Mip protein of Legionella pneumophila exhibits peptidyl-prolyl-cis/trans isomerase (PPlase) activity

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Summary

Legionella pneumophila is an intracellular parasite which is able to survive and multiply in human monocytes and alveolar macrophages. The Mlp (macrophage infectivity potentiator) protein has been shown to be an essential virulence factor. A search of translated nucleic acid data bases has shown that the Mip protein from strain Wadsworth possesses regions homologous to those found in the FK506-binding proteins (FKBPs) of several different eukaryotic organ-Isms. FKBPs are able to bind to the immunosuppressant macrolide FK506 and possess peptidyl-prolyl cis/trans isomerase (PPlase) activity. The gene coding for the Mip protein was cloned from the chromosome of L. pneumophila strain Philadelphia I and sequenced. It was synthesized in Escherichia coll K-12 and after purification it exhibited PPlase activity catalysing the slow cisitrans isomerization of prolyl peptide bonds in oligopeptides. Mip is inhibited by FK506 and fully resistant to cyclosporin A, as was also found for the recently characterized FKBP-type PPlases of eukaryotes. However, the N-terminal extension of Mip and/or the substitutions of the variable amino acids in the C-terminal FKBP core lead to variations, when compared with eukaryotic FKBPs, in substrate specificity with the oligopeptide substrates of type Suc-Ala-Xaa-Pro-Phe-4-nitroanilide. Nevertheless, the Legionelia Mip factor represents a bacterial gene product which shares some characteristics normally found in eukaryotic proteins. In view of the activity of PPlases in protein-folding reactions, such prokaryotic FKBP analogues may represent a new class of bacterial pathogenicity factors.

Introduction

Legionella pneumophila is the aetiological agent of Legionnaires' disease, a distinct form of pneumoniae in humans (McDade et al., 1977; Winn, 1988; Horwitz, 1988). The bacterium multiplies intracellularly in human blood monocytes, human alveolar macrophages and tissue-culture cells. L. pneumophila is phagocytosed by an unusual coiling phagocytosis mechanism (Horwitz, 1984) and is able to inhibit phagosome—lysosome fusion and phagosome acidification of the phagocytes (Horwitz, 1983; Horwitz and Maxfield, 1984; Horwitz, 1988).

In order to identify the gene products involved in the pathogenesis of L. pneumophila (for a review, see Engleberg and Eisenstein, 1991) the determinants coding for two toxins and for three different protein antigens have been cloned and sequenced (Black et al., 1990; Wintermeyer et al., 1992; in preparation; Hoffman et al., 1990; Engleberg et al., 1989; Ludwig et al., 1991). To date, only the Mip (macrophage infectivity potentiator) protein, a polypeptide of 24 kDa, is known to contribute to the virulence of L. pneumophila (Cianciotto et al., 1989; 1990b). Using isogenic Legionella strains with and without an intact mip gene it was shown that Mip potentiates the uptake of L. pneumophila by macrophages and/or contributes to early survival processes in the phagocytic cell. The amino acid sequence of the C-terminal part of the Mip protein was deduced from the DNA sequence of the mip gene from strain Wadsworth (Engleberg et al., 1989) and revealed homology to the sequences of FK506-binding proteins (FKBPs) found in various eukaryotic organisms (Standaert et al., 1990; Maki et al., 1990; Tropschug et al., 1990; Wiederrecht et al., 1991; Jin et al., 1991; Lane et al., 1991). The eukaryotic FKBPs are able to bind to the immunosuppressant macrolide, FK506. FKBPs and cyclophilins form the substance class of the immunophilins (Schreiber, 1991). The latter have the capacity to bind cyclosporin A (Fischer et al., 1989; Price et al., 1991). Immunophilins possess peptidyl-prolyl cis/trans isomerase (PPlase) activity and catalyse the slow isomerization of prolyl bonds in oligopeptides and proteins in

Received 22 November, 1991; revised and accepted 10 February, 1992. *For correspondence. Tel. (931) 31378/31575; Fax (931) 571954.

