as regulators of gene expression on the level of mRNA translation (Gebauer and Hentze, 2004).

It has indeed been demonstrated that phosphorylation of eIF4E reduces the binding affinity to capped mRNA \textit{in vitro} (Scheper et al., 2002; Zuberek et al., 2003). However, the physiological effects of eIF4E phosphorylation on mRNA translation are under debate, but it seems that the overall protein synthesis is not affected by eIF4E phosphorylation (McKendrick et al., 2001; Morley and Naegele, 2002; Ueda et al., 2004). Instead, several reports suggest that effects of Mnk activity is restricted to regulate the translation of specific mRNAs (Nikolcheva et al., 2002; Walsh and Mohr, 2004; Buxade et al., 2005). It has been suggested that transcripts with a complex 5’-untranslated region (UTR) structure are susceptible for Mnk-mediated regulatory mechanisms (Sonenberg and Gingras, 1998).

A recent study indicates an impact of Mnks on mRNA translation which is independent of eIF4E (Buxade et al., 2005). In this study, it was shown that Mnk1 phosphorylates the heterogenous nuclear ribonucleoprotein A1 (hnRNP A1). hnRNP A1 binds AU rich sequences within the 3’-UTR structures of many eukaryotic mRNAs and thereby affects various aspects of mRNA regulation including translation, splicing and subcellular localization (Allemand et al., 2005). As a result of Mnk1 mediated phosphorylation of hnRNP A1, the translation of the tumor necrosis factor alpha (TNF-\(\alpha\)) was downregulated (Buxade et al., 2005). Hence, Mnks target not only eIF4E but also other components of the translation initiation machinery. This way, it appears to regulate the translation of a distinct set of transcripts rather than translation in general.

The recent observation that mice homozygous for a targeted disruption of either one or both Mnk genes show no signs of abnormal development is in line with the conclusion that
Mnks are not essential for overall protein synthesis (Ueda et al., 2004). However, *Drosophila* flies, which have lost the single functional Mnk1/2 homolog, termed Lk6 develop a diet-dependent growth phenotype (Reiling et al., 2005): The *lk6* gene is dispensable when the flies were kept on a high protein diet, whereas a significant and eIF4E-mediated growth reduction has been observed with *Lk6* mutant flies raised on food low in amino acids (Arquier et al., 2005; Reiling et al., 2005). Taken together, the observations of a diet-dependence of Lk6 activity in the fly (Reiling et al., 2005), the indication of Mnk substrates in addition to eIF4E (Hefner et al., 2000; Buxade et al., 2005) as well as the specific role of Mnks in viral translation (Walsh and Mohr, 2004) and in the translation of TNF-α (Buxade et al., 2005) suggest intricate not yet unraveled mechanisms of Mnk action with relevance for human health. Here I will present the crystal structures and biochemical data of the catalytic domains of the two human subfamily members, Mnk1 and Mnk2, and of a Mnk2 active site mutant. The results show that Mnks represent a prototype variation of PKs. The structures provide evidence that Mnks evolved a novel autoinhibitory mechanism. This mechanism is guided by a structurally rearranged activation segment (*Chapter I, II*).
Chapter I: Crystal Structures of the Mnk2 Kinase Domain Reveal an Inhibitory Conformation and a Zinc-Binding Site

Citation:


Statement of Contribution:

crystallization, data collection, structure solution, structure refinement, model building, structure analysis, examination of and comparison to available protein kinase structures, analytical gel filtration, light scattering, ATP binding assay, manuscript preparation
Crystal Structures of the Mnk2 Kinase Domain Reveal an Inhibitory Conformation and a Zinc-Binding Site

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Human mitogen-activated protein kinases (MAPK)-interacting kinases 1 and 2 (Mnk1 and Mnk2) target the translational machinery by phosphorylation of the eukaryotic initiation factor 4E (eIF4E). Here we present the 2.1 Å crystal structure of a non-phosphorylated Mnk2 fragment that encompasses the kinase domain. The results show unique Mnk-specific features such as a zinc-binding motif and an atypical open conformation of the activation segment. In addition, the ATP binding pocket contains an Asp-Phe-Asp (DFD) in place of the canonical magnesium-binding Asp-Phe-Gly (DFG) motif. The phenylalanine of this motif sticks into the ATP binding pocket and blocks ATP binding as observed with inhibitor-bound and thus inactive p38 kinase. Replacement of the DFD by the canonical DFG motif affects the conformation of Mnk2, but not ATP binding and kinase activity. The results suggest that the ATP binding pocket and the activation segment of Mnk2 require conformational switches to provide kinase activity.
Introduction

Control of cellular growth is tightly linked to the regulation of protein synthesis through the eukaryotic initiation factor 4E (eIF4E). eIF4E facilitates the recruitment of mRNA to the translational machinery by binding of the mRNA 5’-cap structure (von der Haar et al., 2004) and is subject to regulation. Nutrition sensing and growth factors regulate formation of productive translation initiation complexes (Richter and Sonenberg, 2005), a process in which eIF4E activity is modulated through phosphorylation by mitogen-activated protein kinases (MAPK)-interacting kinases 1 and 2 (Mnk1 and Mnk2) (Scheper et al., 2001; Waskiewicz et al., 1997; Waskiewicz et al., 1999). Both eIF4E and Mnks bind to eIF4G within the initiation complexes. Mnks are activated in turn by MAP kinases such as extracellular signal regulated protein kinases (ERK) and p38 in response to cytokines, mitogens or cellular stress (Fukunaga and Hunter, 1997; Scheper et al., 2001; Waskiewicz et al., 1997).

The regulatory effects of Mnk-mediated phosphorylation of eIF4E on translation are still controversial (Knauf et al., 2001; Nikolcheva et al., 2002; Ueda et al., 2004) as are the biological consequences of the lack of Mnk activity in mice and fly. Mice homozygous for a targeted disruption of either one or both Mnk genes show no signs of abnormal development (Ueda et al., 2004), whereas loss of the single Drosophila Mnk homolog, Lk6, produces a diet-dependent and eIF4E-mediated growth phenotype (Arquier et al., 2005; Reiling et al., 2005).

Humans possess two mnk genes, both of which encode two alternatively spliced transcripts, resulting in the proteins Mnk1a, Mnk1b, Mnk2a and Mnk2b (Figure 1A; (O’Loghlen et al., 2004; Scheper et al., 2003)). Sequence alignments reveal a central kinase domain in Mnks, which can be placed into the group of Ca²⁺/calmodulin modulated protein kinases (CaMK) (Manning et al., 2002) (Figure 1B) and which is unaffected by alternative splicing (Figure 1A). We have investigated human Mnk2 as a representative of the Mnks, both structurally and functionally. We present the crystal structure of a Mnk2 fragment spanning its kinase domain at 2.1 Å resolution, showing a Mnk-specific zinc-binding module and an open conformation of the activation segment (Nolen et al., 2004). In addition, Mnks contain an Asp-Phe-Asp (DFD) motif in place of the otherwise strictly conserved magnesium-binding Asp-Phe-Gly (DFG), which exhibits a conformation that obstructs ATP binding. Thus, Mnks are architecturally distinct from other protein kinases. We also present structural and functional
data on a Mnk2 mutant containing a canonical DFG motif. Our analysis revealed novel features relevant for the regulation of activity and inhibition of Mnks and demonstrate that Mnks represent a prototype variation of CaMK group kinases.

Results and Discussion

Organization and Atypical Elements of Mnk2

The central region of Mnks harbors the kinase domain of the proteins (residues 81-368 in human Mnk2; Figure 1A, B), according to which Mnks belong to the CaMK group. The N-terminal 80 residues contain a nuclear localization signal (NLS) and a binding motif for eIF4G (Figure 1A). Residues beyond position 368 contribute to the interaction with MAP kinases (Slentz-Kesler et al., 2000) (Figure 1A). Like several other CaMK group members, Mnk’s are not regulated by Ca$^{2+}$/calmodulin in agreement with the lack of a sequence element for the interaction with that protein (Slentz-Kesler et al., 2000).

Protein kinases contain conserved structural elements to bind and activate their protein substrates and the co-substrate ATP (Johnson et al., 1996; Nolen et al., 2004; Taylor and Radzio-Andzelm, 1994). In addition to those, Mnks contain atypical elements such as an Asp226-Phe227-Asp228 (DFD) motif (sequence numbering is according to human Mnk2) in place of the canonical magnesium-binding Asp-Phe-Gly (DFG) motif, and three insertions (I1-I3; Figure 1B), one of which contains a cluster of four conserved cysteines (I3). We asked whether and how these features affect the structure and function of the kinase domain. While we were able to express and purify the recombinant full-length Mnk2, the protein failed to crystallize. We therefore screened for alternative expression constructs, identifying a truncated version (residue 72-385) that encompasses the kinase domain (Figure 1A) and could be produced in soluble form (see Experimental Procedures). This construct is referred to as Mnk2-kinase region (Mnk2-KR) herein. Significantly, non-phosphorylated Mnk2-KR is inactive but can be activated by phosphorylation via ERK2 (Figure 2), thus recapitulating the behavior of the full-length protein (Waskiewicz et al., 1997). The non-phosphorylated Mnk2-KR could be crystallized and yielded diffraction data to 2.1 Å resolution (Table 1).
Figure 1

**A**

NLS/ eIF4G Binding

Kinase Domain

MAPK Binding

Mnk2a

Mnk2b

Kinase Region

(Investigated)

Mnk2b-Specific C-Terminus

**B**

<table>
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<th>Protein</th>
<th>Sequence</th>
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**Figure 1**
Figure 1. Mnk2 Organization and Sequence Alignment
(A) Schematic comparison of the two splice variants of human Mnk2 indicating the arrangement of functional domains (as labeled). The region investigated herein (Mnk2 kinase region, Mnk2-KR) is boxed. Alternative splicing does neither affect the N-terminus nor the kinase domain. NLS - nuclear localization signal. eIF4G – eukaryotic initiation factor 4G, the scaffolding protein of the translation initiation complex which binds Mnk1 and Mnk2 (Pyronnet et al., 1999; Scheper et al., 2001).
(B) Sequence alignment of the kinase domains of human Mnk1 and Mnk2, the Drosophila and C. elegans Mnk orthologs (Lk6 and R166.5, respectively) and three human CaMK group kinases of known structure (MAPKAP - MAP kinase-activated protein-kinase). Mnk2 numbering refers to a recently reported sequence (Slentz-Kesler et al., 2000). Secondary structure elements as found in Mnk2-KR are indicated below the alignment. Stars indicate phosphorylation sites (Scheper et al., 2001). The catalytic loop (i), the DFD motif (DFG in other kinases; ii) and the P+1 loop (iii) are marked with colored bars. Insertions characteristic for Mnks are boxed (I1-I3). Open circles mark Gly91 and Gly93 of the glycine rich loop, Lys113 and Glu129 known to be important for ATP binding (Taylor and Radzio-Andzelm, 1994), filled circles mark Gly164 and Gly165 of the hinge region separating the N-terminal and C-terminal lobes.

Structure Solution and Architecture of the Mnk2 Kinase Region

The structure of Mnk2-KR was solved by molecular replacement using as a search model the structure of the death-associated protein kinase 1 (DAPK1; PDB ID 1JKS; (Tereshko et al., 2001)), another CaMK group member that exhibits about 30% amino acid identity with Mnk2-KR. The final Mnk2-KR model and the DAPK1 structure compare with an average rmsd of 1.32 Å over 211 matching Cα atoms. The Mnk2-KR crystal structure was refined to a final R/Rfree value of 21.5/25.4 % maintaining good overall geometry (Table 1). 16 residues at the C-terminus of Mnk2-KR, which do not pertain to the core kinase domain (Figure 1A), as well as residues 232-250 (part of the activation segment) and 306-309 (part of insertion I3), which are situated in flexible regions, did not show up in the final 2Fo-Fc electron density and have been omitted from the model.

Mnk2-KR displays the typical bilobal arrangement of a protein kinase with the ATP binding cleft sandwiched between an N-terminal and a C-terminal lobe (Figure 3A). The N-terminal lobe contains a twisted β-sheet (β2- β5; numbering of secondary structure elements is according to the standard nomenclature of kinase topology; (Knighton et al., 1991a)) and a α-helix (αC). While the N-terminal part of the protein runs in antiparallel fashion along the edge of strand β2, it does not engage in continuous β-type backbone pairing to the β-sheet and the present structure of Mnk2-KR therefore formally lacks strand β1 (Figures 1B, 3A). The conserved residue Glu129 of helix αC participates in an ionic interaction with Lys113 in β3, as required for stabilizing the ATP substrate (Taylor and Radzio-Andzelm, 1994). The glycine rich loop (residues 90-94) is flexible and contains the conserved residues Gly91 and Gly93 playing key roles in ATP binding (Taylor and Radzio-Andzelm, 1994). The C-terminal lobe
Figure 2. Catalytic Activity of Mnk2-KR and Mnk2-KR\textsuperscript{D228G}
Kinase activity profiles of Mnk2-KR (wild type and D228G mutant) before and after activation by phosphorylation through ERK2. Kinases were titrated in the indicated concentration range into the Mnk kinase assay (see Experimental Procedures). Data points represent means of triplicates, error bars indicate one standard deviation. The data have been subjected to non-linear regression analysis using the four parameter logistic equation (GraphPad Prism software). Results demonstrate that while activation is necessary for both wild type Mnk2-KR and the D228G mutant, the activated wild type and mutant kinase domains exhibit almost identical kinase activity (calculated EC\textsubscript{50}S for Mnk2-KR and Mnk2-KR\textsuperscript{D228G} are 3.8 and 2.8 nM, respectively).
(residues 166-368) starts downstream of the ‘hinge’ connector region (Gly164, Gly165) and contains a predominantly hydrophobic four-helix bundle ($\alpha$D-$\alpha$F, $\alpha$H). Arg204, which defines the RD-kinases that are often activated by phosphorylation (Johnson et al., 1996), the catalytic aspartate (Asp205) and Asn210, which serves in active kinases to coordinates a catalytic magnesium ion (Adams, 2001), are situated in the catalytic loop (C-loop, residues 204-210). In addition, the C-terminal lobe harbors the activation segment (residues 226-260; (Nolen et al., 2004); Figure 3A), which is required for binding the protein substrate.

**Inhibitory Conformation of the Activation Segment**

The magnesium-binding loop (residues surrounding the DFD motif), the activation loop (disordered residues preceding the P+1 loop) and the P+1 loop (residues 250-259) constitute the activation segment (Figure 3A; (Nolen et al., 2004)). It is followed C-terminally by the short helix $\alpha$EF (residues 260-265) and the long helix $\alpha$F (residues 270-290), the latter being part of the four-helix bundle in the C-terminal lobe (Figure 3A). The P+1 loop was shown to interact with a residue adjacent to the phosphorylation site (Knighton et al., 1991b). Together with the adjoining helix $\alpha$EF, the P+1 loop represents subdomain VIII (Hanks and Quinn, 1991), which in active protein kinases is cradled in a predominantly hydrophobic environment provided by $\alpha$F, $\alpha$G and the C-loop. In the present Mnk2-KR crystal structure helix $\alpha$F is N-terminally extended due to a Mnk-specific insertion (I2; Figure 1B). As a consequence, the protein disengages the canonical intramolecular interaction between subdomain VIII and the $\alpha$F/$\alpha$G/C-loop region and instead two symmetry-related, neighboring molecules in the crystal reciprocally insert their activation segments into the C-terminal lobe of the symmetry mate (Figure 3B). This intermolecular interaction observed for Mnk2-KR maintains conserved polar interactions between subdomain VIII and the $\alpha$F/$\alpha$G/C-loop region, which are normally built up intramolecularly (as seen, e.g., in DAPK1; Figure 3C, Supplementary Figure 1), including a hydrogen bond between Ser253 (P+1 loop) and Asp205 (C-loop) as well as a Glu260-Arg356 ion pair (Figure 3C).

Because protein kinases require the interaction of the P+1 loop with the C-loop for substrate binding and since such an interaction is fostered intermolecularly in the present crystal structure, the arrangement of two Mnk2-KR molecules within the crystals suggested a functionally relevant dimer formation. However, multi-angle laser light scattering and
analytical gel filtration chromatography showed that Mnk2-KR is a monomer in solution (data not shown). Therefore, the interaction of the P+1 loop and the C-loop as required for substrate binding is not promoted by intermolecular contacts in solution. An extruded activation segment as observed in the present structure could therefore be one reason, why non-phosphorylated Mnk2 and Mnk2-KR are inactive (Figure 2). Upon phosphorylation the proteins acquire kinase activity (Figure 2), consistent with the notion that after phosphorylation the activation segment of Mnk2 flips inwards so that it can establish the necessary intramolecular contacts with the $\alpha_F/\alpha_G/C$-loop region. Significantly, in the present inactive conformation Arg204 from the C-loop forms a salt bridge with Asp273 in helix $\alpha_F$ (Figure 3C) that immediately follows the Mnk-specific insertion I2 (Figure 1B). Thus, by positioning Asp273, Arg204 stabilizes the extruded conformation of the activation segment in inactive Mnk2-KR. As known from other RD-kinases (Johnson et al., 1996), Arg204 engages in contacts with a phosphate group after activation. The engagement of Arg204 in an alternative interaction with a phosphate after phosphorylation would release Asp273, which in turn could allow for the structural rearrangement of the activation segment required for activity.

**Inhibitory Conformation of the Magnesium-Binding Loop**

Protein kinases contain a conserved DFG motif (residues 226-228; Figure 3A). The first aspartate of this motif is known to coordinate a Mg$^{2+}$-ion, which is needed to activate the $\gamma$-phosphate of ATP through polarization (Taylor and Radzio-Andzelm, 1994). The Mnk2-KR contains a DFD motif (residues 226-228) in place of the canonical DFG motif normally found in protein kinases. This DFD fingerprint is unique within the human kinome but conserved within the Mnk subfamily (Figure 1B). As compared to the DFG motif of, for example, DAPK1, the DFD motif of the Mnk2-KR is rotated by almost $180^\circ$ (Figures 4, 5A). As a result, Phe227 is displaced from its standard position in active kinases and sticks into the ATP binding pocket (Figure 4A), thus excluding ATP from its binding site. The residue also blocks an access route for the nucleotide to the pocket (Figure 4B). Thus, the arrangement of the DFD motif in Mnk2-KR interferes with the generic mode of ATP binding (Figure 4C).

The present conformation of the DFD motif is stabilized (i) by a hydrogen bonding interaction of the carbonyl oxygen of Phe227 with the side chain amino group of Lys113; and
Figure 3
Figure 3. Overall Structure of Mnk2-KR and the Extruded Activation Segment

(A) Stereo ribbon plot displaying the overall Mnk2-KR structure. Prominent structural elements are displayed in colors and labeled according to standard nomenclature (Knighton et al., 1991a). The magnesium-binding loop (around the DFD motif), the activation loop and the P+1 loop constitute the activation segment. Subdomain VIII is in yellow (Hanks and Quinn, 1991). Note that the glycine rich loop (pink) exhibits partially disordered side chains and that residues Gly232 to Cys251 are missing in the final electron density (dotted line). The missing portion encompasses part of the activation loop including Thr244 and Thr249 that are phosphorylated by upstream activating kinases. Another region with partially missing electron density and high temperature factors corresponds to the insertion with the specific cysteine cluster (residues 304-310; tip of the zinc finger). Green sphere: zinc ion.

(B) Side view of two neighboring Mnk2-KR molecules in the crystal lattice (Mol1 and Mol2, light and dark gray, respectively), which show a reciprocal interaction via the P+1 loop and helix αΕF (red for Mol1, blue for Mol2). The left molecule (Mol1) is rotated 20° clockwise about the vertical axis compared to the view in (A).

(C) Stereo ribbon plot showing details of the interaction of the αEF/P+1 loop portion of one molecule (Mol 2, blue) with the C-terminal lobe of a neighboring molecule in the crystal (Mol1, gray with αF in red). Residues involved in conserved polar interaction are highlighted as sticks. Atoms are color-coded by atom type (carbon – as the respective protein backbones; oxygen – red; nitrogen – blue). The view is on the boxed region in (B) rotated 90° about the horizontal axis as indicated.

(ii) by positioning of the Phe 227 side chain in a hydrophobic pocket formed primarily by Leu143 and the non-conserved Phe159 (which corresponds to the so called gatekeeper residue that acts as selectivity filter for ATP-competitive drugs (Cohen et al., 2005); Figure 4A). The Mnk-specific Asp228 does not participate in stabilizing this conformation by direct interactions.