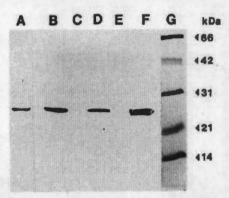


Fig. 1. Western blot analysis of cell extracts and Mip protein using antiserum prepared against the Mip-positive *E. coli* K-12 clone HB101 (pBLL3039). Lanes: A, cell extract of *L. pneumophila* Philadelphia I; B, cell extract of *E. coli* K-12 (pBLL3039); C, cell extract of *E. coli* K-12 (pLAFR2) (control); D, cell extract of *E. coli* K-12 (pBLL106); E, cell extract of *E. coli* K-12 (pBR322) (control); F, purified Mip protein of *E. coli* K-12 (pBLL106); G, molecular weight markers (BioRad Laboratories).

vitro (Lang et al., 1987; Fischer and Schmid, 1990; Tropschug et al., 1989). Recently, it has been suggested that the timing of the triple-helix formation of procollagen I in chick-embryo tendon fibroblasts is related to PPlase activity in vivo (Steinman et al., 1991). In addition, Immunophilins and their complexes with FK506 or cyclosporin A seem to play a key role in the signal-transduction pathways of T cells (Schreiber, 1991).

In order to test whether both PPlase activity and FK506 sensitivity are conserved within the *Legionella* Mip protein, we have cloned and sequenced the corresponding

gene from strain *L. pneumophila* Philadelphia I and have purified the protein. It is shown that Mip has PPlase activity which approximates the same level as that found in the eukaryotic FKBPs. The PPlase activity of Mip is strongly inhibited by 100 nM FK506 but is completely resistant to micromolar concentrations of cyclosporin A.

Results

Cloning and expression in Escherichia coli K-12 of the mip gene from L. pneumophila strain Philadelphia I

A genomic library of L. pneumophila Philadelphia I (Hacker et al., 1991; Ludwig et al., 1991) was screened by Immuno-colony blotting using anti-Legionella-specific antiserum. Seventy-six clones reacted strongly and the cell lysates were further analysed by SDS-polyacrylamide gel electrophoresis (PAGE) and Western blotting. Proteins of 24 kDa were produced by 21 recombinant clones. DNA isolated from each of these clones was found to react in a DNA-DNA dot blot using a 30-mer oligonucleotide probe with a mip-specific sequence (Engleberg et al., 1989). A BamHI-Clai DNA fragment of 4.5 kb was subcloned from one of the cosmid clones, HB101(pBLL3039), into the vector pBR322. Western blots with a polyclonal monospecific anti-Mip-antiserum showed that the resulting clone, HB101(pBLL106), produced the Mip protein (Fig. 1).

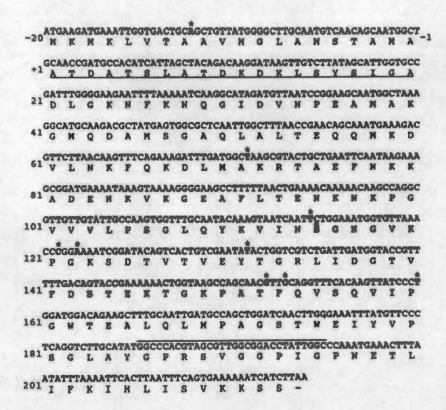


Fig. 2. Nucleotide and protein sequence of the mip gene from strain L. pneumophila Philadelphia I. The nine nucleotides and the amino acid at position 115 which differ from the published sequence of strain Wadsworth are indicated in bold type and by asterisks. The N-terminal amino acid sequence underlined was determined by Edman degradation. The overlined nucleotides represent the oligonucleotide sequence used for DNA-DNA hybridization. These sequence data will appear in the EMBL/GenBank/DDBJ Nucleotide Sequence Data Libraries.

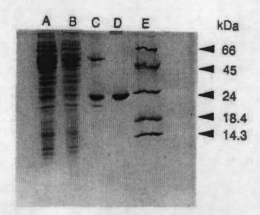


Fig. 3. Purification of recombinant Mip protein. The various fractions were subjected to SDS-PAGE and protein was stained with Coomassie brilliant blue. Lanes: A, crude cell extract of subclone HB101(pBLL106); B, Mip containing fractions after DEAE-sephacei passage; C, Affigel-blue chromatography; D, combined phenyl-sepharose fractions; F, molecular weight markers (Sigma).

DNA sequence of the mip determinant of strain L. pneumophila Philadelphia I

The DNA sequence of the *mip* gene from strain Philadelphia I was determined (Fig. 2). This sequence revealed nine nucleotide substitutions relative to the *mip* sequence of strain Wadsworth (Engleberg et al., 1989). One of these substitutions resulted in the replacement of an alanine at amino acid position 115 with serine. The other eight nucleotide substitutions did not lead to alterations of the Mip amino acid sequence.

Isolation of the recombinant Mip protein

As shown in Fig. 3 and lane F of Fig. 1, the Mip protein of approximately 24 kDa produced by clone HB101 (pBLL106) was isolated and obtained in a purified form using a four-step procedure. Table 1 summarizes the steps and yields as well as the enrichment factors thus obtained. No contaminating cyclophilin-like E. coli PPI-ases could be detected in the final Mip preparation.

Table 1. Purification steps leading to homogeneous Mip-PPlase.

Purification step	Total protein (mg)	Total Mip activity (arbitrary units)	Specific activity (arbitrary units per mg protein)	Recovery (%)	
DEAE-Sephecel*	275	4152	15	100	
TSK Affigel Blue	4.7	1020	215	30	
Phenyl-Sepharose	0.61	346	568	12	

a. Mip activity measured toward Suc-Ala-Pro-Phe-4-nitroanilide; differentiation of Mip enzymatic activity from E. coli PPlase is only possible after the DEAE-Sephacel step

N-terminal protein sequence of Mip

In order to demonstrate that the isolated 24 kDa polypeptide was indeed Mip, N-terminal sequencing of the mature protein was undertaken. As indicated in Fig. 2, the initial amino acids of the mature protein are alanine, threonine, and aspartate. This clearly demonstrates a cleavage of the first 20 amino acids following transport. Each of the 20 N-terminal amino acid residues of the mature Mip protein are identical to those predicted from the DNA sequence of mip indicated in Fig. 2.

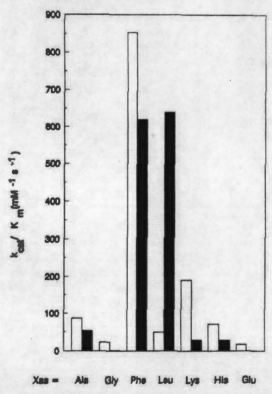


Fig 4. Comparison of the specificity constants $k_{\rm cat}/k_{\rm m}$ of Mip PPtase (40–293 nM) activity (open columns) and human FKBP (shadowed columns, data from Harrison and Stein, 1990) toward Suc-Ala-Xaa-Pro-Phe-4-nitroanilide (pH 7.8; 0.035 M⁻¹ HEPES buffer) at 10.0°C.

Determination of PPlase activity of Mip

The Mip protein was tested for PPlase activity upon Suc-Ala-Xaa-Pro-Phe-4-nitroanilide with the aid of isomer-specific proteolysis using chymotrypsin as the protease. The particular conditions of the assay result in $k_{\rm cut}/k_{\rm m}$ for the *cis/trans* isomerization of the -Xaa-Pro bond.

We found enzymatic activity which slightly exceeds the highest value of k_{cet}/K_m observed for eukaryotic FKBPs (Fig. 4) using the most favoured substrate, Suc-Ala-Phe-Pro-Phe-4-nitroanilide. Mip additionally shows an elevated activity towards a substrate with a Lys in the P₁ position (using the nomenclatural system proposed by Schechter and Berger (1967) for the reactive peptide

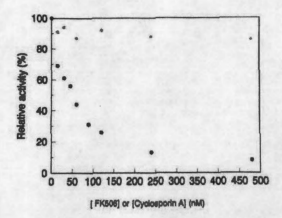


Fig. 5. PPlase activity as a function of the cyclosporin A (*) and of the FK508 (o) concentration.

bond of oligopeptide substrates). The characteristic 5–10-fold drop in activity for the entire set of substrate peptides that was found in prokaryotic and eukaryotic cyclophilin-like PPlases (G. Fischer, H. Bang, M. A. Marahiel, R. Schönbrunner and F. X. Schmld, submitted for publication) is not observed with FKBPs.

More interestingly, MIp lacks the pronounced Leu specificity for the P_1 position demonstrated for human FKBP (Harrison and Stein, 1990). A 600-fold decrease of $k_{\rm out}/K_{\rm m}$ was determined for human FKBP in Xaa=Gly cis/trans isomerization. This value, like that obtained using the human FKBP optimal substrate, was significantly reduced in Mip. This suggests that the hydrophobicity of the amino acid side chains cannot satisfactorily explain the influence of the P_1 -amino acid on catalysis in Mip. The control strain HB101(pBR322) does not exhibit any PPlase activity with a specificity pattern similar to the Mip enzyme.