The non-canonical arrangement of the DFD motif in Mnk2-KR is referred to as the ‘DFG/D-out’ conformation, as opposed to the standard ‘DFG/D-in’ conformation as found e.g. in DAPK1 (Tereshko et al., 2001). A DFG/D-out conformation in the Mg^{2+}-binding loop was initially found in the apo-form of the insulin receptor kinase (IRK), a tyrosine kinase (TK) (Hubbard et al., 1994). A similar DFG/D-out conformation is observed in co-crystals of tyrosine kinases such as the cellular protein encoded by Abelson gene (c-Abl), cKIT and the vascular endothelial growth factor receptor (VEGF-R) kinase with their inhibitors Gleevec and AAL-993, respectively, (Nagar et al., 2002). It is also found in the Ser/Thr kinase p38 when bound to diaryl urea scaffold molecules (Figure 4D; (Pargellis et al., 2002)). The latter substances, termed ‘allosteric inhibitors’, switch the DFG/D-in conformation found in the inhibitor-free p38 (Wilson et al., 1996) into a DFG/D-out conformation, in which the central phenylalanine blocks the ATP binding by sterical hindrance as observed with Mnk2-KR.

Since the DFG/D-out conformation would prevent productive ATP binding (Figure 4C), a switch of DFG/D-out into DFG/D-in is prerequisite for ATP binding in Mnk2-KR. The proposed structural rearrangement can be modeled without sterical clashes (Figure 4A).
Figure 4
**Figure 4. Neighborhood of the DFD Motif**

(A) Close-up stereo view of the DFD region and the ATP binding cleft. The DFG/D-out conformation of wild type Mnk2-KR is indicated by a stick representation for Asp226, Phe227 and Asp228 on the upper left with Phe227 and Asp228 poking into the ATP binding cleft (carbon – cyan). A DFG/D-in conformation (lower right; carbon – green) has been modeled according to the DFG/D-in conformation seen in other kinases and as observed for the Asp228Gly mutant of Mnk2-KR (see Figure 5B). A backbone trace of Mnk2-KR is shown as a semitransparent gray tube. Residues within a radius of 4Å around the of the DFD motif in either the DFG/D-in or -out conformations are displayed as sticks (carbon – gray). Direct interactions with the protein matrix, which stabilize the DFG/D-out conformation, are indicated by dashed lines. Phe227 comes to lie in two different hydrophobic pockets in the two different conformations. No obstacle for adoption of a DFG/D-in conformation is visible. Relative to Figure 3A, the molecule has been rotated 90° clockwise about the vertical axis.

(B) Stereo view of the molecular surface of Mnk2-KR color-coded by electrostatic potential (blue – positive charge; red – negative charge) with the two conformations of the DFD motif as a stick representation (color-coding as in (A)). The ATP binding cleft is pointed out. Asp228 in either conformation is well accessible to the aqueous solvent. The DFG/D-out conformation not only positions Phe227 and Asp228 in the ATP binding cleft but also obstructs access to this cleft from the front. The molecule has been rotated by 30° about the horizontal axis (N-terminal lobe to back) relative to (A) in order to afford an unobstructed view into the DFD pockets.

(C) Same view as in (A) with a non-hydrolyzable ATP analog (adenosine 5’-[β,γ-imido]-triphosphate (AMPPNP); carbon – beige; phosphorus – violet) superimposed as seen in the co-crystal structure with DAPK1 (PDB ID 1IG1). In the DFG/D-out conformation the adenine base clashes with the side chain of Phe227, the phosphate groups clash with the side chain of Asp228.

(D) The same view as in (A) and (C) with only the DFG/D-out conformation shown. The DFG-region of a p38-BMU inhibitor complex (PDB ID 1KV1) is shown for comparison (magenta tube; DFG in stick representation; carbon – magenta) as seen after global superpositioning of the protein structures. The BMU inhibitor (carbon beige; chloride – green) occupies part of the DFG/D-in binding pocket and induces a DFG/D-out conformation in p38.

### Replacement of DFD by the Canonical DFG Motif

As described, we did not observe any direct interactions of the Mnk-specific Asp228 of the DFD motif with other protein residues, questioning its importance for the DFG/D-out conformation. To clarify this point, we replaced Asp228 by a glycine and solved the corresponding Mnk2-KR<sup>D228G</sup> structure at 3.2 Å resolution. The Fo-Fc ‘omit’ electron density map of the DFG region suggests that the Mnk2-KR<sup>D228G</sup> molecules in the crystals adopt both the DFG/D-in and the DFG/D-out conformation (Figure 5B). Therefore, the replacement of the DFD motif in Mnks by the generic DFG motif destabilizes DFG/D-out in favor of DFG/D-in. The alternative conformation observed in the Mnk2-KR<sup>D228G</sup> mutant also shows that there are indeed no structural constrains that would prevent the switch of the DFD motif into DFG/D-in (Figure 4A).

Together, the above observations suggest that Asp228 moderately stabilizes DFG/D-out although it is not involved in direct intramolecular interactions in the present crystal structure. When modeling a DFD/G-in conformation of a DFD motif based on the structure of the Mnk2-KR<sup>D228G</sup> mutant (Figure 4A) no direct interactions involving Asp228 can be discerned either. The overall exposure of Asp228 to the aqueous environment in either conformation...
Figure 5. Conformation of the DFD Motif and ATP Binding

(A) Stereo plot showing the Fo-Fc ‘omit’ electron density contoured at the 3σ level covering the DFD motif (sticks; carbon - cyan) in wild type Mnk2-KR (gray ribbon). Relative to Figure 3A, the molecule has been rotated 90° clockwise about the vertical axis and 90° about the horizontal axis (N-terminal lobe to front), affording a view from the N-terminal lobe onto the motif. Only the DFG/D-out conformation is visible in wild type Mnk2-KR.

(B) Stereo plot showing the Fo-Fc ‘omit’ electron density contoured at the 2.5σ level covering the DFG motif in Mnk2-KRD228G. The DFG motif is found in both a DFD/G-in (carbon green) and in a DFD/G-out (carbon cyan) conformation in the mutant.

(C) Dot blot assaying the binding of α32-ATP to the indicated proteins at concentrations of 2.8 µM (I), 1.4 µM (II) and 0.7 µM (III). GST – negative control using glutathione S-transferase.
seems to be similar as well (Figure 4B). In addition, we cannot discern decisive differences in the electrostatic potentials around the DFD motif that would favor the DFG/D-out or -in conformation (Figure 4B). One possibility is that interactions involving Asp228 ensue in a slightly modified conformation in solution and that the DFG/D-out conformation has been trapped in the crystals. Gly228 in place of Asp228 in the Mnk2-KR D228G mutant may foster increased backbone flexibility, which could then allow the protein to adopt both conformations.

We next asked whether the DFG/D-out to DFG/D-in switch that is necessary to allow ATP binding requires factors other than ATP itself. ATP binding assays with non-phosphorylated Mnk2-KR, phosphorylated and thereby activated Mnk2-KR and with Mnk2-KR D228G showed that both Mnk2-KR and Mnk2-KR D228G were able to bind ATP and that binding occurred irrespective of the phosphorylation-dependent activation of the proteins (Figure 5C). However, only the phosphorylated forms of Mnk2-KR and Mnk2-KR D228G exerted kinase activity (Figure 2). These results establish that phosphorylation of Mnk2-KR is a prerequisite for catalytic activity but not for ATP binding, and that neither aspect depends on Asp228 of the DFD motif.

Since our solution studies showed that ATP is able to bind to non-phosphorylated Mnk2-KR (Figure 5C), we attempted to co-crystallize the protein with ATP, ATP/Mg²⁺ or the non-hydrolyzeable analog AMPPNP with and without Mg²⁺ or to soak preformed crystals with these substances in order to directly visualize conformational changes taking place upon nucleotide binding. Despite extensive efforts, we were not able to obtain crystals of the ATP-bound form. Indirectly, these negative results suggest that indeed conformational changes take place upon ATP binding.

A Zinc-Binding Motif in Mnk2

Among the Mnk-specific insertions, a region containing four conserved cysteine residues stands out (I3; Figure 1B). In the present crystal structure, the four cysteines are arranged in a manner suggesting a metal ion binding site (Figure 6A). In order to verify the presence of a metal ion in the protein, we recorded the X-ray fluorescence spectrum of frozen Mnk2-KR crystals. Kα and Kβ emission lines characteristic of a zinc ion were observed (Figure 6B). Significantly, Zn²⁺ was never added during the expression, purification or crystallization of the protein and, therefore, must have been scavenged spontaneously by the
protein from the cellular environment. To locate the zinc ion within the Mnk2-KR structure, we reprocessed the original Mnk2-KR diffraction data keeping the Friedel pairs separated. While the diffraction data were collected at a wavelength of 1.05 Å, i.e. remote from the Zn K-edge (about 1.28 Å), the anomalous signal expected at that wavelength (~2.7e) is still strong compared to the maximum signal around the edge (~3.9e). Indeed, the Fourier map calculated from the model phases and the anomalous intensity differences revealed a single peak at ~6σ above the background, which maps to the center of the four cysteine SH-groups (Figure 6A). Therefore we conclude that Mnk2 bears a zinc finger-like module within the kinase domain.

We do not expect a direct effect of the zinc finger on the catalytic activity since it is positioned far away from the catalytic center. In zinc fingers of other proteins, zinc ions contribute to the shaping and compaction of short structural motifs, which in the following often provide a binding platform for proteins and/or nucleic acids (Krishna et al., 2003). Thus, we assume that the Mnk-specific zinc finger serves as a docking site for phosphorylation targets and/or regulators. Studies to test the significance of the Mnk specific zinc finger in vivo are in progress.

Conclusions

Herein, we have investigated a truncated form of human Mnk2 (Mnk2-KR) that covers the kinase domain and exhibits phosphorylation-dependent activity indistinguishable from the full-length protein. Our results provide evidence that Mnks represent a distinct protein kinase subfamily. This subfamily is characterized by three insertions which lead to an open conformation of the activation segment in the non-phosphorylated Mnk2-KR and a zinc-coordinating cysteine cluster. Adoption of the active conformation of the activation segment after phosphorylation may be triggered by a phosphate-mediated disruption of an Arg204-Asp273 salt bridge that otherwise stabilizes the inactive, extruded conformation of the activation segment. In addition, we observed a DFD motif that favors an inhibitory DFG/D-out conformation in contrast to other unliganded Ser/Thr kinases. Interestingly, the DFG/D-out conformation has no qualitative effect either on the ATP binding properties of non-phosphorylated Mnk2-KR or on the catalytic activity after phosphorylation, suggesting that it is marginally stable in the present structure. The latter notion is in agreement with the lack of direct interactions of the Mnk specific Asp228 that would fasten the DFG/D-out conformation. Since both, ATP binding and kinase activity, require an open ATP binding pocket, the switch
Figure 6. Zinc-Binding Site

(A) Stereo ribbon plot of the zinc finger motif in Mnk2-KR (blue). The cluster of four cysteines is indicated by sticks at the tip of the finger (carbon – blue; sulfur - yellow). They surround a zinc ion (green sphere). A Fourier map using the anomalous intensity differences and phases from the refined model is shown contoured at the 5σ level and identifies the position of the anomalous scatterer. The region from Trp305 to Glu309 lacks unambiguous backbone density and has been omitted from the model. The molecule has been rotated 90° counterclockwise about the vertical axis compared to Figure 3A.

(B) X-ray emission spectrum of native Mnk2-KR crystals irradiated with X-rays of 1.05 Å. Peaks in the spectrum correspond to I=Zn-Kα line, II=Zn-Kβ line, III Compton scattering, IV elastic scattering.
into the DFG/D-in conformation might be imposed by ATP binding per se, again consistent with a weak intramolecular stabilization of the DFG/D-out conformation in the Mnk2-KR structure. The question remains whether a regulatory potential via interconversion of DFG/D-in and -OUT conformations could be realized in vivo, e.g. by interaction with other regulatory proteins or by the protein adopting other conformations in solution.

Since the DFG/D-out conformation of Mnk2 is rarely observed in inhibitor-free protein kinases, it provides a handle for specific inhibitor design that would extend the range of generic ATP-competitive compounds (Mol et al., 2004), a speculation that is supported by the binding of the DFG/D-out-inducing p38-inhibitor BIRB-796 to Mnk2 (Fabian et al., 2005). Also, the Mnk-specific Asp228 may aid inhibitor design because it renders the Mg\textsuperscript{2+}-binding loop unique among Ser/Thr kinases.
# Tables

## Table 1. Crystallographic Data

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<sup>a</sup> Values for the highest 0.1 Å resolution shell in parentheses.

<sup>b</sup> $R_{sym}(I) = \left( \sum_{hkl} \frac{\mid I(hkl) - <I(hkl)> \mid}{\sum_{hkl} I(hkl)} \right)$; $I(hkl)$ – intensity of the $i^{th}$ measurement of hkl; $<I(hkl)>$ – average value of hkl for all i measurements.

<sup>c</sup> $R_{work} = \sum_{hkl} \frac{\mid F_{\text{obs}} \mid - k F_{\text{calc}} \mid}{\sum_{hkl} \mid F_{\text{obs}} \mid}$; $R_{free} = \sum_{hkl \in T} \frac{\mid F_{\text{obs}} \mid - k F_{\text{calc}} \mid}{\sum_{hkl \in T} \mid F_{\text{obs}} \mid}$; $hkl \in T$ – test set.
Supplementary Figure 1. Open Conformation of the Activation Segment

(A) View from the N-terminal lobe onto the arrangement of the two molecules (rotated 90° about the horizontal axis compared to the view in Figure 3B). The N-terminal lobe and the upper part of the C-terminal lobe have been cut away to allow an unobstructed view onto the interaction regions. The P+1 loop region of DAPK1 is cradled intramolecularly within the body of the C-terminal lobe and is shown for comparison (brown ribbon) after global superpositioning of the DAPK1 structure (PDB ID 1JKS) onto Mol1.

(B) Stereo ribbon plot comparing the conformations of the P+1 loop/αEF/αF regions in Mnk2-KR (red), MAP kinase-activated protein-kinase-2 (MAPKAP2; yellow; PDB ID 1NXK) and DAPK1 (brown; PDB ID 1JKS). The segments are shown after global superpositioning of the protein structures. The remainder of a Mnk2-KR molecule is indicated by a semi-transparent gray ribbon. The view is the same as in (A).
Experimental Procedures

Cloning

A cDNA fragment of human Mnk2 (amino acid residues 72-385; Entrez AAG26336), which encompasses the kinase domain, was amplified using the forward/reverse primer pair 5’CGGGATCCACCGACAGCTTCTCGGGCAGG/5’ACGCGTCGACACTACCTCTCGCAGGACCATGGGAG (restriction sites underlined) and cloned into the BamHI and SalI sites of the vector pGEX-4T1 (Amersham, Uppsala, Sweden). The amino acid substitution Asp228Gly was introduced into the GST-Mnk2-KR construct employing the Stratagene (Heidelberg, Germany) Quik Change Site Directed Mutagenesis kit according to the manufacturers’ instructions. Mutagenesis oligonucleotides were 5’GAAGATCTGTGACTTCGGCCTGGGCAGCGCATCAAACTC and 5’GAGTTTGATGGCCGCCCATCAAACTC.

Protein Preparation

Expression of GST-Mnk2-KR or GST-Mnk2-KR^D228G was performed in E. coli BL21. Cells were grown in LB medium (Merck, Darmstadt, Germany) supplemented with ampicillin (100 µg/ml; 37°C). Expression was induced at OD_{600} of 0.8 with 1 mM isopropyl thiogalactoside (4 h; 25°C). Cells were harvested by centrifugation and resuspended in 10 ml lysis buffer (50 mM Tris/HCl, pH 7.5, 200 mM NaCl, 5 mM DTT) per gram wet weight. Lysates were prepared by sonication and subsequent clearing in a Sorvall SS34 rotor (18000 rpm, 45 min, 4°C). The supernatant was applied to two GSTPrep FF 16/10 columns (Amersham) connected in series and equilibrated with lysis buffer.

Wild type or mutated Mnk2-KR were eluted by on-column thrombin cleavage from the GST tag. 1000 units of thrombin (Amersham) were dissolved in 60 ml wash buffer and cycled over the columns (over night, 8°C). After collection, the thrombin eluate was diluted (1:5 in 50 mM Tris/HCl, pH 8.0), applied to a 25 ml Q sepharose HP column (Amersham) and eluted with a linear gradient of sodium chloride (50 mM Tris/HCl, pH 8.0, 0-1 M NaCl). Fractions were pooled, concentrated (approx. 16 mg/ml) and transferred into 10 mM Tris/HCl, pH 7.5, 50 mM NaCl, 1 mM DTT via a PD10 column (Amersham). Typically the final protein concentration was approximately 12 mg/ml.


**Crystallographic Analysis**

Crystals of wild type Mnk2-KR and of Mnk2-KR$^{D228G}$ were grown by vapor diffusion with a reservoir solution containing 23% (w/v) polyacrylic acid 5100, 2% (v/v) 2-methyl-2,4-pentane-diol and 0.1 M HEPES/NaOH, pH 7.7 and were frozen (liquid nitrogen) in reservoir solution supplemented with 12.5% glycerol and 12.5% ethyleneglycol. Diffraction data (Table 1) were collected on beamline BW6 (DESY, Hamburg, Germany) at 100K on a MarResearch (Norderstedt, Germany) CCD detector and processed with the HKL package (Otwinowski and Minor, 1997).

The Mnk2-KR structure was solved by molecular replacement (MolRep; (Collaborative Computational Project, 1994) using the structure coordinates of DAPK1 (PDB ID 1JKS) as the search model. Model building was conducted automatically using arp/warp (Cohen et al., 2004) and manually using Xfit (McRee, 1999). Refinement was performed with CNS (Brunger et al., 1998) and Refmac5 (Collaborative Computational Project, 1994) using standard protocols. All data between 15.0 and 2.1 Â resolution were employed in the refinement, setting aside 5% of the reflections to monitor the free R-factor (Table 1). In the last refinement cycles, different overall anisotropic temperature factor corrections were applied to the N-terminal lobe and the C-terminal lobe using the TLS refinement option of Refmac5. The structure of Mnk2-KR$^{D228G}$ was subsequently solved by molecular replacement using the refined Mnk2-KR structure and refined with CNS.

**Gel Filtration and Light Scattering**

Gel filtration chromatography was carried out with a SMART system using a Superdex 75 PC 3.2/30 column (Amersham). Experiments were performed at room temperature in 20 mM Tris/HCl, pH 7.5, 100 mM NaCl, 1 mM DTT at a flow rate of 0.04 ml/min. The molecular weight of Mnk2-KR was estimated using standard proteins (BioRad, Munich, Germany). Multi-angle laser light scattering with a 30 µM solution of Mnk2-KR in buffer A was performed as described elsewhere (Jauch et al., 2003).

**Kinase and Dot Blot Assays**

Mnk2, Mnk2-KR and Mnk2-KR$^{D228G}$ (2.5 µM) were activated using 140 nM pre-activated ERK2 and 50 µM ATP in 20 mM HEPES/KOH, pH 7.4, 10 mM MgCl$_2$, 0.25 mM DTT and 0.05 % (w/v) polyoxyethylene 20 stearylether (Brij 78; Sigma, Munich, Germany).
for 45 min at 30°C. In vitro kinase activity was assayed in the presence of 30 µM eIF4E-derived substrate peptide and 20 µM ATP in a buffer of 20 mM HEPES/KOH, pH 7.4, 10 mM MgCl₂, 0.5 mM DTT, 0.1 % (w/v) bovine serum albumin and 0.01 % (w/v) Pluronic F127 (Sigma). Reactions (40 min, 30°C) were terminated by EDTA addition. Substrate phosphorylation was detected based on a competitive fluorescence polarization format (Seethala and Menzel, 1998).

For the dot blots, 0.7-5 µM protein were incubated with a 10 µM mixture of ATP and α³²P-ATP (0.2 µCi; Hartmann Analytic, Braunschweig, Germany) in 10 mM Tris/HCl, pH 7.5, 100 mM NaCl, 10 mM MgCl₂ (30-60 minutes at room temperature). Proteins were transferred onto nitrocellulose membranes using the Minifold I vacuum filtration system (Schleicher&Schüll, Dassell, Germany), washed and detected by autoradiography.