Inhibition studies with FK506 and cyclosporin A

As it was shown that all FKBPs possessing PPlase activity could be inhibited with nanomolar amounts of FK506 (Tropschug et al., 1990; Siekierka et al., 1990), we decided to examine the effect of increasing concentrations of FK506 as well as those of cyclosporin A, the inhibitor of cyclophilin-like PPlases, upon the enzymatic activity of Mip (Fig. 5). PPlase activity could be considerably influenced by FK506 although it was not significantly inhibited by cyclosporin A or by the non-immunosuppressive cyclosporin H. The IC₅₀ value (50% inhibitory concentration) calculated from the data of Fig. 5 is about 45 nM, indicating that the K₁ value may be even smaller.

Discussion

The amino acid sequence of the Mip protein, a virulence

factor of *L. pneumophila* (Cianciotto *et al.*, 1989), the aetiological agent of Legionnaires' disease in humans, shows
homology to amino acid sequences of human, *Neurospora* and yeast FK506-binding proteins (Standaert *et al.*, 1990; Maki *et al.*, 1990; Tropschug *et al.*, 1990;
Wiederrecht *et al.*, 1991; Jin *et al.* 1991; Lane *et al.*, 1991;
see Fig. 6). Cyclophilins and eukaryotic FKBPs represent
a family of peptidyl-prolyl *cis/trans* isomerases (PPlases)
termed immunophilins which can catalyse slow *cis/trans*interconversion of prolyl imidic peptide bonds in oligopeptides and proteins (Lang *et al.*, 1987; Fischer *et al.*, 1989;
Fischer and Schmid, 1990; Tropschug *et al.*, 1989; Price *et al.*, 1991; Schreiber, 1991).

When we tested the recombinant Mip protein cloned from L. pneumophila strain Philadelphia I we found PPIase activity comparable in magnitude to other cyclophilintype prokaryotic PPlases and to eukaryotic FKBPs. The comparison of the P, subsite specificity of the recombinant Mip with human FKBP (Fig. 4) results in a slightly different pattern of k_{cat}/K_m. The highest catalytic constant that could be obtained for the peptide with Xaa=Phe (850 000 M-1 s-1) exhibits an 18-fold lower activity when compared with the highest activity yet measured for a PPlase (pig kidney 17.8 kDa cyclophilin acting on Suo-Ala-Ser-Pro-Phe-4-nitroanilide). The clear dependence of koat/Km on hydrophobicity of the P1 amino acid side chain using human FKBP (Harrison and Stein, 1990) is not valid in Mip catalysis as even charged side chains can fulfil the requirements for favourable catalytic interactions.

The inhibitory effect of FK506 on Mip is similar of that observed for human FKBP although the interaction may be slightly weaker. However, the large PPlase concentration of 47 nM used in our inhibition assay does not allow us to determine a K_i range much smaller than this value. As in the case of P_1 subsite specifity, this may result from either the N-terminal extension of Mip or the variable amino acids located in the FKBP analogue C-terminal part (Fig. 6).

As indicated in Fig. 6, unidentified open reading frames (URFs) in the genome of the bacterial pathogens Neisseria meningitidis (Perry et al., 1987) and Pseudomonas aeruginosa (Kato et al., 1989) resembling FKBPs are located in close proximity to virulence-associated loci, near two silent copies of truncated pilin genes in N. meningitidis and a region encoding regulatory proteins involved in the alginate biosynthesis of P. aeruginosa. Furthermore, Mip of L. pneumophila is closely related to a homologous protein of Legionella micdadei (Bangsborg et al., 1991) and to the L2 protein of Chlamydia trachomatis (Lundemose et al., 1991). Both are pathogenic organisms that exhibit virulence features similiar to those of L. pneumophila (Moulder, 1985; Horwitz, 1988). The possibility that putative gene products of the URFs and of

L.pneumophila phill Mip	AT	DATSLAT	DEDELSTRIG	ADLGENTENQ	GIDVNPEAKA	KGMQDAMSG-	AQLALTEQQN	KDALWKLÖKD
Human (cy) FKBP		*********		********				********
Human (mem) FKBP			********	********	********	********	********	
Bovine FKBP							********	
S.cerevisiae FKBP							********	
N.crassa FKBP	********		*********	********	********	********	********	********
N.meningitidis URF	********	********	********	********	********	********		********
C. trachomatis L2protein					LVEV	IKCHOSEIDG	QSAPLTDTE.	YEKQ
L. micdadei Mip	ATATTDAT	TSAPCTSLITT	DTEKLBYSIG	ADLGKNIKKO	GIEISPAAMA	KGLODGM8G-	GOLLLITDDON	KDALMKLÖKD
P.aeruginosa URF		AP	PEDELAYAVG	ARLGTRLQQE	MPDLELSELL	LGLRQAYRG-	EALEIPPERI	EQLLLQHE
Consensus sequence	AT	*TSL	*.L.Y.*G	A.IG * *			*.*	**FNK***
L. pneumophila phill Mip	- 41 4500 000	PPS DEWELLEA		DAWN/I BEAT.	-AVENTNEAN	GUTTATED	PUPURYNADI.	TOGTVFDATE
n' buadmoburre burre wib	THUNKINDEN	*********	DES DI BUNETA	TALLINEDAN	Asuasunan	4141410		
Human (cy) FKBP			No.	MAN	-OVETTS POD	GRTPDERGO-	TCVVHYTGHL	EDGEKTDESR
Human (mem) PKBP	********	********	**********	TCARGEDY	TOTCOKKDOD	HCDINGDACH	VLHMHYTOKL	EDGTETDESL
Bovine FKBP				May	-OVETTSPED	GRTFPERCO-	TCVVHYTGHL	EDGKKFDESR
S. cerevisiae FKBP		********		MSEVTRON	VKIDRISPAD	GATEPETCD-	LVTIHITGTL	ENGOKEDESV
N. crassa FKBP	********			MTT POT DOT	-OTEVOORGO	GTRETERGD-	NADAHAKGAF	TSCKKTDASY
N.meningitidis URF				Mag	LITEDIOESE	GERAVECE	EITVHYTGWL	EDGTKFD8SI.
C. trachomatis Liprotein	MARUONACAR	APCOMULACE.	ERPI.KEWERE	AGUTET. PONE	THUBANKEGT	GRVI.SCEP	TALLHYTGSF	IDGKVFDSSE
L.micdadei Nip							VYTVEYTOKL	
P. aeruginosa URF	MARKADARIA	DATABASAS	WIANDYAD	ECVPRIM	CCULACATOD	COCHETCAAT	QVHVRTRGLL	ADGOVEDOSE
Consensus sequence	KATIBIPKII	N +	P PT. SSY S	FGANGEL	GGIDIDELEGA	aKauarawar	Y.G	GFD
Consensus sequence		********	DIFM. "-K."					
L.pneumophila phill Mip	KTGKPATPQV	SQVIPGWI	BALQLHPAGS	TWEITVPSGL	AYGPREVGGP	IGPHETLIFE	IHLISVEKSS	
Human (cy) FKBP	DENEDEKTMI.	GEORVIRGUE	RGVAONSUGO	BAKI/TIRPDY	AYGATCHPGI	IPPHATLUED	VELLKLE	
Ruman (mem) FKBP	PONOPPVPSI	GTGOVIKOWD	OGLIGHERGE	KRKI.VI PARL	GYGERGAPPK	IPCGATLVFE	VELLKIERRT	BL
Bovine FKBP	DENEBEKEUT	GKOEVTROWE	BGVAOMSVGO	PAKITTEPDY	AYGATGHPGI	IPPNATLIFO	VELLKLE	
S. cerevisiae FKBP	DEGSPECCHT	GUGOVIKOWD	VGTPKTAVGE	KARLTIDGDY	AYGPEGFPGL	IPPMSTLVED	VELLKAN	
N. crassa FKBP	DEGERINATA	COCONTROWD	MCLICHKTOR	KPKI/TIAPHI	AYGNRAVGGI	IPANSTLIFE	TELVGIEGVO	KGE
N.meningitidis URF	DEBORITTE	GVGOVIKGWD	BGFGGIOCEGG	KRKI/TI PSEM	GYGATRRGGV	IPPHATLIFE	VELLKAYE	*******
C. trachomatis L2protein	ENKEPTI.T.DI	TKVIPOFS	OGMOGNERGE	VRVLYIHPDL	AYGTAGO	LPPMSLLIFE	VKLIEANDON	VSVTE
L.micdadei Hip	KTGKPATPKV	BOVI POWT	BALOLMPAGE	TWEVYIDANI	AXGPREVGGP	IGPNETLIFE	IHLISYKKAN	A
P.aeruginosa URF								
Consensus sequence								*******
Advingtions and anima				V 50 00 00 000 000	A DESCRIPTION		The second secon	The state of the s