Accession Numbers

Coordinates and structure factors have been submitted to the RSCB Protein Databank (http://www.rcsb.org/pdb/) with accession codes 2AC3 and 2AC5 and will be released upon publication.

Acknowledgments

We thank Monique Erling for excellent technical support in protein production and purification and Gleb Bourenkov (DESY, Hamburg) for help with diffraction studies. Work was in part supported by the Max-Planck Gesellschaft (HJ and MCW). CN is a fellow of the Boehringer Ingelheim Fonds. This work is part of the scientific research conducted in the International Ph.D. Program Molecular Biology - International Max Planck Research School at the Georg August University, Göttingen, Germany.
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Chapter II: The MAPK Interacting Kinase (Mnk) Subfamily is Regulated via a Novel Autoinhibitory Mechanism

Citation:


Statement of Contribution:

crystallization, data collection, structure solution, model building, structure refinement, structure analysis, examination of and comparison to available protein kinase structures, manuscript preparation
The MAPK Interacting Kinase (Mnk) Subfamily is Regulated via a Novel Autoinhibitory Mechanism

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MAP kinase interacting kinases 1 and 2 (Mnk1/2) regulate the expression of certain genes via phosphorylation of key factors involved in translation initiation. Here we present the crystal structure of the unphosphorylated Mnk1 catalytic domain showing that an otherwise structurally conserved element at the C-terminus of the activation segment of Mnk1 occupies a novel binding site alongside helix $\alpha$C. This reorganization seeds a circuit of structural rearrangements. Firstly, it causes closing of the interlobal cleft by reorienting the N-terminal lobe through interactions with $\alpha$C. Secondly, a phenylalanine which derives from a Mnk specific insertion is plunged into a hydrophobic pocket that is normally occupied by the phenylalanine of the Asp-Phe-Gly (DFG) motif (DFD in Mnks) of the Magnesium binding loop. The DFD motif, in turn, flips by almost 180° and occupies the ATP binding site. This conformation is stabilized by carboxyl-carboxyl side chain interactions of the DFD motif with the catalytic aspartate and a conserved glutamate residue of $\alpha$C. The C-terminal end of the activation segment is therefore employed as an autoinhibitory domain similar to the juxtamembrane domains of c-Kit and FLT3, respectively. Comparison of the Mnk1 structure with Mnk2 allows us to propose a model for the mechanistic dynamics that accompany the activation of Mnk subfamily members. It predicts a novel regulatory principle that is unique for the action of Mnks.
Introduction

Mitogen activated protein kinases (MAPK) interacting kinases (Mnk’s) comprise a small subfamily of serine-threonine kinases (Manning et al., 2002; Jauch et al., 2005). Mnk’s are activated through phosphorylation by the growth factor stimulated Ras- extracellular signal regulated protein kinases (ERK)1/2 and the stress induced p38-pathway (Fukunaga and Hunter, 1997; Waskiewicz et al., 1997). The two mammalian Mnk isoforms, Mnk1 and Mnk2, phosphorylate the eukaryotic initiation factor 4E (eIF4E) in vitro and in vivo (Waskiewicz et al., 1999; Scheper et al., 2001; Ueda et al., 2004). eIF4E binds the CAP structures of eukaryotic messenger RNA’s (Marcotrigiano et al., 1997) suggesting that Mnk dependent control of eIF4E phosphorylation affects CAP dependent translation (Waskiewicz et al., 1999). However, the effect of eIF4E phosphorylation by Mnks on global protein synthesis is under debate (Scheper and Proud, 2002). At the organismic level, it has been shown that mice deficient for Mnk1/2 develop normally implying that Mnks have no essential function (Ueda et al., 2004). However, in Drosophila the single Mnk orthologue, Lk6, was shown to regulate growth, body mass and lipid metabolism (Arquier et al., 2005; Reiling et al., 2005). The Lk6 phenotype was dependent on amino acid availability and has been interpreted to be a result of an intricate interplay with the mTOR pathway (Reiling et al., 2005). Furthermore, a most recent study indicates that Mnks control the translation initiation of specific mRNAs also by phosphorylating different targets (Buxade et al., 2005). This study demonstrates, that phosphorylation of the RNA binding heterogenous nuclear ribonucleoprotein A1 (hnRNPA1) diminishes the translation of the tumor necrosis factor alpha (TNF-α). Hence, Mnks have the potential to regulate inflammation by controlling the TNF-α levels.

Mnks have also been reported to affect the translation of RFLAT-1 as well as of viral RNAs (Nikolcheva et al., 2002; Walsh and Mohr, 2004). Therefore Mnk mediated phosphorylation appears to regulate the translation of specific mRNAs but the molecular details of the role of Mnks in translation initiation are still unkown. However, the involvement of Mnks in growth control, in lipid metabolism, viral translation and TNF-α regulation defines them as a potential targets for pharmaceutical intervention.
Recently we reported the crystal structure of the Mnk2 kinase region (Mnk2-KR) (Jauch et al., 2005) showing an extended conformation of the activation segment and a novel zinc coordinating site in a Mnk specific insertion, two features absent from other protein kinases. In addition, we noted the substitution of the highly conserved Magnesium binding Asp-Phe-Gly (DFG) motif by Asp-Phe-Asp (DFD) which adopts a conformation that obstructs ATP binding. However, the significance of the extended activation segment and the trigger for adoption of the inhibitory DFD motif conformation remained elusive. Here we present the crystal structure of the Mnk1 kinase region (Mnk1-KR), showing that it adopts a conformation different from the Mnk2-KR. The Mnk1-KR structure reveals a structurally rearranged activation segment that serves as an autoinhibitory module in the non-phosphorylated form. Buttressing of the activation segment against the N-terminal lobe closes the interlobal cleft and leads to the adoption of an ATP-competitive conformation of the DFD motif. Furthermore the results reveal how the Mnk-specific Asp of the DFD motif stabilizes this autoinhibited state. We propose a model of Mnk activation in which the N-terminal lobe, the Magnesium binding loop and the activation segment undergo distinct structural rearrangements to proceed in a stepwise fashion from an autoinhibited to an active state.

Results

Structure Solution and overall Architecture of Mnk1-KR

We produced a polypeptide spanning residues 37 to 341 of Mnk1 in *Escherichia coli* BL21 cells and purified it to homogeneity. The construct includes the catalytic domain of Mnk-1 and is henceforth referred to as Mnk1 kinase region (Mnk1-KR). As shown for Mnk2-KR (Jauch et al., 2005) this fragment recapitulates the activity of the full length protein, i.e. it phosphorylates an eIF4E peptide after activation by ERK2 *in vitro* (Mnk1 data not shown). Needlelike, tetragonal crystals of Mnk1-KR could be grown at 20°C using the vapor diffusion method. A 2.8Å dataset was collected at the Swiss-Light-Source (Table 1). After molecular replacement using a truncated model of the Mnk2-KR (Jauch et al., 2005) followed by density modification, an interpretable electron density was obtained. The model could subsequently be refined to R/R\textsubscript{free} factors of 23.5/28.0% (Table 1). The asymmetric unit contains two Mnk1-
Figure 1: Overall structure of Mnk1 in stereo representation (A) and primary sequence (B). If not indicated otherwise the coloring scheme is maintained in the following figures: N-lobe: black; C-lobe: gray; C-loop: yellow; DFG/D-motif orange; $\alpha$C helix and Lys-Glu pair: blue; activation segment: green. Connections which could not be traced in the 2Fo-Fc electron density are indicated with dotted lines. (B) residues known to interact with ATP are marked with closed circles, residues comprising the DFG/D-in pocket: green empty circles; residues comprising the DFG/D-out pocket: red empty circles. Mnk specific amino acid insertions are boxed and Mnk specific residues with functional relevance are highlighted with a red arrow. Phosphorylation sites are indicated with stars. Regions expected to form conserved secondary structure elements in the active state are show with quotation marks: ‘P+1 loop’, ‘$\alpha$EF’ and the ‘$\alpha$EF/$\alpha$F loop’. Note that those elements are not present in their functional configuration in the inactive Mnk1 structure. Regions not traced in the 2Fo-Fc electron density are shown as empty letters.
KR molecules which are related by a non-crystallographic two-fold axis. Molecule A exhibits lower temperature factors and a clearer electron density in several regions. The functionally important regions, however, are virtually identical in molecule A and molecule B. The final model spans residues 39-335 of Mnk1.

The Mnk1-KR preserves several global features of protein kinase (PK) architecture including the bilobal makeup (Knighton et al., 1991a). The N-terminal lobe contains a twisted sheet of five antiparallel β-strands (β1-β5) and the regulatory helix αC (Figure 1). It contains the key elements essential for ATP binding such as the Gly rich loop, the Lys78 from β3 and the Glu94 emanating from αC. The larger C-terminal lobe is predominantly α-helical and contains the elements required for substrate binding and phosphate transfer. It includes the catalytic loop (C-loop, residues 170-175), the DFD motif (residues 191-193) and the activation segment (Figure 1). Two segments within the Mnk1-KR exhibit strong conformational flexibility and could thus not be traced in the electron density. The first segment comprises the core of the activation segment including the activation loop and the P+1 loop (residues 197-222). The second segment spans the Mnk specific cysteine cluster and helix αG (residues 261-290).

*The activation segment reorients the N-lobe via helix αC*

In its canonical conformation as seen in most other protein kinases the P+1 loop as well as the succeeding helix α-EF are buried within the kinase core in an environment provided by the catalytic loop and the helices αF, αG (Knighton et al., 1991a). As a result, the P+1 loop is in a position to interact with the peptide substrate, e.g. with the residue downstream of the phosphorylation site (hence the name P+1 loop). The short helix α-EF is connected to helix αF through the α-EF/αF loop which varies in size and primary sequence between individual PKs (Nolen et al., 2004). In contrast, the region corresponding to α-EF and the α-EF/αF loop of the Mnk1-KR adopts a different conformation. This region is unwound and traverses the cleft between the N-terminal and C-terminal lobes. This way, the residues of the highly conserved Asp-Pro-Glu (APE) motif that usually form the helix α-EF come to lie at the N-terminal lobe, lining up with helix αC (Figure 2A). This conformation is stabilized by interaction of Glu225 (APE motif) and Glu228 (α-EF/αF loop) with Arg90 and Arg93 form αC. Significantly, residues corresponding to Arg90 and Arg93 are often bound to phosphate
Figure 2: N-lobe movement induced by the activation segment. Overall structure of Mnk1 (A) and Mnk2 (B). The DFD-motif, Phe-230, Arg-90, Arg-93, Glu225 and Glu-228 are shown in the ball-and-stick representation (Mnk1 numbering). Arg90 and Arg93 correspond to residues known to interact with phospho amino acids (Krupa et al., 2004). (C, D) Comparison of the N-lobes of Mnk1 (black, αC blue) and Mnk2 (yellow) after the superposition of the C-lobes (C-lobes not shown). (C) orientation as in (A,B). (D) rotated by 90° around the vertical axis.
moieties in active protein kinases (Krupa et al., 2004). Therefore, in inactive Mnk1, the region corresponding to \( \alpha\)-EF and the \( \alpha\)-EF/\( \alpha\)F loop is apparently positioned next to helix \( \alpha\)C by residues which in active PKs engage in alternative interactions. This observation suggests that phosphorylation will provide a trigger that releases the activation segment from helix \( \alpha\)C.

As a consequence of the unusual positioning of the \( \alpha\)-EF-\( \alpha\)-EF/\( \alpha\)F loop region, helix \( \alpha\)C and the remainder of the N-terminal lobe are displaced towards the C-terminal lobe and thereby narrow the interlobal cleft (Figure 2A). Thus, in the Mnk1 structure, the structural region corresponding to the P+1 loop and \( \alpha\)-EF adopts a conformation that is incompatible with catalysis and must therefore be rearranged for activity. Hence, the activation segment for Mnks spans the region located between the DFD motif and helix \( \alpha\)F. It therefore includes, in contrast to previous conventions (Nolen et al., 2004), \( \alpha\)-EF and the prolonged \( \alpha\)-EF/\( \alpha\)F loop (Figure 1).

The unusual conformation of the activation segment of Mnk1 may be the result of two amino acid insertions which are not present in most other Ca\(^{2+}\)/Calmodulin modulated kinases (CAMK) group PKs. Insertion 1 (I1, Figure 1) is located C-terminal of the DFG/D-motif and is disordered in the present structure. Insertion 2 (I2) affects a region corresponding to the \( \alpha\)-EF/\( \alpha\)F loop. Notably, residues of insertion I2 participate in the stabilization of the conformation seen in Mnk1 (e.g. Glu228 and Phe230).

The activation segment induces an ATP-competitive conformation

The vast majority of protein kinases bears an DFG motif at the beginning of the activation segment (subdomain VII) which shapes the ‘lip’ of the ATP binding ‘mouth’ of protein kinases at the interlobal cleft (Hanks, 2003; Hanks et al., 1988; Taylor and Radzio-Andzelm, 1994). The first aspartate of this motif is invariant among catalytically active protein kinases and is known to coordinate a magnesium ion that is essential for phosphoryl transfer (Adams, 2001). The region encompassing the DFG motif has therefore been termed the magnesium binding loop. Mnks contain a DFD motif at the corresponding position. We thus refer to the motif as DFG/D. In active kinases, the Phe of the DFG/D is situated in a structurally conserved pocket. In Mnk1, this pocket is provided by Leu98 and Thr97
(emanating from αC), His168 (upstream of the C-loop), Ile107, Ile189 and Leu161. This active DFG/D motif conformation is referred to as DFG/D-in conformation and its corresponding binding site will henceforth be referred to as DFG/D-in pocket.

The conformation of the activation segment as seen in the present Mnk1 structure has a dramatic effect on the DFD. As a result of the activation segment repositioning Phe230, comes to lie in the DFG/D-in pocket (Figure 3). It thereby displaces the DFGD. The DFD motif is in turn rotated by ~180° around the Φ angle of Asp191 (Φ_{Asp191} = -120) with respect to the DFG/D-in conformation of active state protein kinases (e.g. DAPK1 Φ_{Asp161}=55°; Figure 4A; Tereshko et al., 2001). As a result the Phe of the DFG/D (Phe192) occupies a hydrophobic pocket provided by Val63, Leu108, the ‘gatekeeper residue’ Phe124 and Leu177 which would normally accommodate the adenosyl moiety of ATP. The resulting DFG/D conformation is referred to as DFG/D-out. The DFG/D-out conformation has also been described for Mnk2 (Jauch et al., 2005).

*The Mnk-specific DFD stabilizes the DFD-out Conformation*

The DFD motif of Mnk1 participates in a network of polar interactions that explains the preference for the DFG/D-out conformation. Both, the invariant Asp191 and the Mnk specific Asp193 are engaged in tight carboxyl-carboxyl side chain interactions with active site residues (Figure 4B): (i) Asp191 binds to Glu93; (ii) Asp193 binds to Asp170. Glu94 and Asp170 correspond to residues which are invariant among catalytically active protein kinases (Hanks et al., 1988). Glu-94 emanates from the regulatory helix αC and is known to form an ion pair with Lys78 that is essential for productive ATP binding (Adams, 2001). This pairing is obstructed in Mnk1, since Asp191 of the DFG/D motif interacts with Glu94 (OD-Asp191:OE-Glu94) and with Lys78 (O-Asp191:Nz-Lys78). Asp170, which interacts with Asp193, corresponds to the catalytic aspartate of the C-loop.

Although carboxyl-carboxyl side chain interactions appear unusual, they have often been observed in proteins structures and are particularly abundant within the catalytic centers of enzymes (Flocco and Mowbray, 1995). The pH of the crystallization conditions for Mnk1 (pH 5.6) may have favored the stabilization of those interaction. However, acid-acid interactions have been observed even at basic buffer conditions (Flocco and Mowbray, 1995). The O-O distances between the two carboxylic acid groups reside in proximity of 2.6 Å and
Figure 3: The ATP binding pocket of active DAPK1 (A; PDB-id: 1ig1; (Tereshko et al., 2001)) and inactive Mnk1 (B). The molecules are in the same orientation as in Fig. 1 with the ATP binding regions blown up. (A) active DAPK1 contains the non-cleavable ATP analogon ANP-PNP (ball-and-stick) at the functional site. Note the permissive DFG/D-in conformation of the Magnesium binding DFG-motif. The ATP site blockade of Mnk1 (B) and Mnk2 (Jauch et al., 2005) is achieved by the inhibitory DFG/D-out conformation. The DFD-motif, Lys-78, Glu94, Asp-170, Asn-175 and corresponding residues of DAPK1 are shown in the ball-and-stick representation.
2.5 Å which is significantly shorter than the average O-O distance between non-acidic hydrogen donor/acceptor pairs. The latter observation was attributed to a proton sharing binding mode (Flocco and Mowbray, 1995). As described for other such carboxyl-carboxyl interactions Asp191:Glu94 as well as Asp193:Asp170 are stabilized by amines (Lys78) or amides (Asn175), respectively (see e.g. (Werten et al., 2002) for comparison).

**Discussion**

*The activation segment of Mnk1 is reorganized*

Previous studies have shown that in most protein kinases, the flexible portion of the activation segment is restricted to the activation loop, which is located between the DFG/D motif and the P+1 loop (Nolen et al., 2004). The P+1 loop interacts with the residue adjacent to the phosphorylation site of substrate peptides and therefore plays an important role in substrate peptide positioning (Knighton et al., 1991b). Within the Mnk subfamily, however, the activation segment is extended through two amino acid insertions (Figure 1). In addition, the region subject to structural rearrangements is strongly expanded since the flexible portion not only includes the activation loop but also the succeeding P+1 loop, the region corresponding to the short helix \( \alpha \)-EF and the \( \alpha \)-EF/\( \alpha \)F loop. Both the P+1 loop and the \( \alpha \)-EF helix occupy conserved sites within the vast majority of reported protein kinase structures, no matter whether the active or the inactive version of the enzymes were crystallized (Nolen et al., 2004). In contrast, the \( \alpha \)-EF helix and the \( \alpha \)-EF/\( \alpha \)F loop of Mnk1 unfold and traverse the interlobal cleft. Hence, the activation segment of Mnk1 is not only longer as compared to other PKs but also structurally rearranged in an unprecedented manner. This observation has profound consequences for the function of Mnk1.

*Variegation of the Juxtamembrane Theme: Autoinhibtion of Mnk1*

The previously reported Mnk2 structure displays an inactive conformation which is characterized by the ATP competitive DFG/D-out conformation (Jauch et al., 2005). The autoinhibition could be destabilized in Mnk2 by introducing the canonical DFG motif in place
Figure 4: Autoinhibition in c-KIT (A; PDB-id: 1t45) and Mnk1 (B). The autoinhibitory JM domain of c-KIT crosses the interlobal cleft coming from the N-terminus of the molecule (colored red). A region corresponding to helix αEF and the αEF/αF loop traverses the interlobal cleft in the opposite direction in Mnk1. Residues occupying the DFG/D-in pocket and the Phe of the DFG/D-motif are displayed in the ball-and-stick representation. Corresponding structural elements are colored equally. Note that the activation segment of c-KIT is fully visible in contrast to Mnk1.
of the Mnk specific DFD motif, suggesting a role of the Mnk-specific Asp. However, the reason for adopting DFG/D-out remained largely speculative in the case of Mnk2. The present Mnk1 structure sheds light on the structural basis for Mnk regulation and clarifies the role of the DFD motif.

Autoinhibition is a prominent strategy for protein kinase regulation which is imposed differently in individual cases and may affect various functional sites of the molecules (Huse and Kuriyan, 2002). Autoinhibiting regulatory domains outside the core of the protein kinase domain are employed, for example, in CAMKI (Goldberg et al., 1996), Twitchin (Kobe et al., 1996), c-KIT (Mol et al., 2004b) and Flt3 (Griffith et al., 2004). In the case of c-KIT and FLT3, two receptor tyrosine kinases of the platelet derived growth factor receptor (PDGFR) family, the autoinhibition is due to a juxtamembrane (JM) domain. This domain connects the transmembrane helix to the cytoplasmatic catalytic domain and functions as a regulatory element in many receptor tyrosine kinases (Hubbard, 2004). In c-KIT and FLT3, the JM domain causes autoinhibition by plunging a hydrophobic residue into the DFG/D-in pocket. As a result, the DFG/D-out conformation is adopted and catalytic activity is repressed (Figure 3; Griffith et al., 2004; Mol et al., 2004a). Although Mnk1 does not contain a JM domain, it is likewise autoinhibited by the induction of the DFG/D-out conformation (Figure 3, 4). The ‘molecular trick’ employed by Mnk1 parallels the mechanism described for c-KIT and FLT3. In contrast to c-KIT and Flt-3, however, Mnk1 does not bear an autoinhibitory domain outside the PK domain. Instead, Mnk1 induces DFG/D-out through its activation segment. To this end, Mnk1 inserts Phe230 emanating form the Mnk-specific insertion I2 into the DFG/D-in pocket. As a result, the DFG/D-in pocket is inaccessible for the DFG/D motif and the DFG/D-out conformation is enforced. The reorganized activation segment, in particular Phe230, may therefore constitute an autoinhibitory element which plays a key role for a Mnk specific regulatory mechanism. This element serves formally the same function as the JM domain of PDGFRs.