Fig. 6. Comparison of the amino acid sequences of FKBPs exhibiting PPlase activity (Neurospora crassa, Saccharomyces cerevisiae, human Jurkat T cells [Cy FKBP], human cancer cell line [mem FKBP], and bovine calf thymus cells), and amino acid sequences of the mature Mip-like protein of L. micdadei, the C-terminal part of the L2 protein of Chlamydia trachomatis, parts of unidentified open reading frames (URFs) of Neisseria meningitidis and Pseudomonas aeruginosa and the mature L. pneumophila (L.p.) Mip protein of strain Philadelphia I. Identical amino acid residues are indicated in bold type. Dashes represent gaps for optimal alignment. Isofunctional replacements are indicated by asterisks. Abbreviations: cy, cytoplasmic; mem, membrane-bound.

the L. micdadei mip as well as of the L2 locus also influence the pathogenicity of the respective organisms cannot be excluded.

It has been shown recently that a genetically engineered mip mutant of L. pneumophila strain Wadsworth was defective in its ability to initiate macrophage infection (Cianciotto et al., 1989; 1990b). Cloning of the mip locus from another pathogenic Legionella isolate. Philadelphia I (Fig. 2), confirms the view that mip is highly conserved among all Legionella strains (Cianciotto et al., 1990a). This suggests a common function for the protein in different isolates. The role of Mip enzymatic activity in Legionella virulence has yet to be elucidated. It is possible that the Mip protein modifies bacterial cell structures or surface proteins of the phagocytic cell, thereby enhancing their phagocytic capacity. A further possibility is that the Mip protein activity in the phagocytic cell could modify proteins to inhibit phagosome-lysosome fusion and acidification of the vacuole (Horwitz, 1983; Horwitz and Maxfield, 1984). Alternatively, a deregulation of signal-pathway proteins in phagocytes cannot be ruled out.

A large number of extracellular substances with enzymatic activities such as phospholipases, lecitinases, proteases or sugar transferases, some of which are membrane-associated, have been shown to contribute to the pathogenicity of bacteria (see Finlay and Falkow, 1989). The Legionella Mip protein, exhibiting PPlase activity reminiscent of that found in FKBPs, is an example of a bacterial virulence factor possessing characteristics normally observed in eukaryotes. A further example of a protein in pathogenic bacteria with enzymatic capacity normally ascribed to eukaryotes was recently detected in Yersinia. This genus comprises the species Yersinia pestis, the causative agent of plague. The YopH protein of Yersinia exhibits tyrosine phosphatase activity which is able to influence the phosphorylation pattern of the eukaryotic cell (Guan and Dixon, 1990; Bliska et al., 1991). Like Legionella, Yersinia is also able to replicate intracellularly. One may speculate that prokaryotic proteins with enzymatic functions related to eukaryotic cell signal factors represent a new general class of gene products which contribute to survival and dissemination of intracellular bacterial pathogens.

Experimental procedures

Media, enzymes and chemicals

Legionella strains were cultivated on buffered charcoal-yeast

extract (BCYE) agar plates at 37°C in a 5% CO₂ atmosphere for 48 h prior to use. *E. coli* K-12 strains were grown in L-broth. Radiochemicals were purchased from NEN Research Products. Restriction enzymes, T4 polynucleotide kinase and T4 ligase were obtained from Pharmacia, Boehringer, Biolabs, and Gibco BRL. All other chemicals were obtained from Merck, Difco, BRL, Oxoid, Roth, and Serva. Antibiotics were from Serva or Sigma.

Bacterial strains and plasmids

The genomic library was constructed from *L. pneumophila* strain Philadelphia I (McDade *et al.*, 1977). For cloning experiments, the *E. coli* K-12 strain HB101 was used. Plasmid pLAFR2 is a cosmid vector described previously (Knapp and Mekalanos, 1988). For subcloning, vector pBR322 (Bolivar *et al.*, 1977) was used. Antibiotics were added at concentrations of 50 µg ml⁻¹ (ampicillin) and 10 µg ml⁻¹ (tetracycline).

Cosmid cloning procedure

Legionella-specific chromosomal DNA was isolated as described (Ludwig et al., 1991; Wintermeyer et al., 1991). Large (20 kb) chromosomal Sau3A fragments were ligated into the BamHI site of cosmid pLAFR2 and transduced into HB101 with the help of the cosmid packaging system (Hohn and Collins, 1980).