The Ins and Outs of DFG/D

Both Mnk1 and Mnk2 display the DFG/D-out conformation, which excludes ATP from entering its binding pocket. This observation suggests that the DFG/D-out conformation is a default state of inactive Mnk kinases. The Mnk1 structure provides an explanation for the
consequences of the Mnk specific Asp within the DFG/D-motif. This Asp interacts with catalytically important residues and thus stabilizes the DFG/D-out conformation (Figure 3). Hence, the DFD motif may facilitate the specific regulation of Mnks by stabilizing an inactive conformation.

Our observation highlights a possible significance of the DFG/D-motif for PK regulation. To date, the DFG/D-out conformation has been observed in only a fraction of the more than 50 protein kinases for which structural data are available (Mol et al., 2004b; Vieth et al., 2005). In some cases, the DFG/D-out conformation has been induced and/or stabilized by the binding of selective inhibitors such as in the structures of Birb796 bound to p38 (Pargellis et al., 2002), Cleevec bound to c-Abl (Nagar et al., 2002) and AAL-993 bound to VEGFR-2 (Manley et al., 2004). Notably, inhibitors targeting the DFG/D-out conformation are expected to bind with higher selectivity when compared to classical compounds which target the ATP binding pocket directly (Mol et al., 2004b). Hence, Mnks display a selective binding surface for specific inhibitors in their unphosphorylated state. The selectivity of this site is further pronounced by the presence of the Mnk specific Asp within the DFG/D motif.

*The Mnk1 structure differs from Mnk2*

Although their primary sequence is largely identical the crystal structures of Mnk1 and Mnk2 display pronounced differences in structural modules which are essential for kinase function (Figure 2, 4). These differences include (i) the conformation of the activation segment, (ii) the orientation of the two lobes towards each other and (iii) atomic details of the DFG/D-out conformation (Figure 2, 4). In the Mnk2 structure the activation segment protrudes from the PK core and helix $\alpha$F is prolonged by one turn. Mnk2 does also display the P+1 loop and helix $\alpha$-EF, a region which is either disordered or structurally rearranged in Mnk1. However, the P+1 loop and $\alpha$-EF are inserted into the symmetry equivalent neighbor molecules instead of becoming positioned at their canonical intramolecular binding site (Jauch et al., 2005). As a consequence, the interaction between the activation segment and the N-lobe is relieved and the occupation of the DFG/D-in pocket is reversed in Mnk2. As a result, the N-lobe of Mnk2 could snap back, the interlobal cleft could open up and the Lys-Glu ion pair would be formed (Figure 2B). When compared with Mnk1, the N-lobe of Mnk2 is tilted by approximately 10° leading to an opening of the ATP binding mouth of the kinase (Figure 2C,
D). A N-lobe movement would explain the absence of the polar interactions of the DFD motif in the Mnk2 structure.

**The mechanistic dynamics of Mnks**

Mnk1 and Mnk2 exhibit a sequence identity of 78% within their catalytic domains and their cellular function is largely redundant (Waskiewicz et al., 1999; Scheper et al., 2001). Hence, we expect that they behave similarly at the atomic level. We therefore assume that both structures provide snapshots of different structural states. These states are common among the Mnk subfamily members and can therefore be adopted by Mnk1 and Mnk2. Based on this assumption, we propose a model describing the structural basis for Mnk regulation. The model comprises 3 states: (I) the inhibited state, (II) the intermediate state and (III) the active state (Figure 5). State I is represented by the structure of Mnk1. In this state the PK is autoinhibited through a novel positioning of the reorganized activation segment. It displays an activation segment which traverses the interlobal cleft and aligns with helix $\alpha C$. As a result DFG/D-out is adopted and the interlobal cleft is closed. Several basic residues which are known to interact with phospho-residues in activated PKs engage in unique intramolecular contacts in state I. First, the Arg upstream of the catalytic loop (RD-Arg) undergoes a polar sidechain interaction with Asp-238 (Asp-273 in Mnk2). Second, basic residues from $\alpha C$ (Arg-90 and Arg-93) interact with the repositioned activation segment. Both, the RD-Arg and basic residues from $\alpha C$ are known to form the so-called RD-pocket which accommodates phospho-residues emanating from the activation segment (Krupa et al., 2004; Nolen et al., 2004). The interaction between such phospho-residues and the RD-pocket has been reported to have at least two effects: it positions the activation segment for substrate binding and it induces a closed conformation by re-orientating the N-lobe (Nolen et al., 2004).

It is well established that Mnk’s require phosphorylation at two sites within the activation loop to become activated (Waskiewicz et al., 1999). Phosphorylation is expected to disrupt the intramolecular interactions of RD-pocket residues as seen in state I. It is therefore likely that state I is destabilized in favor of the states II/III as a result of such phosphorylation events. We assume that state II, which is depicted by the structure of Mnk2, constitutes an intermediate state on the way to full activation (Jauch et al., 2005). The fact that state II is adopted by Mnk2 in the absence of phosphorylation can be explained by the unusual crystal
**Figure 5: A model of the Mnk activation cascade.** The different states are depicted schematically. The N- and C-terminal lobes are indicated with oval circles. The ATP binding site is highlighted with a transparent red box. αC (blue) and αF (gray) as well as the magnesium binding loop (orange) and the region corresponding to P+1 (state 2 and 3 only), αEF and the αEF/αF loop (green) are shown as cartoon. The Phe of the DFG/D-motif and the catalytically important Lys-Glu pair are shown as ball-and-stick. Arrows indicate movements of structural elements (see text for details of the model).
packing interaction. Despite this consideration, state II illustrates the structural consequences which occur, when the activation segment leaves its binding site seen in state I: On the one hand, the N-lobe is subjected to a pronounced reorientation leading to lobe opening and the formation of the Lys-Glu ion pair. On the other hand, the DFG/D-in pocket is no longer occupied by Phe-230. However, state II still displays several features of inactive state kinases.

For the adoption of the active state III, further structural rearrangements are required. Firstly, to enable productive ATP binding, the DFG/D-motif has to adopt the DFG/D-in conformation. This switch has in parts been observed in the previously reported structure of the Mnk2D228G active site mutant (Jauch et al., 2005). Secondly, an inward switch of the P+1 loop and α-EF is required. Without this switch, the canonical mode of peptide substrate binding as described for many PKs can not be pursued (Hubbard, 1997; Brown et al., 1999; Yang et al., 2002). We further assume a re-closure of the interlobal cleft which brings N-lobe residues such as the Gly rich loop and the Lys-Glu pair to their catalytic position. The lobe re-closure is probably induced by interactions of phospo-residues with RD-pocket residues from αC.

**Experimental Procedures**

**Cloning and Protein Preparation**

A cDNA fragment of human Mnk-1, corresponding to amino acid residues 37 to 341 and encompassing the kinase domain (KD) was amplified using the forward/reverse primer pair 5’CGGGATCCACTGACTCCTTGCCAGGAAAG/5’ACGCGTCGACCTATCCCTTTTCTGGAGCTTGCC (utilized restriction sites underlined) and was cloned into the BamHI and SalI sites of the vector pGEX-4T1 (Amersham, Sweden, cat. no. 27-4580-01). This construct allows prokaryotic expression of Mnk1-KR as a fusion protein with a N-terminal, thrombin cleavable glutathione S-transferase (GST) tag. The expression and purification of the Mnk-1KR was essentially identical to method described for Mnk2-KR (Jauch et al., 2005).
**Crystallization and Data Collection**

Needlelike crystals of wild type Mnk1-KR were grown at 20°C by vapor diffusion with a reservoir solution containing 20% (w/v) PEG3350, 0.2M Ammonium sulfate and 0.1 M Na-Citrate, pH 5.6 and were frozen (liquid nitrogen) in reservoir solution supplemented with 20 % glycerol. Diffraction data (Table 1) were collected on the PXII beamline (SLS, Villingen, Switzerland) at 100K on a MarResearch (Norderstedt, Germany) CCD detector and processed with the HKL package (Otwinowski and Minor, 1997).

**Structure Solution and Refinement**

The Mnk1-KR structure was solved by molecular replacement (MolRep; (Collaborative Computational Project, 1994) using a truncated version of Mnk2-KR as the search model. The resulting electron density was further modified by density modification and non-crystallographic symmetry (NCS) averaging. Model building was performed manually by interpreting the 2Fo-Fc electron density using Xfit (McRee, 1999). Refinement was performed with the CNS program excluding NCS restraints (Brunger et al., 1998). All data between 15.0 and 2.8 Å resolution were employed in the refinement, setting aside 5% of the reflections to monitor the free R-factor (Table 1). The molecular figures were prepared with PyMol (http://pymol.sourceforge.net/).
References


Tables

Table 1. Data collection and refinement statistics

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Supplementary Chapter III: Structures of Escherichia coli NAD Synthetase with substrates and products reveal mechanistic rearrangements

Citation:


Statement of Contribution:

structure refinement, model building, structure analysis and structural comparison, manuscript preparation
Structures of *Escherichia coli* NAD Synthetase with Substrates and Products Reveal Mechanistic Rearrangements*

Received for publication, November 22, 2004, and in revised form, January 26, 2005
Published, JBC Papers in Press, February 7, 2005, DOI 10.1074/jbc.M413195200

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Nicotinamide adenine dinucleotide synthetases (NADS) catalyze the amidation of nicotinic acid adenine dinucleotide (NAAD) to yield the enzyme cofactor nicotinamide adenine dinucleotide (NAD). Here we describe the crystal structures of the ammonia-dependent homodimeric NADS from *Escherichia coli* alone and in complex with natural substrates and with the reaction product NAD. The structures disclosed two NAAD/NAD binding sites at the dimer interface and an adenosine triphosphate (ATP) binding site within each subunit. Comparison with the *Bacillus subtilis* NADS showed pronounced chemical differences in the NAAD/NAD binding sites and less prominent differences in the ATP binding pockets. In addition, the *E. coli* NADS structures revealed unexpected dynamical rearrangements in the NAAD/NAD binding pocket upon NAAD-to-NAD conversion, which define a catalysis state and a substrate/product exchange state. The two states are adopted by concerted movement of the nicotinysyl moieties of NAAD and NAD, Phe-170, and residues 224–228, which may be triggered by differential coordination of a magnesium ion to NAAD and NAD. Phylogenetic structure comparisons suggest that the present results are relevant for designing species-specific antibiotics.

Nicotinamide adenine dinucleotide (NAD)\(^1\) is a ubiquitous carrier of reduction equivalents and functions as a cofactor in numerous metabolic reactions. NAD plays further roles in processes as diverse as calcium mobilization, DNA repair, and post-translational modification of proteins in eukaryotes. The final step of NAD biosynthesis is catalyzed by the enzyme NAD synthetase (NADS) and involves the conversion of nicotinic acid adenine dinucleotide (NAAD) to NAD (1) (see Fig. 1A). NAD synthetase activity was first described in the late 1950s (2, 3). Two types of NADS enzymes have been discovered; the first using preferentially ammonia as the amide donor, the second employing glutamine instead. The corresponding reaction schemes have been reported in 1967 (4).

\[
\text{ATP} + \text{NAAD} + \text{NH}_3 \rightarrow \text{AMP} + \text{PPi} + \text{NAD} \\
\text{REACTION 1}
\]

\[
\text{ATP} + \text{NAAD} + \text{l-glutamine} \rightarrow \text{AMP} + \text{PPi} + \text{NAD} + \text{l-glutamate} \\
\text{REACTION 2}
\]

In the course of the above reactions, the enzymes mediate an adenylyl transfer from ATP to the nicotinysyl moiety of NAAD, resulting in NAD-adenylate (see Fig. 1B) as a reaction intermediate. The second step of the reaction is the amidation of the nicotinysyl moiety using either ammonia (Reaction 1) or glutamine (Reaction 2) as an amide source.

NADSs from several prokaryotes, including *Bacillus subtilis* (bsu) (5), *Escherichia coli* (eco) (6, 7), and *Mycobacterium tuberculosis* (mtu) (8) have been cloned and characterized. Both *eco*NADS and *bsu*NADS exhibit a clear preference for ammonia over glutamine (2, 5, 7), whereas the *mtu*NADS can use both glutamine and ammonia as amide donors (8). Recently, the human (NADsyn1, here referred to as NADS1) and yeast (Qns1) NADSs have been characterized (9–11). Eukaryotic NADSs confer an additional N-terminal domain with nitrilase activity, which enables them to use glutamine as an amide donor (10).

NADS from *Bacillus subtilis* has been subjected to intensive crystallographic investigations. At first, the crystal structures of the *bsu*NADS apoenzyme, the complex of *bsu*NADS with ATP, and the complex of *bsu*NADS with the reaction intermediate NAD-adenylate had been reported (12–14). Subsequently, the picture was expanded by co-crystallizing *bsu*NADS with NAAD, AMP + PP\(_i\) (a combination of NAAD and ATP), and also with the ATP analog AMP-CP\(_i\) (15). More recently, a 1.0-Å resolution structure of *bsu*NADS in complex with the NAD-adenylate intermediate has been described (16). The combined structural studies unraveled the overall architecture of a NADS and delineated its mode of substrate binding. They defined separate binding sites for the ATP co-substrate (ATP binding pocket) and the substrate NAAD (NAAD/NAD binding pocket), and they revealed some details of the catalytic mechanism. NADS blocking agents have been suggested to be promising antibiotics for pathogens, such as *Bacillus anthracis*, which may be developed employing the *bsu*NADS crystal structures (17, 18).

In the present study, we provide high resolution crystal structures of NADS from *E. coli*, which shed light on evolutionary relationships among the enzymes *eco*NADS and *bsu*NADS.
In addition to the structure of the apoenzyme and complexes of the enzyme with AMP, NAD, and AMP+PP\textsubscript{i}, we present also the structure of NADS in association with its reaction product, NAD. This structure exhibits some unexpected rearrangements both in the active site and in the product molecules upon conversion of NAAD to NAD. The results reveal novel dynamical aspects for the reaction mechanism for ammonia-dependant NADSs. This work should also aid in the design of species-specific inhibitors for this class of enzymes.

**EXPERIMENTAL PROCEDURES**

Cloning, Expression and Purification—The *E. coli* nadE gene encoding NADS was PCR-amplified with *E. coli* DH5α cells as the template source and was inserted into the EcoRI site of the expression plasmid pRSETA (derived from pRSET6d (19)). The expression plasmid was designed to contain an N-terminal hexahistidine tag (tag sequence MAHHHHHHEF). The DNA construct was verified by automated sequencing, which predicted a protein product in accordance with SwissProt accession number P18843.

The protein was expressed in shaking cultures of *E. coli* BL21(DE3) cells at 32 °C overnight by the addition of 1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) mid-log phase of growth. The cells were harvested by centrifugation, resuspended in buffer A (50 mM Tris-HCl, pH 7.0, 100 mM NaCl, 0.2 mM phenylmethylsulfonyl fluoride) and lysed by sonication (Branson sonifier, Schwebach Gmünd, Germany). The His-tagged protein was captured on a nickel-nitriotic acid column (Qiagen, Duesseldorf, Germany) in buffer A, washed with buffer B (50 mM Hepes-NaOH, pH 7.0, 100 mM NaCl, 50 mM imidazole, 0.2 mM phenylmethylsulfonyl fluoride), and eluted in a linear gradient buffer C (50 mM Hepes-NaOH, pH 7.0, 100 mM NaCl, 800 mM imidazole, 0.2 mM phenylmethylsulfonyl fluoride). The NADS-containing fractions were pooled, dialyzed against buffer D (30 mM Hepes-NaOH, pH 8.0, 0.1 mM EDTA) and applied to a DEAE-cellulose column (DE-52, Whatman, Brentford, UK), washed with buffer E (20 mM Tris-HCl, pH 8.0), and eluted in a linear gradient to buffer F (20 mM Tris-HCl, pH 8.0, 0.400 mM NaCl, 0.2 mM dithiothreitol; 0.2 mM phenylmethylsulfonyl fluoride). The NADS-containing fractions were pooled, dialyzed against crystallization buffer (20 mM Tris-HCl, pH 7.0, 10 mM NaCl, 0.2 mM dithiothreitol, 0.2 mM EDTA), and concentrated by ultrafiltration (10-kDa molecular mass cutoff membrane; Amicon, Billerica, MA) to 19 mg/ml.

**Crystallographic Data Collection**—*E. coli* NADS apoenzyme was crystallized in hanging drops by vapor equilibration at room temperature against 12.5–15.5% polyethylene glycol 400, 50 mM MgCl\textsubscript{2}, 100 mM sodium acetate, pH 5.0–5.4, with a drop volume ratio of 2:1 (protein solution:reservoir). The addition of 1 μl of a suspension of lithium-meta-titantate significantly increased the nucleation rate and quality of the crystals. Crystals appeared and grew within a period of 1–14 days. NADS-ligand complexes were obtained by co-crystallization, adding 1.5–2 mM AMP (form II), ATP (form III), NAAD (form IV), or NAD (form V) to the mother liquor on one side. Data were collected using the HKL software package (20). The addition of 1 μl of a suspension of lithium-meta-titantate significantly increased the nucleation rate and quality of the crystals. Crystals appeared and grew within a period of 1–14 days. NADS-ligand complexes were obtained by co-crystallization, adding 1.5–2 mM AMP (form II), ATP (form III), NAAD (form IV), or NAD (form V) to the mother liquor on one side. Data were collected using the HKL software package (20). Crystals appeared and grew within a period of 1–14 days. NADS-ligand complexes were obtained by co-crystallization, adding 1.5–2 mM AMP (form II), ATP (form III), NAAD (form IV), or NAD (form V) to the mother liquor on one side. Data were collected using the HKL software package (20). The crystals of the enzyme complexes were obtained by co-crystallization and pertained to two different crystal systems (Table I). Employing diffraction data to maximum resolutions between 1.7 and 2.3 Å without intensity cutoffs, the models were refined to R\textsubscript{work} and R\textsubscript{free} values between 17.1 and 22.3%, maintaining good stereochemistry (Table I). All crystals comprised the full-length protein (residues 2–275), excluding the first methionine. No residues for the N-terminal hexahistidine affinity tag could be located in the electron densities. The short flexible loop P1 (residues 88–90) was not visible in the structures in which the ATP binding site is empty (forms I, IV, and V). A large loop P2 (residues 208–224; Fig. 2) could not be traced in either of the five crystal structures. Arg-186, located in a loop between β5 and α6 (see below for a description of the topology), exhibited a disallowed dihedral angle conformation in all five models, which was maintained because of unambiguous 2F\textsubscript{o} − F\textsubscript{c} electron density for this residue. The unusual conformation is stabilized through hydrogen bonds and ion interactions with Glu-22, Ser-26, Pro-183, and Leu-184.

**Global Structure and Comparison to the B. subtilis Enzyme**—

ecoNADS exhibits an α/β fold similar to the *B. subtilis* enzyme (Figs. 3 and 4A). 55% of all residues are found in α-helical conformation, 15% belonging to β-structures. A twisted five-stranded parallel β-sheet is sandwiched between the two large helices α5 and α6. All β-strands are separated by single α-helices, with the exception of β3 and β4, which are disrupted by two α-helices (α5 and α6). Thus, the core of ecoNADS comprises a typical Rossmann fold nucleotide binding motif.

In both space groups, C2 and P3\textsubscript{2}1, ecoNADS or its complexes crystallized with one protein molecule per crystallographic asymmetric unit. It has been well established that bacterial NADSs form homodimers (7, 12). In the present crystals, ecoNADS monomers are positioned close to crystallographic two-fold axes, which generate the functional dimers. The globular NADS homodimer has approximate dimensions of 77 × 60 × 56 Å. Based on an analysis with the protein-protein interaction server (26), ~2500 Å\textsuperscript{2} of combined surface area are buried in the dimer interface, encompassing 18% of the solvent-exposed surface area of the molecules. Contacts between the protomers are principally mediated by the α5–α6 region. The two pairs of helices align homotypically in an antiparallel fashion (Fig. 3). The dimer interface is predominantly nonpolar (75% of the participating residues) and involves 14 hydrogen bonds but lacks intervention water molecules. Therefore, ecoNADS qualifies as an obligatory dimer.

ecoNADS and bsuNADS are 55% identical in their amino acid sequences (Fig. 2). Consistently, they exhibit quite similar overall folds. Fig. 4A shows an overlay of the NAAD-containing models from both organisms, which exhibit an average root mean square deviation of 0.55 Å for 241 aligned C-α atoms. Five regions in the *E. coli* protein contain more than one residue deviating by >1 Å from the *B. subtilis* structure (Fig. 4A): region 1 (residues 37–42) (*E. coli* numbering), region 2...
Table 1
Crystallographic data

Data in parentheses for the last 0.05 Å (for form I last 0.1 Å).

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<th>V</th>
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Data collection

Resolution (Å) | 30.0–2.3 | 30.0–1.8 | 30.0–1.7 | 30.0–1.9 | 30.0–1.9 |

Unique reflections | 16,081 | 25,734 | 29,229 | 21,526 | 29,755 |

Re�undancy | 5.4 | 3.7 | 3.8 | 3.6 | 3.8 |

Completeness (%) | 95.1 (97.7) | 95.9 (92.1) | 91.6 (87.6) | 94.5 (91.7) | 99.5 (95.3) |

I/σ(I) | 14.4 (3.0) | 13.0 (2.9) | 14.7 (2.7) | 11.9 (2.3) | 19.6 (3.0) |

R_{cryst} (%) | 8.8 (45.9) | 10.5 (41.8) | 7.4 (42.8) | 11.7 (58.6) | 5.4 (36.7) |

Refinement

Resolution | 30.0–2.3 | 30.0–1.8 | 30.0–1.7 | 30.0–1.9 | 30.0–1.9 |

Re�undancy | 16080 | 25734 | 29229 | 21526 | 29758 |

Protein molecules/asymmetric unit | 1 | 1 | 1 | 1 | 1 |

Solvent atoms | 2008 | 2029 | 2051 | 2008 | 2008 |

Nucleotide atoms | — | 46 | 55 | 44 | 44 |

Mg^{2+} | — | 1 | 2 | 1 | 1 |

R_{work}/R_{free} (%) | 18.3/21.7 | 19.1/21.1 | 17.6/20.5 | 19.9/22.3 | 17.1/19.1 |

Average temperature factors (Å²)

Protein | 36.4 | 18.7 | 18.9 | 20.3 | 28.2 |

Water oxygens | 47.8 | 33.9 | 38.1 | 34.2 | 50.4 |

Nucleotide (NAD/ATP sites) | — | 19.5/45.1 | 17.1/38.6 | 29.0/— | 38.3/— |

Mg^{2+} | — | 42.9 | 59.2 | 33.5 | — |

Wilson B-factor | 39.8 | 26.5 | 26.8 | 24.5 | 33.3 |

Root mean square deviation from ideal values

Bond length (Å) | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 |

Bond angle (°) | 1.08 | 1.20 | 1.10 | 1.13 | 1.15 |

Ramachandran analysis

Preferred (%) | 93.6 | 93.9 | 93.9 | 94.5 | 92.3 |

Additionally allowed (%) | 6.0 | 5.7 | 5.3 | 5.0 | 7.3 |

Disallowed (%) | 0.4 | 0.4 | 0.8 | 0.5 | 0.4 |

Notes:

a R_{cryst} = \Sigma_{|hkl|} ||I_{calc}(hkl) - (|I_{obs}(hkl)|)||/\Sigma_{|hkl|} ||I_{calc}(hkl)|;

b R_{work} = \Sigma_{|hkl|} ||F_{calc}(hkl) - |F_{obs}(hkl)||/\Sigma_{|hkl|} ||F_{calc}(hkl)|;

R_{free} = \Sigma_{|hkl|} ||F_{calc}(hkl) - |F_{obs}(hkl)||/\Sigma_{|hkl|} ||F_{calc}(hkl)|;

hkl \in T - test set.

(c)—, not determined.

(residues 71–76), region 3 (residues 125–129), the far C terminus, and the C-terminal continuation of the P2 loop. Regions 1 and 2 exhibit single or double amino acid insertions in ecoNADS. Significantly, region 1 is intimately involved in binding of the adenylyl moiety in the NAAD/NAD binding pocket (see below). In contrast, the deviating regions 2, 3, and the far C terminus are distant from the substrate binding sites and are therefore not expected to exhibit effects on the substrate or product binding characteristics of the enzymes. The P2 loop has a fundamental impact on the catalytic mechanism (see “Architecture of the NAAD/NAD Binding Site”).

Design of the ATP Binding Site—In the present ensemble of crystal structures, the ATP binding sites of form II and III crystals contained AMP and AMP + PP_i, respectively, whereas the site was empty in the apoenzyme (form I), in the NAAD complex (form IV), and in the NAD complex (form V). Crystal form III was obtained by the addition of ATP to the crystallization drops, but the crystals contained AMP and pyrophosphate, indicating that hydrolysis of ATP took place during crystallization.

Each of the two ATP binding sites in the ecoNADS dimer is formed exclusively from one subunit. In crystal forms II and III, the ribose portions of the adenine nucleotides are engaged in hydrogen bonding interactions with the backbone oxygen of Gly-46 (to ribose O-2’) and the side chain hydroxyl group of Thr-160 (to ribose O-3’). The AMP adenine ring is held in place through stacking interactions with Arg-142, Ile-47, and Ser-48 (Fig. 5A). A hydrogen bond between N-6 of the adenine moiety and Gln-88 leads to a stabilization of loop P1, as indicated by clear 2F_o − F_c electron density for this protein region. In contrast, no electron density developed for loop P1 in the apoenzyme or in other crystals with unoccupied ATP binding sites (forms I, IV, and V). Flexibility in loop P1 is the only recognizable mechanism to yield access to the ATP binding site. Thus, the P1 loop serves as a lid, which is flexibly designed to allow ATP access but closes the ATP pocket upon substrate binding.

The side chain of Arg-82 exists in two conformations in the AMP co-crystal structures. In the major conformation, it interacts either with Gln-88 of the P1 loop and with N-1 of the adenine moiety and Gln-88 leads to a stabilization of loop P1, as indicated by clear 2F_o − F_c electron density for this protein region. In contrast, no electron density developed for loop P1 in the apoenzyme or in other crystals with unoccupied ATP binding sites (forms I, IV, and V). Flexibility in loop P1 is the only recognizable mechanism to yield access to the ATP binding site. Thus, the P1 loop serves as a lid, which is flexibly designed to allow ATP access but closes the ATP pocket upon substrate binding.
strate sensor or gate keeper for the ATP binding pocket.

As in most ATP-binding proteins, several magnesium ion binding sites have been reported for bsuNADS (14). In the present ecoNADS structures, a Mg\(^{2+}\) binding site (site I) is clearly occupied as well in crystal forms II (AMP), III (AMP+PP\(_i\)), and IV (NAAD), i.e. in structures that contain at least AMP at the ATP binding site or NAAD at the NAAD/NAD binding site. The Mg\(^{2+}\)-I ions show a typical octahedral coordination sphere, which in the AMP+PP\(_i\) co-crystal structure (form III) is composed of oxygens from the AMP α-phosphate, the two phosphate groups of the pyrophosphate moiety (former β- and γ-phosphates of ATP), the side chain carboxyls of Asp-51 and Glu-165 and a water molecule (Fig. 5C). In the AMP co-crystal structure (form II), which lacks the pyrophosphate molecule, and in the NAAD co-crystal structure (form IV), in which the ATP binding pocket is unoccupied, missing phosphate oxygens are replaced by water molecules in the coordination sphere.

In crystals of form IV (NAAD co-crystal), the nicotinylsyl portion of NAAD is oriented toward Mg\(^{2+}\)-I. The nicotinylsyl carboxyl group of NAAD is engaged in a hydrogen bond with a water oxygen, which belongs to the octahedral coordination sphere of Mg\(^{2+}\)-I (Fig. 5C). The latter water oxygen coincides with the position of an α-phosphate oxygen in forms II (AMP) and III (AMP+PP\(_i\)) structures. In contrast, the form V structure (NAAD co-crystal) does not contain the Mg\(^{2+}\)-I ion. Correlating with this observation, the nicotinylsyl moiety of NAAD is differently oriented from the corresponding portion of NAAD and is pointing away from the Mg\(^{2+}\)-I binding site (see “Architecture of the NAAD/NAD Binding Site”). Therefore, in the NAAD co-crystal structure, Mg\(^{2+}\)-I is seen as a direct link between the ATP binding pocket and the NAAD/NAD binding pocket. This direct interaction is lost upon formation of the product NAD.

A second magnesium ion (Mg\(^{2+}\)-II) has been reported in bsuNADS crystal structures (14, 15), where it is coordinated by the AMP α-phosphate, two pyrophosphate oxygens, two water molecules, and the backbone carbonyl oxygen of Thr-208 (B. subtilis numbering). In the present ensemble of structures, the Mg\(^{2+}\)-II ion is only seen in form III crystals (AMP+PP\(_i\) complex). Because Mg\(^{2+}\)-II is present only in the AMP+PP\(_i\) complex, the pyrophosphate (or by inference the β- and γ-phosphate groups of ATP) is required to capture a magnesium ion at site II in ecoNADS.

Comparison of the ATP Binding Sites of E. coli and B. subtilis NADS—Ordering of the P1 loop upon occupation of the ATP binding site has also been seen in bsuNADS. Therefore, loop P1 seems to be a conserved flexible element, the function of which is to close the ATP binding site upon the addition of a nucleotide. In ecoNADS, we have observed an additional conformational change in the side chain of Arg-82 upon AMP+PP\(_i\) binding, which is consistent with Arg-82 constituting a sensor for the occupation of the ATP pocket (Fig. 5A). In contrast, according to available models for the B. subtilis protein, the Arg-82 conformation of the nucleotide-bound forms is preformed in the bsuNADS apoenzyme. Thus, despite a high degree of similarity in the chemical composition of the ATP binding pockets, ecoNADS and bsuNADS seem to respond differently to nucleotide binding.

Significantly, in bsuNADS, the N-terminal part of loop P2 contributes the backbone carbonyl oxygen of Thr-208 (B. subtilis numbering) for coordination of Mg\(^{2+}\)-II in the ATP binding pocket. Loop P2 is ordered in structures of the Bacillus enzyme, in which nucleotides occupy the ATP pocket, portraying how this region can fold back onto the bulk of the protein (Fig. 4B). In contrast, loop P2 is disordered in the present structures of ecoNADS. The sequence of loop P2 is highly conserved between the two enzymes, and ecoNADS also comprises a threonine residue corresponding to Thr-208\(^{bsu}\) (Thr-210\(^{eco}\); superscripts identify the organism). However, although the AMP+PP\(_i\) co-crystal structure (form III) of ecoNADS contains a Mg\(^{2+}\)-II, the backbone of the P2 loop remains disordered. It is possible that one ligand of the Mg\(^{2+}\)-II coordination sphere in the ecoNADS form III structure, which has been modeled as a water molecule, actually corresponds to the backbone carbonyl of Thr-210\(^{eco}\) and that the P2 loop becomes partially ordered upon occupation of the Mg\(^{2+}\)-II site. However, full ordering of the P2 loop seems to be supported by crystal packing contacts. In the
bsuNADS crystal structures, residues Glu-214\textsuperscript{bsu} and Lys-215\textsuperscript{bsu} from the central portion of loop P2 are involved in salt bridges to a neighboring molecule of the crystal lattice. In contrast, the P2 region in both \textit{ecoNADS} crystal systems (C2 and P3121) faces a large solvent channel, creating a solution-like environment around this region of the enzyme.
In conclusion, we note that the N-terminal part of loop P2 can adopt various degrees of ordering, completely fixed (as in the structures of bsaNADS with nucleotides in the ATP pocket), partially ordered (as in the structure of AMP+PPi of ecoNADS), and completely disordered (as in the structures of ecoNADS, in which β- and γ-phosphates are lacking from the ATP binding site). To some degree, ordering of loop P2 thus seems to be dependent upon occupation of the Mg$^{2+}$·II site, the fixation of which in turn depends on the presence and mode of binding of the ATP β- and γ-phosphates or, after hydrolysis, the PP, leaving group. The interplay between the second Mg$^{2+}$ ion and the N-terminal part of loop P2 therefore correlates with the state of the nucleotide in the ATP binding pocket.

Architecture of the NAAD/NAD Binding Site—The two identical NAAD/NAD binding sites per homodimer are located at the dimer interface (Fig. 3). Both subunits provide residues for NAAD/NAD binding (in the following termed subunit A and subunit B). In the present structures, we observed three different nucleotides at the NAAD/NAD binding site: AMP in forms II (AMP co-crystallization) and III (ATP co-crystallization), NAAD in form IV, and NAD in form V. Thus, co-crystallization in the presence of adenine nucleotides (but without NAAD or NAD) leads to the occupation of the NAAD/NAD binding site by AMP.

Binding of the adenosyl portions of the nucleotides by the enzyme is identical in the co-crystal structures with NAAD (form IV) and NAD (form V), and the adenosyl parts are tightly fixed at the dimer interface by these interactions. The adenosine ring of the AMP portion of the nucleotides is sandwiched between His-260 from subunit A and Ile-40/Val-156 from subunit B. Tyr-33B (superscripts identify the subunit) is engaged in a hydrogen bonding interaction with the adenine N-1 atom. The adenylyl ribose interacts via its 3’-OH with Asp-180B (carbonyl oxygen) and via O-4’ with His-260A (atom N2’). The phosphate groups interact with the side chains of Lys-261A, Arg-140A, and Asn-136B and with the backbone nitrogen of Lys-173A.

In contrast, the nicotinylsyl parts of both NAAD and NAD are fixed to a lesser extent compared with the adenosyl parts, indicated by higher temperature factors. However, the conformation of both NAAD and NAD could be unambiguously determined in $F_{o} - F_{c}$ “omit” electron density maps (Fig. 5B). The nicotinic acid ring of NAAD is stabilized by the side chains of Phe-170A, Thr-172A, and Asp-176A. The NAD conformation in form V crystals deviates significantly from the NAAD conformation (Fig. 3C). Relative to the NAAD structure, the nicotinamide moiety of NAD is rotated around the O-5’-C-5’ bond (Fig. 5C). Supporting the alternate NAD conformation, Phe-170 undergoes a concomitant structural change, from stacking on the NAAD nicotinylsyl ribose to stacking with the nicotinamide ring of NAD (Fig. 5C). The conformation of Phe-170 in the apoenzyme structure resembles its preferred conformation in the NAD co-crystal structure. In contrast, the structures of ecoNADS, in which AMP is bound to the NAD site (forms II and III), exhibit a NAAD-like Phe-170 conformation.

The conformation of loop P2 also appears to be subjected to rearrangements in the course of the enzymatic reaction. Although loop P2 could be traced neither in the form IV (NAAD) nor in the form V (NAD) structures as mentioned above, the conformation of the residues 224–228, which comprise the first traceable part after the disordered stretch 208–223, is strikingly different in the NAD complex compared with the NAAD co-crystal structure (Fig. 5C). The regions of the two structures exhibit C-α root mean square deviation values of >3 Å. Resembling the behavior of Phe-170, the conformation of region 224–228 in form V (NAD) recalls the conformation of the apoform (form I), whereas its conformation in form IV (NAAD) is highly similar to the models with occupied ATP sites (forms II and III). Significantly, the preferred conformation of NAD in the NAAD/NAD pocket and the conformation of the surrounding enzyme...
regions (Phe-170 and residues 224–228) in the NAAD co-crystal structure, and vice versa, are mutually exclusive. Superposition of the NAD-ecoNADS and NAAD-ecoNADS co-crystal structures clearly show a steric hindrance for adoption of the NAAD-bound conformation of region 224–228 when NAD is present and vice versa (Fig. 5C). In this regard, it is important to note that the change in conformation of ecoNADS region 224–228 and of Phe-170 also correlates with the presence of Mg^{2+}, which we have shown above to be involved in orientation of the substrate NAAD, but which is lacking when NAD is present. Therefore, release of AMP + PP_i and concomitant loss of Mg^{2+}–1 may be involved in the adoption of the NAD/apoform structures of the enzyme.

In conclusion, two alternative states of the NAAD/NAD binding site exist in ecoNADS, represented by alternative conformations observed for Phe-170, residues 224–228 (and most likely the remainder of the P2 loop), and the nicotinyl moiety of NAAD and NAD, as well as the differential coordination of NAAD and NAD to Mg^{2+}–1. One conformation, which is observed in the apoform and in the NAD-bound form of the enzyme and which we term the “exchange (E) state,” seems to allow substrate binding and product release. The other conformation, which is formed upon occupation of the ATP binding site or in the presence of the substrate NAAD and which we term the “catalysis (C) state,” has to be adopted for substrate conversion. It is noteworthy that these differences affect both the ATP binding pockets and the NAAD/NAD binding pockets and are therefore indicative of a structural communication between these sites during catalysis.

**Comparison to the NAAD/NAD Binding Site of bsuNADS**—Region 1, pointed out in the global comparison above (Fig. 4A), provides the docking environment for the adenylyl moiety in the NAAD/NAD binding site (Fig. 4C). Its amino acid composition is decisively different in ecoNADS and bsuNADS (Figs. 2 and 4C). *E. coli* region 1 contains a higher number of bulky, hydrophobic, or aromatic residues: Tyr-37 instead of Thr, Pro-38 instead of Gly, a Phe-39 insertion and Ile-40 instead of Ala (Figs. 2 and 4C). Despite these chemical differences, similar binding of the adenylyl moiety of NAAD or NAD is achieved by the two enzymes. Because of the phylogenetic variability in region 1, the NAAD/NAD pocket may be of particular interest for the development of species-specific antibiotics. Additional minor variations at the NAD binding environment, such as the replacement of Val-156 and Trp-256 in ecoNADS by Leu-153 and Arg-253 in bsuNADS, could be of further relevance for rational inhibitor design.

We have outlined above how the N-terminal portion of loop P2 (residues 208–224) is involved in nucleotide binding at the ATP site through coordination of the Mg^{2+}–1 ion. In bsuNADS, Ala-209 and Leu-211 from the same region of the P2 loop...
stabilize the nicotinysyl of bound NAAD. These interactions are not seen in ecoNADS. Supported by the crystal packing (see above), these interactions may therefore contribute to the fixation of loop P2 in bsuNADS structures. Interestingly, the C-terminal part of loop P2 also participates in substrate binding in bsuNADS. It is involved in the interaction with the nicotinysyl moiety of NAD-adenylate in the Bacillus enzyme (PDB code 1KQP). The diol of the nicotinysyl ribose hydrogen bonds to the carboxyl group of Glu-223. Glu-223 is substituted by alanine in the E. coli enzyme, which consequently loses the interaction site to the nicotinysyl ribose.

**DISCUSSION**

In the present manuscript, we presented the high resolution crystal structures of ecoNADS as well as the structures of the enzyme with various substrates and products. The ensemble of

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**Fig. 5. Structural rearrangements.**

A, close-up stereo view of the ATP binding pocket of the ecoNADS apoenzyme (carbon atoms are in brown, and other atoms are colored as described in the legend to Fig. 4) superimposed onto the same region of the AMP complex structure (carbons in light blue). Residues lining the binding pocket are labeled in single letter style, and Arg-82 (R82) is highlighted. The arrow indicates repositioning of the Arg-82 side chain upon nucleotide binding. The dashed lines indicate hydrogen bonds of Arg-82 and of Gln-88 to the adenine base of AMP. B, Fo–Fc difference omit map contoured at the 2.5σ level with a radius of 3 Å around NAAD (top) and NAD (bottom). The cofactors are shown as sticks. Arrows around NAAD bonds indicate the main rotations necessary to produce the NAD conformation. The arrow in NAD indicates the nitrogen of the amide group introduced in the course of the reaction. C, comparison of the NAAD/NAD binding pocket when complexed with NAD (carbon atoms and backbones traces in light blue) and when complexed with NAAD (carbon atoms and backbones traces in brown) after global superpositioning of the enzyme structures. Other atoms are color-coded as above. Mg²⁺–I is indicated as a green sphere coordinated to the Asp-51 and Glu-165 carboxylate groups and to water molecules (small red spheres). One water molecule also hydrogen bonds to the carboxylate of NAAD (dashed line). The different conformations of regions 224–228, C-terminal of the disordered loop P2, are indicated by the backbone traces with side chains.
structures encompasses the first co-crystal structure of a NADS with its enzymatic product, NAD. The structures confirm and significantly expand the picture that has emerged from the structural studies of \textit{bsu}NADS and provide evidence for a wide ranging conformational plasticity in this enzyme family. This plasticity manifests itself in (i) structural reorganizations upon binding of different substrates, (ii) a conformational cross-talk between the ATP binding site and the NAAD/NAD binding pocket, and (iii) conformational adjustments during substrate conversion.