Recombinant DNA techniques

Plasmid DNA was isolated as described (Birnboim and Doly, 1979). For restriction enzyme analysis, DNA was treated with appropriate enzymes and the resulting fragments were separated by gel electrophoresis on 0.8–1.0% agarose gels (Sambrook et al., 1989). DNA fragments were isolated from the agarose gel with the help of the GeneClean kit from Bio101 as described by a protocol of the manufacturer. For cloning, DNA fragments were ligated into suitable vector molecules after heat-inactivation of the restriction endonucleases at 65°C for 10 min (Sambrook et al., 1989). E. coli K-12 strains were transformed by the CaCl₂ method (Lederberg and Cohen, 1974).

Oligonucleotide synthesis

Oligonucleo' les used for DNA sequencing and DNA-DNA dot blotting were synthesized with an Applied Biosystems 380A DNA synthesizer using the phosphoramidite method of Beaucage and Caruthers (1981). Oligonucleotides were purified with oligonucleotide purification cartridges from Applied Biosystems.

DNA sequencing

The sequence of the *mip*-specific DNA region was determined with the help of a sequencing kit from Boehringer, as described by a protocol of the manufacturer.

Computer analysis

The programs used for compiling the nucleotide sequence data

were from J. Devereux (University of Wisconsin Genetics Computing Group).

DNA-DNA dot blotting

As a *mip*-specific probe the oligonucleotide 5'-GGCCCACG-TAGCGTTGGCGGACCTATTGGC-3' (see Fig. 2) was used. The probe was labelled by T4 polynucleotide kinase using $[\gamma^{32}-P]$ -ATP. The DNA dot blotting procedure and hybridization were carried out according to Sambrook *et al.* (1989).

Preparation of antisera

The antigen used for production of the *L. pneumophila*-specific antiserum was prepared from heat-killed *L. pneumophila* strains grown on agar plates as described (Hacker *et al.*, 1991). Antiserum was prepared by intravenous injection of the antigen into New Zealand rabbits. The final immunofluorescence titre (IFT) of the antiserum was 1:1024. Antiserum specific for the cloned Mip protein of *L. pneumophila* was prepared from *E. coli* K-12 clone HB101(pBLL3039). The final IFT was 1:512. To remove the *E. coli* K-12-specific antibodies the antiserum was absorbed with a suspension of HB101(pLAFR2).

Immuno-colony blotting

Immuno-colony blots were made as described by Van Die et al. (1985).

Western blotting

For Western blotting, 10 ml of overnight culture of the recombinant *E. coli* K-12 clones or an equal amount of *L. pneumophila* cells harvested from agar plates was pelleted by centrifugation at 4°C for 10 min and washed with 10 mM ice-cold HEPES buffer (pH 7.4). Cell pellets were suspended in 1 ml 10 mM HEPES buffer (pH 7.4). The suspensions were passed through a French press (3×6210 kPa) and residual cells were removed by centrifugation. Proteins were analysed by SDS-PAGE as described by Laemmli (1979). The Western blots were performed according to the method of Kyhse-Andersen (1984).

Protein sequencing

The protein was subjected to 12.5% SDS-PAGE and blotted on to PVDF membranes (Immobilion transfer; Millipore) as described by Choli et al. (1989). The Mip protein was sequenced in an Applied Biosystem 470 A gas-phase sequencer.

Isolation and purification of the Mip protein

For isolation of the Mip protein, cells from 10 I of ovemight cultures of the recombinant *E. coli* K-12 clone HB101 (pBLL106) were pelleted by centrifugation at 4°C for 10 min at 10 000 r.p.m. and washed with 10 mM ice-cold HEPES buffer (pH 7.4). Bacterial cells were suspended in 50 ml of 10 mM HEPES buffer (pH 7.4), passed through a French press (3×6210 kPa) and cell debris were removed by centrifugation for 20 min at 20 000 r.p.m. (in a Kontron T 324 centrifuge; A 8.24 rotor). Polymin P (BASF) was added to the supernatant to

give a final concentration of 0.4%. After being stirred for 30 min, the solution was centrifuged as described above. The supernatant was applied to a DEAE-Sephacel column (2.5 cm i.d. × 35 cm) equilibrated in 35 mM HEPES (pH 7.8). In contrast to E. coll PPlase, the Mip protein does not bind to this material. Enzymatically active fractions were applied to a 1.5 × 15 cm Affigel-blue column (Merck) equilibrated with 35 mM HEPES (pH 7.8). Under such conditions the Mip protein adsorbs to the affinity column. Approximately 80% of the protein could be recovered when eluted with 200 ml of 2 mM Tricin (pH 8.5).