Flexible Elements Mediate Structural Reorganizations upon Substrate Binding—The mode of ATP binding appears to be largely invariant between \textit{E. coli} and \textit{B. subtilis} NADS. In both enzymes, loop P1 becomes fixed upon nucleotide binding to the ATP pocket and thus functions as an evolutionarily conserved lid for the ATP binding site. However, a significant difference between the two enzymes is seen in residue Arg-82, which undergoes side chain reorganization upon ATP or AMP+PP\textsubscript{i}, binding in \textit{eco}NADS, whereas its ATP-bound conformation is preserved in the \textit{bsu}NADS apoenzyme. Thus, Arg-82 represents a hitherto unrecognized dynamical part of the ATP binding site in addition to loop P1.

Different from loop P1, the P2 loop is disordered in all structures of \textit{eco}NADS presented herein. In contrast, it is ordered in various \textit{bsu}NADS structures, where several of its residues engage in direct or indirect contacts to the substrates ATP and NAAD. In its fixed conformation in \textit{bsu}NADS, both the N- and the C-terminal ends of the P2 loop are involved in substrate binding. At the N terminus, the backbone carboxyl of Thr-209\textsubscript{bsu} coordinates to Mg\textsuperscript{2+}-I in the ATP pocket. In addition, residues Ala-209\textsubscript{bsu} and Leu-211\textsubscript{bsu} participate in the positioning of the nicotinysyl moiety of NAAD (15). At the C terminus, both side chain carboxyl oxygens of Glu-223\textsubscript{bsu} hydrogen bond to O-2’ and O-3’ of the nicotinysyl ribose of nucleotides in the NAAD/NAD pocket. The highly similar sequence of the loop in \textit{eco}NADS and the similar modes of substrate binding suggest that P2 can interact in a similar fashion with the substrates in the \textit{E. coli} protein, except for Glu-223\textsubscript{bsu} (which is replaced by alanine), but that these substrate interactions are transient. Thus, the main function of loop P2 seems to be the mediation of access to the substrate binding pockets and perhaps shielding them from the environment in the course of the reaction. Complete ordering of this loop in \textit{bsu}NADS structures seems to be observed, because it is supported by favorable crystal lattice contacts.

d\textit{eco}NADS Dynamics in Response to Substrates and Products—Our data provide the first co-crystal structure of a bacterial NADS in complex with the reaction product, NAD. We have shown that the nicotinysyl moiety of enzyme-bound NAD exhibits higher flexibility than the adenosyl portion and prefers a conformation, which is different from that of NAAD. The conformational difference in the substrate and product coincides with alternate conformations of Phe-170 and residues 224–228 following the C-terminal end of the disordered loop P2. Phe-170 stacks on the nicotinysyl ribose of NAAD and on the nicotineamide ring of NAD in the co-crystal structures. The conformation of residues 224–228 in the NAAD complex sterically interferes with the conformation of bound NAD (Fig. 5C). These observations suggest a structural reorganization prior to or accompanying product release. Catalysis requires an orientation of nicotinic acid toward the ATP binding site (14). We speculate that product release requires a torsion within NAD, which in turn necessitates the reorganization of Phe-170 and the P2 loop. A possible trigger for this reorganization may be the Mg\textsuperscript{2+}-I ion. In contrast to the previously reported complex structure of \textit{bsu}NADS with NAAD (15), Mg\textsuperscript{2+}-I is present in our \textit{eco}NADS-NAAD co-crystal structure. The nicotinysyl carboxylate group of NAAD hydrogen bonds to a water of the Mg\textsuperscript{2+}-I coordination sphere. On the other hand, NAD in the form V structure points away from the Mg\textsuperscript{2+}-I site. Thus, Mg\textsuperscript{2+}-I seems to act as an orientation device for the NAAD nicotinysyl moiety, which is activated by adenyl transfer from ATP in the course of the reaction. Because Mg\textsuperscript{2+}-I under physiological conditions presumably binds to the enzyme in complex with ATP, it prepares the NAAD conformation for this adenyl transfer when the ATP pocket is ready to deliver the AMP moiety.

The data presented herein for \textit{eco}NADS, including the conservation of the mode of substrate and metal ion binding, are in general agreement with the reaction mechanism described for \textit{bsu}NADS (16), suggesting that it is conserved for bacterial NADSs. However, our ensemble of \textit{eco}NADS structures allows us to expand the mechanism to structural changes taking place in the enzymes upon substrate conversion. We have found that the active center of \textit{eco}NADS exists in different conformations depending on the occupation of the ATP and NAAD/NAD binding sites. We refer to these conformations as the catalysis (or C-) state and the substrate/product exchange (or E-) state. The C-state is displayed in three of our crystal structures (forms II, III, IV), i.e. when \textit{eco}NADS is complexed with AMP, AMP+PP\textsubscript{i}, or NAAD. The E-state predominates in the apoform of \textit{eco}NADS (form I) and in the complex with NAD (form V). The two states are distinguished by three major features: (i) the side chain conformation of Phe-170, (ii) the conformation of the P2 loop as represented by its C-terminal extension, comprising residues 224–228, and (iii) the occupation of the first magnesium site (Mg\textsuperscript{2+}-I). The two states reflect the structural requirements in pursuing two interdependent processes. The E-state allows substrate binding and product release, and the C-state provides the environment for adenylation and amida
tion of the substrate NAAD. As described above, Mg\textsuperscript{2+}-I may be the element crucial for switching between the two states, because it seems to orient the NAAD nicotinysyl moiety. Following the enzymatic reaction, AMP, PP\textsubscript{i}, and Mg\textsuperscript{2+}-I are released, shifting the equilibrium toward the E-state.

The \textit{NAAD/NAD Binding Pocket May Be a Target Site for Future Design of Species-specific Antibiotics—}NADS has been proposed to be a promising target for the development of antibiotics to cure \textit{e.g.} anthrax (17, 18). Knowledge of the results of structural comparisons among NADS from different organisms may allow development of inhibitory compounds that exhibit organism specificity. Although the structures of \textit{eco}NADS expectedly revealed a similar overall fold as \textit{bsu}NADS (Figs. 3 and 4A), significant structural differences can be observed in detail. In particular, although the architectural alterations in the NAAD/NAD binding sites of \textit{E. coli} and \textit{B. subtilis} NADS do not affect the binding of the natural substrate, their chemically quite different makeup may provide a handle for specific inhibitor design (Fig. 4C).

In contrast to bacterial NADSs, which use exclusively ammonia as the amide donor (5, 7), NADS1 from human uses both ammonia and glutamine (11) (the ammonia-specific human NADS2 was later found to derive from a bacterial contamination (9)). Although many catalytic residues are invariant, human NADS1 contains several amino acid substitutions and peptide insertions at the ATP and the NAAD/NAD binding sites (Fig. 2). These differences confirm the suitability of the NADS enzymes for the design of antibiotics, because it should be possible to alleviate side effects in humans.

Acknowledgments—We thank Petra Birle and Tatjana Krywcun for help in enzyme preparation and crystallization, Dr. Hans-Georg Beisel and Dr. Peter Hof for initial help in structure solution, and Prof. Dr. Herbert Jackie for support in data analysis.
REFERENCES
Supplementary Chapter IV: The zinc finger-associated domain of the Drosophila transcription factor Grauzone is a novel zinc-coordinating protein-protein interaction module

Citation:


Statement of Contribution:

cloning, protein expression, protein purification, biochemical assays, light scattering, circular dichroism, structure refinement, model, building, structure analysis, manuscript preparation
The Zinc Finger-Associated Domain of the *Drosophila* Transcription Factor Grauzone Is a Novel Zinc-Coordinating Protein-Protein Interaction Module

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Introduction

A large number of protein domains, collectively referred to as zinc fingers, bind zinc ions through various combinations of four cysteine and/or histidine residues (Berg and Shi, 1996; Klug and Schwabe, 1995). The coordination of a zinc ion allows such modules to adopt compact three-dimensional structures which are otherwise too small to maintain extensive hydrophobic cores (Grishin, 2001). The prototype zinc finger proteins contain a 30 amino acid C2H2 motif (Miller et al., 1985; Pavletich and Pabo, 1991). In addition to this motif, a number of variants were found which are distinguished by zinc-coordinating residues, the secondary structural elements contributing the zinc ligands and/or topology (Krisha et al., 2003; Laitly et al., 2001). While a large number of zinc fingers function as DNA binding domains (Pavletich and Pabo, 1991; Rosenberg et al., 1986; Wolfe et al., 2000), it becomes increasingly clear that they can also mediate interactions of the protein with RNA, other proteins and small molecules of other kinds (Krisha et al., 2003; Mackay and Cossley, 1998; McCarty et al., 2003). Furthermore, zinc-coordinating folds may confer specific enzymatic activities on proteins (Lorick et al., 1999).

The C2H2 zinc finger motif characterizes one of the most abundant eukaryotic protein families (Lander et al., 2001). C2H2 zinc finger proteins frequently contain additional protein modules in their N termini (Collins et al., 2000) such as the mammalian Krüppel-associated box (KRAB) or the insect zinc finger associated domain (ZAD) (Lespinet et al., 2002). Both KRAB- and ZAD-containing C2H2 zinc finger proteins are characterized by lineage-specific expansions in the respective genomes (Chung et al., 2002; Lespinet et al., 2002; Looman et al., 2002). The KRAB domain acts as a transcriptional repressor module (Margolin et al., 1994; Peng et al., 2000), whereas the function of ZAD is unknown.

In silico studies have recently identified more than 90 ZAD proteins in *Drosophila*, accounting for ~28% of the total C2H2 zinc finger proteins of this organism (Chung et al., 2002). This family of proteins is characterized by a conserved constellation of four cysteines within the ZAD, the chromosomal clustering of the corresponding genes, and the lack of homologs in noninsect genomes (Chung et al., 2002). Although the identification of expressed sequence tags ensured that the majority of ZAD-encoding genes of the *Drosophila* genome are transcribed, only four annotated members of the gene family have been examined in some detail. Two of them, *Serendipity delta* (Sry-δ) and *grauzone* (grau), were shown to encode transcriptional activators of the egg-polarity gene *bicoid* (Payre et al., 1994) and the gene *cortex* which encodes a Cdc20-like APC subunit, respectively (Chen et al., 2000). The third factor, the Dorsal-interacting protein 1 (DIP1), was identified in a yeast two-hybrid screen as an interaction partner of the dorso-ventral polarity transcription factor Dorsal (Bhaskar et al., 2000). Finally, *Zeste-white-5* (Zw5) was reported to confer enhancer-blocking activity by association with the boundary element *scs* and may thereby participate in chromatin structuring by providing an insulating activity (Blanton et al., 2003; Gaszner et al., 1999). *grau*, *Zw5*, and *Sry-δ* have also been characterized by mutant analysis showing that they carry vital functions (Chen et al., 2000; Crozatier et al., 1992; Gaszner et al., 1999). In two cases, single amino acid replacements within the ZAD, such as cysteine 7 by tyrosine in Sry-δ (Crozatier et al., 1992) and arginine 4 by glycine in Zw5 (Gaszner et al., 1999), were the cause of lethality, implying that conserved amino acids within the ZAD are essential for its function.

Here we present the structure and biochemical features of the ZAD of Grauzone (ZADgrau). Grauzone is a 570 amino acid long transcriptional regulator characterized by the N-terminal ZAD and an array of eight C-terminal C2H2 zinc finger domains which mediate the binding of Grauzone to a promoter element of its target gene *cortex* (Chen et al., 2000). In genetic terms, *grau*
is necessary for the proper transcriptional activation of the gene cortex, coding for a Cdc20-like APC subunit (Chu et al., 2001). Maternal loss-of-function mutations of grau cause an irregular growth arrest in meiosis II, whereas homozygous grau mutant embryos, which have received maternal grau activity, develop into normal adults. The grau mutant phenotype suggests that grau is exclusively required during oogenesis and cortex is its only target gene (Chen et al., 2000). Our results show that ZADgrau exhibits a C4 zinc-coordinating fold with a novel treble-clef-like structure. The crystallographic data and additional biochemical evidence indicate that ZADgrau is a protein interaction module with the capability to form homodimers and suggest that ZAD provides a taxon-specific means for the assembly of C2H2 transcription factor complexes.

Results and Discussion

Sequence alignments of a total of 91 Drosophila C2H2 proteins and corresponding proteins of a variety of other insects such as Anopheles gambiae revealed a conserved N-terminal sequence motif that contains two invariant pairs of cysteines (Figure 1A) (for details see Chung et al., 2002). Due to the specific association with C2H2 family members of zinc finger proteins, this motif was termed zinc finger-associated domain (ZAD). In order to characterize this protein motif, we studied the ZAD of the transcription factor Grauzone (ZADgrau). The ZADgrau open reading frame (amino acid residues 2–90; Figure 1A) was fused to Glutathione S-transferase (GST) and the recombinant fusion protein was produced in bacteria (see Experimental Procedures). After an initial capture step, the ZADgrau portion of the fusion protein was liberated by protease digestion, purified by column chromatography, and crystallized (see Experimental Procedures). SDS-PAGE analysis of dissolved crystals revealed that they contain only the ZADgrau protein (data not shown). The fact that ZADgrau crystallizes provides first evidence that ZADs encompass an independently folding protein module.

ZAD is a Zinc Binding Protein Module

Based on the four conserved cysteine residues, ZADs have been suggested to comprise metal binding modules (Chung et al., 2002; Lander et al., 2001; Lespinet et al., 2002). Because metal association is expected to contribute significantly to the stability of the fold, heat denaturation experiments were performed in the presence and absence of EDTA. In the absence of EDTA, ZADgrau was stable for at least 15 min at 60°C, whereas in the presence of EDTA (5 mM) a considerable portion of the protein precipitated when heated above 50°C for 15 min (Figure 1B). In these experiments, care was taken to exclude divalent metal ions from the purification procedure. The result therefore suggests that ZADgrau contains an endogenous metal center that confers stability on its structure. In order to test this proposal, we performed an X-ray fluorescence scan. With this approach, the characteristic emission lines for zinc were observed (Figure 1C). This result and the finding that the anomalous diffraction from the zinc center could be exploited to solve the structure with a native crystal (see below) demonstrate that ZADgrau includes a zinc ion.

Structure Solution and Quality of the Model

The primary sequence of ZADgrau is shown in Figure 1A. The crystal structure of ZADgrau was solved de novo by a two-wavelength multiple anomalous dispersion experiment around the K-edge of an endogenous Zn2+ ion. A segmented poly-alanine model could be manually fitted to the experimental electron density map and allowed the subsequent incorporation of side chains. The sequence assignment was guided by well-defined aromatic residues and by the two pairs conserved cysteines (C4, C7, C53, and C56; see Figures 1A and 2A), which coordinated the Zn2+ ion (Figure 1D). Three regions in flexible loops (V11-C17, D22-E27, and E41-E47; see Figures 1A and 2A) initially displayed weak densities and were fitted during the subsequent rounds of refinement and manual model building. These loop regions, which are made up of nonconserved residues, lack substantial intra- and interchain contacts. Thus, they gained above-average temperature factors during the refinement. The main chain electron density of the molecule remained fragmented around position S12–A14 and D22 in the final 2Fo–Fc map, consistent with high internal flexibility and lack of crystal packing contacts in these regions. All other side chains of the refined model, except those of some hydrophilic surface residues, were entirely covered by the final 2Fo–Fc map. No electron density developed for the C-terminal 9 residues (positions 82–90; Figure 1A) of ZADgrau and the 5 N-terminal vector-derived residues that are contained in the recombinant protein. The final model therefore encompasses the residues D2–S81 of ZADgrau that could be traced unambiguously. Presumably because of the considerable fraction of flexible residues at the termini, which could not be accounted for in the final model, the refinement converged with an R factor/Rfree factor of 24.1%/26.7%. Convergence at these numbers is consistent with a rather high B factor extracted from a Wilson plot (52.6 Å²), which is comparable to the averaged B factor of the final structure (52.9 Å²).

During all refinement steps, 5% of the reflections were set aside to monitor the Rfree factor (Table 1). Of the final model, 91.9% of the residues resided in the preferred regions of the Ramachandran plot, 6.8% in the additionally allowed areas. Only the single residue K29, which was well defined in both the experimental and the final model-derived maps assumed an unconventional φ/ψ conformation even after manual interference. The mean positional error of the model was estimated at 0.15 Å (Luzzati, 1952).

Structural Properties of ZADgrau

The structural portion of ZADgrau resembles the letter “b.” Its approximate dimensions are 60 × 30 × 30 Å (Figures 2A and 2B). The N-terminal body of the “b” comprises a globular fold structured around a zinc ion. The C-terminal stem is formed by a long α helix (α2, positions 54–80) that contains almost one-third of all residues of the domain. Residues 82–90 are not included in the ZAD consensus sequence (Chung et al., 2002).
Figure 1. ZAD as a Zinc-Coordinating Fold

(A) Alignment of the Grauzone ZAD with its closest paralog, CG15073, in the Drosophila melanogaster genome and with a putative Anopheles gambiae homolog (ZAD only fragment, see Experimental Procedures). Asterisks mark 100% conservation in all three ZADs; red characters, invariant cysteine pairs; yellow boxes, conserved hydrophobic or aromatic amino acid residues.

(B) Portion of ZADGrau remaining soluble after incubation at the indicated temperatures in the absence (-EDTA) and in the presence of 5 mM EDTA (+EDTA). Reduction of the soluble fraction is clearly visible when ZADGrau is heated above 50°C in presence of the chelator.

(C) X-ray emission spectrum of a native ZADGrau crystal irradiated with X-radiation of \( \lambda = 1.0 \text{ Å} \). I, ZnKα line; II, ZnKβ line; III, Compton scattering; IV, elastic scattering.

(D) Electron density maps around the zinc center in ZADGrau. Cyan, experimental MAD electron density map (1σ); magenta, anomalous difference Fourier map (10σ), generated with the anomalous differences at the peak wavelength and the phases obtained after solvent flattening. The zinc-coordinating cysteine residues are labeled and shown in ball-and-stick. If not mentioned otherwise, all structural figures were prepared with Bobscript (http://www.strubi.ox.ac.uk/bobscript/) and rendered with Raster3D (Merritt and Bacon, 1997).

and are disordered in the present structure. Presumably, they provide a flexible linker to the remainder of the molecule, i.e., an acidic region followed by an array of eight C2H2 zinc fingers in the case of Grauzone (Chen et al., 2000).

The arrangement of the different ZAD structural elements is summarized in Figures 2A–2C and detailed in the Figure 2 legend. The fold of ZADGrau appears to be critically dependent on the zinc coordination, a result that is consistent with the EDTA-dependent precipitation of ZADGrau shown above. The two pairs of coordinating cysteines are approximately 50 residues apart. Zinc coordination, therefore, links the \( \beta2-\alpha2 \) transition region at the center of the molecule with the N terminus of the domain likely to solidify the structure. In this view, the position of the zinc ion allows the fold to constrain the slanting angle of the C-terminal helix, \( \alpha2 \), with respect to the long axis of the \( \beta \) sheet (Figures 2B–2C). The relative orientation of this long helix and the \( \beta \) sheet is further defined by helix \( \alpha1 \), which rests with one of its surfaces on the end of the \( \beta \) sheet distal to the zinc ion and the adjacent loop regions. With a neighboring surface, helix \( \alpha1 \) is in contact with the N-terminal and central portions of helix \( \alpha2 \). Both the \( \alpha1 \) sheet and the \( \alpha1-\alpha2 \) associations are based on extensive hydrophobic contacts forming a considerable hydrophobic core (\( \alpha1 \): V29, V33, L34, H37, and F38; \( \beta1/\beta2 \): L18, I20, and I52; \( \alpha2 \): W57, V60, F63, H64, and Y67; Figures 1A and 2A–2C). In addition, the conserved H37 residue (atom NE2) of helix \( \alpha1 \) hydrogen bonds at the edge of this core to the
conserved H64 (atom ND1) of helix α2. In conclusion, the L2/α1/L3 region between the two central β strands and the zinc ion provide platforms which orient the C-terminal helix relative to the globular portion of the ZADGrau molecule.