Mip activity was eluted by applying a linear gradient (300 ml total) of 0-400 mM KCl in 2 mM Tricine (pH 8.5). Active fractions (at approx. 240 mM KCI) were pooled and KCI was added to a total concentration of 0.5 M to the pool prior to pumping the sample on to a 0.5 x 5 cm phenyl-sepharose column in 2 mM Tricine, 0.5 M KCI, (pH 8.5). The following elution was then carried out using a 50-ml step gradient, starting with 2 mM Tricine (pH 8.5), followed by addition of 0.1% Triton X-100 and ending with 3% Triton X-100. The homogeneity of Mip was verified by silver-staining after SDS-PAGE. The recovery of Mip was about 10% with respect to the Mip enzymatic activity found after separation from the E. coli PPlases. For sequencing, salts were diluted out from the protein with water using Centricon C 10 filtration (Amicon).

PPlase assay

Enzymatic activity was measured in 0.035 M HEPES buffer (pH 7.8) at 10°C using the protease-coupled assay described previously (Fischer et al., 1989). Briefly, with respect to the prolyl bond the substrates Succinyl-Ala-Xaa-Pro-Phe-4nitroanilide (Bachem) exist at pH 7.8 in an equilibrium of about 5-20% cis and 80-95% trans conformer. Alpha-chymotrypsin cannot readily split off the 4-nitroaniline residue in the cis conformer. In the presence of 0.5-1.0 mg ml⁻¹ α-chymotrypsin to perform rapid cleavage of the trans substrate it remains uncleaved in the solution. Because the cis form is subject to a slow isomerization reaction leading to the cleavable trans substrate, consumption of the total amount of the chromogenic peptide is achieved after several minutes. In the presence of PPlase activity the slow kinetic phase of appearance of 4nitroaniline following the trans cleavage is a composite of the uncatalysed (k1) and the PPlase-catalysed (kerz) cis to trans interconversion. Thus, the first-order kinetics obtained can be described by the rate equation $v = k_{obs} [cis]$; $k_{obs} = k_1 + k_{enz}$.

For all substrates used in the assay the range of total substrate concentrations was limited to 0.06-0.02 mM. Under the reaction conditions described above, the following rate constants of the uncatalysed cis to trans isomerization (k_1) have been determined: Xaa = Ala (0.0076 s⁻¹); Phe (0.0043 s⁻¹); Leu (0.0073 s⁻¹); Lys (0.0055 s⁻¹); Gly (0.0060 s⁻¹); His (0.0059 s-4); Glu (0.0034 s-4). Utilizing reaction conditions at high Mip-PPlase concentrations (k1 << kens) and rapid monitoring of the slow phase of the reaction (first data points after 5 s) it could be shown that the time-course of the 4-nitroaniline appearance is strictly first-order in rate throughout the reaction. The result shows that the relationship $k_{obs}/[PPlase] = k_{cat}/K_{m}$ holds true for all of the substrates. This was more directly indicated for bovine FK506-sensitive PPlase since the Michaelis constant, Km (0.52 mM cis isomer) of Suc-Ala-Leu-Pro-Phe-4-nitroanlide could be determined by measuring the V/S dependence at high substrate concentrations (Kofron et al.,

The PPlase assay was performed by adding a 1 µl aliquot of the peptide dissolved in dimethylsulphoxide (DMSO) to 1.1 ml of the solution of α-chymotrypsin (Merck) in the buffer containing Mip-PPlase, too. The concentration of Mip was determined by the Bradford procedure (1976). Stock solutions of the inhibitors were made in EtOH/water (50%/50%, v/v). For the kinetic runs a Hewlett Packard 8452 diode array UV/VIS spectrophotometer was used for monitoring the time course of the difference in absorbance at 390 nm and 520 nm. First-order rate constants (k_{obs}) were calculated on the basis of 100-500 data points covering at least two half-lives of the reaction.

Acknowledgements

This work was supported, in part, by grants from the BMFT (01 Ki 8829 and 30 K 008405) and the Fonds der Chemischen Industrie. We thank Laurence R. Phillips (Würzburg) for critical reading of the manuscript.

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