To further support the overall structure of ZADGrau obtained by X-ray crystallography (Figures 2A–2D), we took circular dichroism spectra of the molecule in solution (data not shown). The calculated secondary structure content (45%–50% helix, 5%–10% strands) obtained from these recordings compare favorably to those seen in the crystal structure (37% helix, 6% strands). The conservation of length, the predicted secondary structure, the arrangement of the annotated secondary structure elements and the critical amino acid residues of ZADs (Chung et al., 2002) (Figure 1A) suggest that the present ZADGrau crystal structure provides a prototype for ZAD folding.

Structure Comparison

Next we compared the ZAD fold with known structures by manual inspection, with special emphasis on reported zinc finger fold groups (Krishna et al., 2003). We found that ZAD exhibits a structural relation to treble clef zinc finger (TCZF) domains. Figure 3A shows a comparison of ZAD with the structure of the C-terminal C4 TCF of chicken GATA-1 (PDB ID code: 2gat) (Tjandra et al., 1997), revealing a similar topology of the secondary structure elements. The treble clef element has been identified among the members of seven different SCOP fold groups including RING fingers, LIM domains, ribosomal proteins L24E and S14, and nuclear receptor-like fingers (Grishin, 2001; Krishna et al., 2003). Thus, despite the lack of significant sequence similarity between ZADs and TCZFs, which prevented recognition of their relation in the past, ZAD can be regarded as a novel, atypical TCFZ that exerts several distinct features. Unlike the canonical zinc knuckle of...
Table 1. Crystallographic Data

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Phasing

| Resolution (Å) | 20.0–2.1 |
| Heavy atom sites | 1 |
| Centrics | 0.50 |
| Acentrics | 0.61 |
| Phasing power |
| Centrics | 1.67 |
| Acentrics | 1.97 |
| FOM |
| Before DM | 0.42 |
| After DM | 0.78 |

Refinement

| Resolution (Å) | 15.0–2.0 |
| Number of reflections | 6590 |
| Number of nonhydrogen atoms | |
| Protein | 644 |
| Zn^{2+} ions | 1 |
| Water oxygens | 56 |
| Rwork (%) | 24.1 (29.6) |
| Rfree (%) | 26.7 (33.7) |
| Rmsd from ideal geometry |
| Bond lengths (Å) | 0.009 |
| Bond angles (°) | 1.20 |
| Average B factors (Å²) | |
| Protein | 50.6 |
| Zn^{2+} ion | 37.2 |
| Water oxygens | 80.3 |
| Wilson B factor | 52.6 |
| Rmsd B factors (Å²) | |
| Main chain bonds | 1.9 |
| Main chain angles | 3.3 |
| Side chain bonds | 3.5 |
| Side chain angles | 5.2 |
| Ramachandran analysis (%) | |
| Preferred | 91.9 |
| Additionally allowed | 6.8 |
| Disallowed | 1.4 |
| Mean residual error (Å) | 0.15 |

Data for the last 0.05 Å in parentheses. DM, density modification (solvent flattening); rmsd, root-mean-square deviation.

**Figure 3. Structural Comparison**

Ribbon (A) and electrostatic surface representation (B) of ZAD_{N\text{term}} (left panels) and the C-terminal TCZF from chicken GATA-1 (right panels). Color coding for the ribbon plot is as in Figure 2. In the surface plot, red indicates negative potential, blue positive potential. The orientation is the same as in Figure 2B. This and all other surface images were prepared with SwissPDBViewer (Guex and Peitsch, 1997).
Figure 4. The ZAD<sub>gra</sub> Dimer

(A) Stereo ribbon plot of a ZAD<sub>gra</sub> dimer as seen in the crystal. The two subunits (Mol I and Mol II) are colored red and blue; the two zinc centers are depicted in ball-and-stick. N- and C termini are labeled. The orientation of the rear subunit is the same as in Figure 2A.

(B) Stereo representation of residue interactions in the dimer interface, 90° from the view in (A). The two ZAD<sub>gra</sub> subunits are colored as in (A) but are rendered semitransparent to clearly reveal the interacting residues in ball-and-stick. All residues and the molecular termini are labeled.

(C) Mapping of the contact residues within the dimer (green) onto the surface of a ZAD<sub>gra</sub> monomer (gray). The orientation of the molecule is the same as that of the red subunit in (A).

(D) Mapping in orange of conserved residues on the surface of ZAD<sub>gra</sub> in two diametric views. The left orientation is the same as in (C), indicating that the largest conserved surface patch and the dimer interface largely coincide.

be a prerequisite for extending the length of helix α2, without loosing its defined orientation relative to the β sheet. Finally, helix α2 displays a conserved pattern of hydrophobic residues in ZAD (Figure 1D; see also below) which is absent from other TCZFs. ZAD can therefore be classified as a distinct subgroup of TCZF domain structures.

Crystal Structure Suggests ZAD<sub>gra</sub> Homodimers

The evolutionary restriction of ZADs to certain C2H2 zinc finger genes in insects, the chromosomal clustering of the majority of these ZAD-containing genes and the unique folding characteristics of the domain next to the DNA binding domain of transcriptional regulators such as Grauzone strongly argue that the ZAD is associated with a specific and distinct biological function. However, the mere similarity to the fold group of the TCZFs does not allow any conclusions about the specific role of ZAD, because treble clef motifs embody functions as diverse as binding to nucleic acids, proteins as well as small ligands and some may even exert enzymatic activity (Grishin, 2001).

However, the crystal structure of ZAD<sub>gra</sub> clearly supports a model, in which ZAD represents a protein-protein interaction module involved in homodimerization. Figure 4 shows that in the crystal two ZAD<sub>gra</sub> molecules are associated through a 2-fold axis in an isologous head-to-tail fashion. As revealed by the protein-protein interaction server (http://www.biochem.ucl.ac.uk/bsm/PP/server/), this contact buries ~1000 Å² of accessible surface area (ΔASA). This value seems on the lower side when compared to known homodimers (Jones and Thornton, 1996). However, the interaction surface of ZAD<sub>gra</sub> covers a total of close to 20% of the entire surface area. More importantly, a large number of amino acid residues, which are strongly conserved among ZAD
family members (for details see Chung et al., 2002), are responsible for the contacts between the two subunits (Figure 4B). When the conserved residues are mapped on the surface of ZAD$_{Grau}$, it becomes obvious that the largest conserved surface patch closely coincides with the presumed dimer interface (Figures 4C and 4D). In particular, hydrophobic residues of the long C-terminal helix $\alpha_2$ (F63, F66, I70, and Y77; Figure 1A) build up major parts of the contact interface (Figure 4B). As a consequence, 72.5% of the amino acid residues of the dimer interface are nonpolar. The presumed dimer interface is thus designed very differently from the remainder of the surface, which is lined with polar residues and exhibits a highly negative electrostatic potential (Figure 3B). In the region where the tip of helix $\alpha_2$ from one molecule contacts the globular portion of the other subunit, some intermolecular hydrogen bonds are observed as well (Q74-R5; Y77-E47; Figure 4B).

Because of the involvement of its hydrophobic side chains, the presumed homodimerization mode provides a straightforward explanation for the amphipathic design and the unusual length of helix $\alpha_2$ (Figure 2A). Since, in addition, both helix $\alpha_2$ and the globular portion of ZAD are involved in this dimerization mode, the importance of restraining their relative orientations by the inserted L2/ $\alpha_2$/L3 module and the zinc coordination becomes obvious. Further supporting its significance, the present dimerization may explain the lethal phenotype observed with the conserved R4 (corresponding to R5 in Grauzone) mutated to glycine in Zw5 (Gaszner et al., 1999). Its side chain is positioned by ionic interactions with D46 to engage in a hydrogen bond with Q74 of the neighboring molecule (Figure 4). We note, however, that there are a number of other contacts between the two subunits that could stabilize subunit interactions. It is therefore possible that the mutation simply perturbs the globular structure of the N terminus which might preclude dimerization. In any case, the analysis of the crystal packing strongly suggests a functional homodimerization of ZAD$_{Grau}$.

Besides the above symmetrical contact between two ZAD$_{Grau}$ molecules, other crystal contacts were observed. Some of these are not functional because of an unreasonably small $\Delta$ASA. Other contacts did not involve conserved residues. One rather intimate alternative association takes place through the crystallographic 4-fold screw axis. However, this symmetry element gives rise to heterologous contacts, leaving the bonding potentials of ZAD unsaturated. The latter interaction mode could therefore lead to the formation of larger oligomers. We therefore examined the arrangement of ZAD in solution.

**ZAD$_{Grau}$ Forms Dimers in Solution**

The quaternary structure of ZAD$_{Grau}$ in solution was investigated by two approaches. First, we performed chemical crosslinking experiments with glutaraldehyde (Figure 5A). In contrast to SNAP-25, which was reported not to self-interact (Fasshauer et al., 1999), and in parallel with the dimerizing GST, ZAD$_{Grau}$ could be efficiently crosslinked to the dimer state, but no higher oligomers were observed. Secondly, multiangle-laser-light-scattering following size exclusion chromatography yielded strong evidence for homodimerization in solution (Figure 5B). ZAD$_{Grau}$ eluted as a single symmetrical peak from various size exclusion columns with different optimal separation regimes. Comparison of the elution times with those of reference proteins was consistent with ZAD$_{Grau}$ dimers (data not shown). The scattering signal at 632.8 nm across the elution peak indicated a molecular weight of 21.11 kDa, which matched the theoretical dimer mass (21.14 kDa) almost perfectly (Figure 5B). This high congruence and the lack of a monomer or a trimer signal, strongly suggested a monodisperse dimer solution.

Collectively, our results provide strong evidence for ZAD$_{Grau}$ homodimerization under near-physiological salt conditions. They leave the 2-fold symmetrical association of ZAD$_{Grau}$ in the crystal (Figure 4) as the only dimerization mode, which is consistent with all observations.
Because ZADGrau is an independently folding unit, dimerization is expected to prevail within the context of the full-length Grauzone transcription factor. By extension of this finding, a general function of ZADs could be to provide dimerization modules that mediate homodimer and/or heterodimer formation among closely related members of the ZAD transcription factor family. In support of this observation, homodimerization was also reported for the ZAD-containing transcription factor Sry-Δ (Payre et al., 1997; Ruez et al., 1998). It is noteworthy that Sry-Δ comprises a deviated version of ZAD family members and could thus not be considered as prototype for ZAD function (Chung et al., 2002).

ZADGrau Lacks DNA Binding Features

The majority of zinc finger proteins were shown to be nucleic acid binding proteins (Berg and Shi, 1996; Laity et al., 2001; Pavletich and Pabo, 1991; Wolfe et al., 2000). Because of this ample documented function, we asked whether the ZADGrau structure has features to support nucleic acid binding as well. In order to obtain first hints, we compared the ZAD with the structure of the C-terminal treble clef module of the chicken erythroid transcription factor GATA-1 (Tjandra et al., 1997), a representative of sequence-specific DNA binding TCZFs. Figure 3B shows that the two molecules display opposite electrostatic surface potentials. Whereas the TCZF of GATA-1 exhibits an almost continuously electropositive surface potential that facilitates a close association with the negatively charged sugar-phosphate backbone of the DNA, the ZADGrau is almost completely wrapped into negative potential. This observation strongly argues against DNA binding features of ZAD.

Superimposition of one subunit of a ZADGrau dimer on the structure of DNA-bound GATA-1 TCZF (Tjandra et al., 1997) revealed that the second ZADGrau module faces the minor groove of the DNA and would thus collide with the DNA backbone (data not shown). Furthermore, Grauzone mutants, which lack the ZAD-containing region of the protein, had unaltered DNA binding properties in vitro, whereas the part of Grauzone protein that contains the ZAD in the absence of C2H2 zinc fingers had no DNA binding activity (Chen et al., 2000), supporting the argument that ZADGrau has no DNA binding features. Since the key positions and features of the amino acid residues are conserved among ZADs (for an alignment, see Chung et al., 2002), it appears unlikely that any of them carries DNA binding properties. We therefore propose that ZADs, in general, are protein interaction modules involved in homodimerization and/or the formation of protein complexes. To further support this conclusion, we also compared the ZADGrau dimer with the DNA-bound dimer of other members of the TCZF fold group, the glucocorticoid receptors (Luisi et al., 1991; Grishin, 2001). In these receptor molecules, the DNA binding helix equivalent to ZAD helix α2 is employed as a recognition element, which is positioned in the DNA major groove. Two DNA binding domains interact through their C-terminal helical extension, which have no equivalent in ZADs. These C-terminal extensions yield a spacing between the two recognition helices from neighboring subunits, which is large enough to position them both into neighboring turns of the DNA major groove. Therefore, the dimerization mode of ZADs is fundamentally different from the interaction mode seen in the DNA binding domains of glucocorticoid receptors, consistent with these molecules serving different functions.

Experimental Procedures

Sample Preparation and EDTA-Dependent Precipitation

The DNA coding for Grauzone ZAD residues 2–90 was PCR amplified, cloned into the Not1/EcoR1 sites of the pGEXT-3 vector, and recombinantly expressed as a GST-fusion protein in a BL21(DE3) E. coli strain. Cells were grown to an OD600 0.7–1.0 at 30°C, shifted to 20°C, and induced with 0.7 mM IPTG overnight. Cells were harvested by centrifugation and resuspended in phosphate-buffered saline containing Complete protease inhibitor tabs (Roche) and traces of lysozyme. Cells were lysed by sonication and the insoluble fraction was removed by centrifugation. GST-ZADGrau was captured using Glutathione (GSH)-Sepharose 4B (Amersham) beads in disposible columns following the manufacturer’s suggestions. Samples were eluted with 50 mM Tris-HCl, pH 8.0, 50 mM GSH. Thrombin was directly added to the eluate to 1 U/100 μg protein and incubated overnight at room temperature. The protein identity was confirmed by peptide fingerprinting and subsequently purified with a 1 ml Resource-Q 15 cm anion exchange column applying a linear 20 ml salt gradient (20 mM Tris-HCl, pH 8.0, 0–1 M NaCl), a second GSH column and a Superdex 25/60 75 pg gel filtration (GF) column (GF buffer: 10 mM Tris, pH 7.5, 100 mM NaCl, 5 mM DT). Samples were concentrated to 8 mg/ml via 20 5000 Da molecular weight cutoff concentrators (Vivascience). For EDTA-dependent precipitation studies, 10 μM ZADGrau samples in GF buffer either lacking EDTA or containing 5 mM EDTA were incubated for 15 min at room temperature, 37°C, 50°C, and 60°C. Samples were pelleted and the supernatant was analyzed by SDS-PAGE.

Crystalization and Data Collection

Crystalization screens were carried in the sitting drop vapor diffusion format on 24-well Crychem plates (Hampton Research, Laguna Niguel). 1–4 μl of an 8 mg/ml ZADGrau solution in GF buffer were combined with 1 μl reservoir buffer and equilibrated against a 400 μl reservoir. Crystals grew at 22°C with 0.2 M ammonium acetate, 0.1 M sodium citrate, pH 5.6, and 30% polyethylene glycol 4000 in 3–10 days.

After transfer into Paratone-N (Hampton Research) and removal of residual mother liquor, crystals could be shock frozen in a liquid nitrogen stream. Diffraction data were collected at 100 K on the HASYLAB beamline BW6 (DESY, Hamburg, Germany; http://www-hasylab.desy.de/facility/experimental_stations/stations/BW6.html). X-ray fluorescence spectra of native crystals clearly indicated zinc emission lines (Figure 1B). A single such crystal yielded complete anomalous data sets at the Γ-maximum of the K-edge (λ = 1.2828 Å) and at a remote wavelength (λ = 1.0500 Å). The data were recorded on a MarResearch (Norderstedt, Germany) CCD detector and processed with the HKL package (Otwinowski and Minor, 1997) (see also Table 1).

Structure Solution, Model Building, and Refinement

If not mentioned otherwise, programs from the CCP4 collection (CCP4, 1994) were employed for the structure solution. The location of a single Zn2+ ion could be deduced from anomalous difference Patterson maps at the Γ-maximum (program RSPS) or by direct methods (SHELX, http://shelx.uni-ac.gwdg.de/SHELX) and was used to phase the data with the remote data set as the reference (MLPHARE). The hand of the heavy atom was revealed during solvent flattening (DM).

Prominent secondary structure elements were manually fitted to the electron density with MAIN (Turk, 1996). Subsequently, these regions were decorated with side chains guided by the prominent features of the aromatic residues and by four cysteine residues, which coordinated the Zn2+ ion. The connecting loops were sequentially traced by alternating refinement of the partial model (CNS) (Brünger et al., 1998) and model building guided by the experimental
and 2F-, F electron density maps. Refinement included standard procedures of positional and B-factor optimization, a round of simulated annealing, and positioning of 56 water molecules into spherical peaks of the F- maps. The final step was a TLS refinement with REFMAC5 with the structure divided into four rigid bodies (the Zn²⁺ ion, residues 2–19 and 51–54, residues 20–50, and residues 55–81). No data within 15.0–2.0 Å were excluded, and during all refinement steps the same set of reflections (5%) was used to monitor the R free (Table 1). The geometry of the final model was analyzed with PROCHECK (Laskowski et al., 1993) (Table 1), secondary structure elements were extracted with PROMOTIF (http://www.biochem.ucl.ac.uk/bsm/promotif/promotif.html).

CD Spectroscopy

For far UV-CD measurements, the sample was dialyzed against 10 mM NaHPO₄, pH 7.9. The spectra (190–260 nm) were recorded with a JASCO J-720 spectropolarimeter (Jasco Corporation, Tokyo, Japan) at 20°C in a 0.1 cm cuvette with a scanning speed of 50 nm/ min. 30 scans were averaged and baseline corrected. The spectra were analyzed using the neural-network-based CD deconvolution software CDNN v2.1 (Böhm et al., 1992).

Chemical Crosslinking

The crosslinking reactions were carried out at 25°C in 20 mM Tris-HCl, pH 7.5, 100 mM NaCl with 15 μM samples of ZADmax GST (28 kDa), and SNAP-25 (25 kDa) (Fasshauer et al., 1999) and 1 mM glutaraldehyde. The reactions were stopped at selected time points by boiling in SDS sample buffer.

Multiangle-Laser-Light Scattering

The experiment was performed on a HR-10/30 Superdex-200 size exclusion column (Amersham) connected to a UV spectrometer and the Dawn and Optilab instruments XY (Wyatt Technology Corp.). 200 μl of a 25 μM ZADmax sample were chromatographed in 20 mM Tris-HCl, pH 8.0, 150 mM NaCl. The UVabsorption, the light scattering at 632.8 nm and the differential refraction of the elution profile were monitored. Spectra were analyzed with the Astra software package (Wyatt, 1990). We also checked ZADmax migration on an analytical Superdex-75 column in comparison to protein molecular weight standards.

Identification of a Putative Grauzone ZAD Homolog in the Anopheles gambiae Genome

We searched the whole Anopheles gambiae genome using the ZAD HMM described in Chung et al., 2002 and the Wise package 2.2.0 (Binney et al., 1996). All identified ZADs were aligned with all Droso phila melanogaster ZADs using ClustalW 1.8.1 (Thomson et al., 1994), false positive hits were eliminated. The alignment was used to construct a neighbor-joining tree with ClustalW. Based on this tree, we identified the closest relative of the Grauzone ZAD in the Anopheles gambiae genome. The alignment of the Grauzone ZAD with the closest paralog, CG15073, of Droso phila melanogaster and the putative Anopheles homolog was performed with ClustalW using default parameters.

Acknowledgments

We thank our colleagues in the labs for their various contributions, T. Siddiqui for kindly providing SNAP-25, G. Dowe for sequencing, and Stefan Pabst for help with MALLS analysis. Work was supported by the Max Planck Society. H.-R.C. is a fellow of the Boehringer Ingelheim Fonds.

Received: June 30, 2003
Revised: July 7, 2003
Accepted: July 9, 2003
Published: November 4, 2003

References


Accession Numbers

The structure has been submitted to the Protein Data Bank (http://www.rcsb.org/pdb) under accession number 1PZW.
General Discussion

Zinc Coordinating Protein Modules and Intermolecular Recognition

Zinc ions are found in a plethora of proteins that belong to a variety of functional and structural classes (Krishna et al., 2003). Besides acting as co-factors in catalytic reactions, e.g. in certain proteases (Hernick and Fierke, 2005), zinc ions function to stabilize structural modules remote from catalytic centers and enable the formation of small and highly compact folds (Krishna et al., 2003). In the latter case, zinc ions contribute to the architectural integrity of protein modules which are often involved in intermolecular recognition (Laity et al., 2001). In this study, zinc ions have been identified in two unrelated classes of proteins (Chapter I, Supplementary Chapter IV). The first class is represented by the zinc finger associated domain (ZAD) of the Drosophila transcription factor Grauzone.

The ZAD constitute a ~80 amino acid sequence motif that is specific for insects and genes encoding the domain are expanded in their genomes (Chung et al., 2002). The ZAD of the Drosophila transcriptional activator Grauzone represents a prototype member of this insect specific domain class (Harms et al., 2000; Chung et al., 2002). It has been demonstrated that the ZAD constitutes a protein-protein interaction module which enables dimerization of ZAD containing proteins in an antiparallel fashion (Supplementary Chapter IV). In the ZAD, the zinc ion is coordinated by a cluster of four Cys residues. This arrangement leads to the formation of a spherical and compact N-terminal portion of the domain from which a C-terminal α-helix emanates. This α-helix accommodates most of the conserved and hydrophobic amino acid residues required for dimerization (Supplementary Chapter IV). Although the ZAD has some architectural similarities with the large group of the so-called treble clef zinc fingers (Grishin, 2001), it exhibits unique features which led to the definition of a novel protein domain family (Zf-AD, synonym ZAD; Pfam accession number PF07776). Since the ZAD is almost exclusively associated with transcription factors of the canonical C2H2 zinc-finger type, it may contribute to the regulation of gene expression likely by mediating the assembly of distinct ZAD bearing proteins on clustered target sequences.
A second zinc coordination site has been detected within the catalytic domain of Mnk2 (Chapter I). In this case, the zinc ion is coordinated by a Cys cluster located within an amino acid insertion upstream of helix αG that is not found in other PKs. This insertion is phylogenetically conserved within the Mnk subfamily. A zinc coordination site as integral part of a PKD is unprecedented for this class of enzymes. The zinc site shapes a C-terminal extension of the enzyme. This extension is present in all protein kinases but absent in, for example, many metabolite kinases (Kannan and Neuwald, 2005). The C-terminal part of PKs and in particular helix αG has therefore been proposed to provide a scaffold for protein-protein recognition (Kannan and Neuwald, 2005; Kim et al., 2005). Hence, although the precise function of the zinc containing structural motifs of Mnks remains elusive, it is likely involved in the binding to substrates and/or regulator proteins. Taken together, the results obtained for both the Mnks and the ZAD highlight the significance of zinc ions in shaping protein domains and provide novel examples of zinc containing protein-protein recognition motifs.

**Mechanistic Plasticity of Biological Catalysts: Lessons from Mnk1/2 and NADS**

The structural studies on Mnk subfamily members and on the ecoNADS (see Chapters I, II and Supplementary Chapter III) provide different snapshots of highly dynamic enzymes which illustrate mechanistic rearrangements of their functional elements. The ecoNADS in complex with different substrates and products, as well as the three structures of Mnk subfamily members, permit to draw a dynamic picture of the respective enzymes. The plasticity of Mnk subfamily members is posed by alternative conformations of closely related isoforms and active site mutants, whereas the structural changes of NADS are depending on the association with different substrates and products. Binding of ATP to NADS is accomplished through an induced fit mechanism that leads to the reorganization of a short loop and amino acid side chains. The NAD site of NADS adopts alternative conformations depending on whether the substrate NAAD or the product NAD is bound. This structural rearrangement is accompanied by a reorientation of the ligand itself and reflects necessities which enable, on one hand, the catalytic reaction, and on the other hand, product release (Supplementary Chapter III).
The crystal structures of the PKDs of Mnk1, Mnk2 and the active site mutant Mnk2^{D228G} reveal a dynamic behavior which distinguishes the Mnk subfamily from other PKs. The dynamic behavior is manifested in pronounced conformational differences of the isoforms Mnk1 and Mnk2 despite their almost identical amino acid sequence. Mnk1 and Mnk2 vary with respect to the relative orientation of structural subdomains (N- and C-terminal lobe), the conformation of the activation segment and the side chain orientation of active site residues (Chapter I and II). These conformational differences can be attributed to distinct snapshots of different states of activity which are common to members of the Mnk subfamily of PKs. The structure of the Mnk2^{D228G} mutant demonstrates how a single amino-acid substitution affects the stability of the catalytically relevant magnesium binding loop of the PK domain. In conclusion, the results suggest a novel regulatory mechanism that includes a gradual interconversion of the conformational states depicted in the structures of Mnk1, Mnk2 and Mnk2^{D228G}.

*The Structures of Mnk1 and Mnk2 Reveal a Novel Mechanism of Protein Kinase Regulation*

Although the more than 500 human PKs exhibit a very similar overall architecture, individual members function in distinct biological pathways and often respond to specific signaling events (Krebs, 1983; Manning et al., 2002; Kannan and Neuwald, 2005). At the structural level, this specificity is achieved by a diverse set of regulatory mechanisms which distinguish individual PKs and adjust their activities (Huse and Kuriyan, 2002). As a consequence, most PKs exhibit intricate structural features which are specific and facilitate their differential regulation. Such features constitute adjoining regulatory domains, appropriate recognition sites for such domains and for regulatory subunits as well as specific phosphorylation sites.

The active site and, in particular, the activation segment, frame structural hot spots at which regulatory events converge (Nolen et al., 2004). Subtle structural changes affecting these regions of a PKD may be a prerequisite for productive substrate peptide and/or ATP binding. The activation of protein kinases is therefore accompanied by dynamic events which alter the conformation of structural elements reaching from individual amino acids to
structural domains. The activation segment, for example, is highly flexible and often subjected to pronounced structural changes as a result of phosphorylation events (Nolen et al., 2004). In addition, regulatory events may influence the orientation of the N- and C-terminal lobes of the PKD relative to each other and, in particular, the conformation of the regulatory helix αC. These rearrangements affect the chemical environment of the ATP binding site and, in turn, alter its accessibility and the catalytic activity (Huse and Kuriyan, 2002). Such mechanisms are pursued, for example, by the binding of cyclins to helix αC at the N-lobe of CDKs (for CDKs this helix is also called PSTAIRE helix). The structure of cyclinA bound to CDK2 revealed that cyclinA binding induces a combination of rotating and translating movements of the αC-helix (Jeffrey et al., 1995). This way, the binding displaces certain backbone atoms by as much as 8.5Å when compared to the cyclinA free structure of CDK2 (Jeffrey et al., 1995). As a result of this movement, active site residues are positioned in a productive configuration and the kinase adopts an active conformation. A further principle of PK regulation is implemented by autoinhibiting domains located outside the core of the PKD. Autoinhibiting domains can obstruct PK function by acting as substrate mimics which occupy the substrate binding site as observed with Twitchin (Kobe et al., 1996), or by distorting the productive configuration of active site elements as observed with CaMKI (Goldberg et al., 1996). The juxtamembrane (JM) domain constitutes a further regulatory element which is found in a subset of receptor protein tyrosine kinases (RTK). The JM domain links the transmembrane helix to the catalytic PKD and functions as an autoinhibitory domain in members of the platelet-derived growth factor receptor (PDGFR) family of RTKs such as c-KIT and FLT3 (Arkin and Wells, 2004; Griffith et al., 2004; Hubbard, 2004). The release of the JM domain-mediated autoinhibition in c-KIT and FLT3 is achieved through phosphorylation of conserved tyrosine residue located within the JM domain (Hubbard, 2004).

The structures of Mnk subfamily members reveal a novel autoinhibitory mechanism of PK regulation (Chapter II). But Mnks do not contain autoinhibitory regions outside the core of the PKD such as a JM domain (Parra-Palau et al., 2003; Scheper et al., 2003). Hence, Mnks employ elements to achieve autoinhibition which are integral parts of the PKD. However, to stay inactive, Mnks employ a ‘molecular trick’ which strikingly similar to the JM domain mediated autoinhibition of PDGFRs. Both classes of PKs obstruct ATP binding by sterically interfering with the productive DFG/D-in conformation and induce the inhibitory DFG/D-out
conformation. This rearrangement is achieved by inserting hydrophobic amino acid residues into a structural region referred to as DFG/D-in pocket. The origin of these inserted residues, however, differs between PDGFRs and Mnks. In contrast to c-KIT and FLT3 where inhibiting amino acid residues emanate form the JM domain, the Mnks make use of a specific Phe from the αEF/αF loop of the extended and structurally rearranged activation segment. Hence, PDGFRs and Mnks employ an analogous strategy to prevent constitutive PK activity. Albeit, the structural elements used for this purpose are of different structural origin.

The occupation of a novel binding site by the activation segment of Mnk1 constitutes the prerequisite for the induction of DFG/D-out. This repositioning of the activation segment has further structural consequences: By binding to residues from helix αC that otherwise interact with phosphor residues the helix αC and the remainder N-terminal lobe is reoriented and the interlobal cleft is closed (see Chapter II for details). Both series of structural changes, i.e. the induction of DFG/D-out and the N-lobe reorientation, crosstalk and enhance each other by intramolecular contacts at the ATP binding site. Notably, the structural data derived from both, the Mnk2 and the Mnk2\(^{D228G}\) active site mutant, display a partial, but yet incomplete, release from autoinhibition (Chapter I). Due to the high degree of conservation within their primary sequences it is likely that Mnk1 and Mnk2 behave similarly on the molecular level. Hence, I assume that the different conformational states as seen for Mnk1 and Mnk2 reflect alternative states which can be adopted by each one of the two isoforms. The combination of the results enables to depict the mechanistic events that accompany activation of Mnk subfamily members. The resulting model illustrates a novel mechanism of protein kinase regulation and expands the functional roles of the activation segment and in particular of the helix αEF and the αEF/αF loop. The Mnk structures also highlight the significance of the DFG/D motif for PK regulation (see discussion of Chapter II for details of the model). The study on Mnk regulation is therefore relevant for mechanistic and regulatory aspects of PKs in general (Huse and Kuriyan, 2002; Nolen et al., 2004).
‘The first protein structures revealed wonderful new faces of nature, but they did not help to cure anyone’. This statement, made by Max Perutz in 1992, illustrates that structural biology is, at least in part, driven by the desire to aid medical research (Perutz, 1992). The availability of an overwhelming amount of structural data and technology advancements on several levels may now indeed ‘help to cure’ since structure-guided drug design has turned into a commonly recognized discipline which is widely applied by the pharmaceutical industry (Nienaber et al., 2000; Williams et al., 2005). The authors of one of the first protein/drug co-structure, which is the bacterial enzyme dihydrofolate reductase in complex with methotrexate, set the trend by saying: ‘Perhaps it is worth noting here that the availability of the kind of detailed structural information ... provides an opportunity for rational drug design ... ’ (Matthews et al., 1977). This prediction had soon been met. Among the first examples of successful applications of structure-guided drug design studies count the development of a potent agent against the influenza virus (von Itzstein et al., 1993) and inhibitors for the HIV protease (Ridky and Leis, 1995). Likewise, the proteins studied in the course of this PhD-work constitute potential drug targets or are related to pharmaceutically relevant proteins, respectively. Inasmuch, their crystal structures may provide valuable templates for structure guided drug design efforts. The availability of inhibitors against the studied proteins could, in addition to their medical applications, also potentiate basic functional understanding on the organismic and the tissue culture level.

PKs are implicated in many severe diseases. However, they have initially been considered to be ‘not drugable’ because of the strong conservation of their ATP binding site and the high intracellular ATP concentration (Cohen, 2002). Nevertheless, the first selective PK inhibitor was launched in 2001 (O'Dwyer and Druker, 2000). Subsequently, PKs turned into a major target of drug development programs (Noble et al., 2004). As a result, several additional PK inhibitors were launched and the pipelines of the pharmaceutical industries are filled with promising candidates (Vieth et al., 2005). Currently, protein kinases constitute the most important drug target in conjunction with the G-protein coupled receptors and investigators make tremendous use of structural data when designing and optimizing small molecule inhibitors (Cohen et al., 2005; Gill et al., 2005). The majority of small-molecule PK inhibitors achieve inactivation through direct binding to the ATP site. However, the desire for
selectivity encouraged the search for alternative inhibitory strategies. Consequently, novel agents have been found comprising monoclonal antibodies and small molecules which recognize binding sites remote from the ATP binding pocket (O’Loghlen et al., 2004; Vieth et al., 2005).

The crystal structures of Mnk1 (Chapter II) and Mnk2 (Chapter I) suggest novel strategies of PK inhibitor development (Vieth et al., 2004). The autoinhibited conformation of Mnk1 and Mnk2 as well as the presence of Mnk-specific residues at the active site, suggest distinct target sites for specific Mnk inhibition and for PK inhibition in general. This conclusion is based on two major observations. Firstly, the autoinhibitory DFG/D-out conformation of the magnesium binding loop resembles a conformation observed in the MAPK p38 which was induced by the inhibitor BIRB-796 (Pargellis et al., 2002). The mode of inhibition seen in the p38/BIRB-796 complex indicates the presence of a novel allosteric binding site which causes indirect inhibition by stabilizing DFG/D-out (Pargellis et al., 2002). This allosteric binding site is structurally less conserved than the actual ATP binding site and may therefore result in an improved selectivity of the corresponding inhibitors (Mol et al., 2004; Noble et al., 2004). Since Mnk1 and Mnk2 adopt DFG/D-out passively, they appear well suited for allosteric inhibition. This assumption is supported by the recent observation that the allosteric p38 inhibitor BIRB-769 binds to Mnk2 in vitro (Fabian et al., 2005). Furthermore, the occurrence of DFD instead of the canonical DFG assigns the chemical distinctiveness to the relevant binding site and could benefit selectivity. A second strategy of PK inhibition can be deduced from the induction of autoinhibition through the structurally rearranged activation segment as seen in the Mnk1 structure. As described above, the occupation of the DFG/D-in pocket through a hydrophobic residue induces DFG/D-out and thereby switches the PK off. Hence, the development of agents which target the DFG/D-in pocket may represent a novel strategy for the design and screening for protein kinase inhibitors. Peptidomimetics resembling the autoinhibitory elements encoded by the αEF helix and the αEF-αF loop region may represent a suitable starting point for such endeavors.

The second type of enzyme under study, the ecoNADS, is likewise of relevance for structure guided drug design. With the availability of structural data for the ecoNADS phylogenetic structure comparisons with the corresponding enzyme of pathogenic bacteria are made possible. Namely, the NADS of Bacillus anthracis is targeted for the development of agents combating anthrax (Sutherland, 2003). Therefore, the availability of the crystal
structure of various forms of the *Escherichia coli* NADS provided in *Supplementary Chapter III* and its detailed comparison to previously investigated *Bacillus subtilis* NADS structures can help to optimize inhibitory compounds by potentiating the selectivity (*Supplementary Chapter III*).

Traditionally, the small-molecule binding sites of enzymes and receptors are targeted by drug design efforts (Cochran, 2000). However, protein-protein recognition surfaces are often chemically more distinct and, thus, receive an increasing attention in the course of drug design efforts (Arkin and Wells, 2004; Sillerud and Larson, 2005). The insect-specific zinc-finger associated domain (ZAD) represents such a potentially drugable protein-protein recognition site (*Supplementary Chapter III*). The ZAD exhibits a unique fold and amino acid substitutions within the ZAD of certain genes result in lethality (Crozatier et al., 1992; Gaszner et al., 1999). Furthermore, the analysis of the genomic sequence of humans and other vertebrates revealed that the ZAD is missing in the respective genomes (Chung et al., 2002). This finding is consistent with the assumption that drugs specifically targeting the ZAD would not be harmful to vertebrates including humans. Hence, the ZAD constitutes a molecular scaffold which could potentially be used to develop insecticides. One of the potential targets of such compounds could be the transmitter of malaria *Anopheles gambiae* (*Supplementary Chapter IV*).
Bibliography


## Abbreviations

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<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>ATP</td>
<td>adenosine tri-phosphate</td>
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<tr>
<td>CAMK</td>
<td>Ca$^{2+}$ - calmodulin modulated protein kinase</td>
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<tr>
<td>CDK</td>
<td>cyclin dependent kinase</td>
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<tr>
<td>C-loop</td>
<td>catalytic loop</td>
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<td>DAPK</td>
<td>death associated protein kinase</td>
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<td>DFG</td>
<td>Asp-Phe-Gly tripeptide motif</td>
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<td>eIF</td>
<td>eukaryotic initiation factor</td>
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<tr>
<td>ERK</td>
<td>extracellular signal regulated kinase</td>
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<tr>
<td>hnRNPA1</td>
<td>heterogenous nuclear ribonucleoprotein A1</td>
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<td>IRK</td>
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<td>JM</td>
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<td>MAPK</td>
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<td>MAPKAP</td>
<td>MAPK activated protein kinase</td>
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<td>Mnk</td>
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<tr>
<td>mRNA</td>
<td>messenger RNA</td>
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<td>NAAD</td>
<td>nicotinic acid adenosine dinucleotide</td>
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<tr>
<td>NAD</td>
<td>nicotinamide adenosine dinucleotide</td>
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<td>NADS</td>
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<td>PDGFR</td>
<td>platelet-derived growth factor receptor</td>
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<td>PK</td>
<td>protein kinase</td>
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<td>PKD</td>
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<tr>
<td>RTK</td>
<td>receptor tyrosine kinase</td>
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<td>TNF-α</td>
<td>tumor necrosis factor alpha</td>
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<td>tRNA</td>
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<td>VEGF-R</td>
<td>vascular endothelial growth factor receptor</td>
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<td>ZAD</td>
<td>zinc finger associated domain</td>
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## Abbreviations - Amino Acids

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Abbreviations - Amino Acids
Acknowledgements

I’d like to express my gratitude to Prof. Dr. Herbert Jäckle for his inspiring advice, continuous interest in my work and seemingly infinite support. I particularly appreciated that he gave me the freedom to develop my personal scientific interest without restrictions. I would also like to thank all members of the ‘Abteilung für Molekulare Entwicklungsbiologie’ for helpful advice, discussions and reagents. In this respect I’m particularly grateful to Dr. Ralf Pflanz. None of my projects would have been possible without the close collaboration with the ‘Arbeitsgruppe Röntgenkristallographie’ led by Dr. Markus Wahl. I enjoyed a tremendous support provided by Markus in most aspects of my biochemical and crystallographic endeavors. I also thank the other members of his group for advice, help with reagents and for tolerating me as a ‘contaminant’ in their lab. I’m particularly grateful to Catharina Netter who helped me with the initial crystallographic work and by setting up the crystallization facility. In addition, she was always available for critically assessing my manuscripts and ideas. I also thank Dr. Gleb Bourenkov (DESY) and Dr. Ehmke Pohl (SLS) for excellent beamline maintenance, help and discussions.

I thank Prof. Dr. Reinhard Jahn and Prof. Dr. Thomas R. Schneider as members of my PhD committee for their support, helpful comments and critical evaluation of my work. The efficient operation of the coordination office of the ‘MolBio’-program led by Dr. Steffen Burkhardt facilitated numerous aspects of the ‘PhD life’.

The NADS project would not have been impossible without the initial work performed by Dr. Andreas Humm and his colleagues. I’m very grateful that I have been allowed to use his data. I thank the Develogen AG and in particular Dr. Stefan Jäkel, Dr. Babette Aicher and Dr. Cord Dohrmann for providing purified Mnk proteins. It’s been a very fruitful and congenial collaboration. For numerous instructive discussions particularly during the lunch breaks I thank Ben Harder, Patrick Müller and Ajaybabu V. Pobbati. I’m grateful to Dr. Jörg Kahle and Anja Uhmann for proofreading this manuscript and many aspects beyond the scientific life.
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Patents
