addenda and errata

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Preliminary crystallographic data of the three homologues of the thiol-disulfide oxidoreductase DsbA in *Neisseria meningitidis*. Corrigendum

Céline Lafaye,^a Thomas Iwema,^b Jean-Luc Ferrer,^b J. Simon Kroll,^c Mickael Griat^a and Laurence Serre^a*

^aLaboratoire des Protéines Membranaires, Institut de Biologie Structurale, CEA/CNRS/ Université Joseph Fourier, 41 Rue Jules Horowitz, 38027 Grenoble CEDEX 01, France, ^bLaboratoire de Cristallogénèse et Cristallisation des Protéines, Institut de Biologie Structurale, CEA/CNRS/Université Joseph Fourier, 41 Rue Jules Horowitz, 38027 Grenoble CEDEX 01, France, and ^cDepartment of Paediatrics, Imperial College London, St Mary's Hospital Campus, Norfolk Place, London W2 1PG, England

Correspondence e-mail: laurence.serre@ibs.fr

A correction is made to the name one of the authors of Lafaye *et al.* [*Acta Cryst.* (2008). F64, 111–114].

In the article by Lafaye *et al.* (2008) one of the authors names was incorrect. The correct name is Thomas Iwema as given above.

References

Lafaye, C., Iwena, T., Ferrer, J.-L., Kroll, J. S., Griat, M. & Serre, L. (2008). *Acta Cryst.* F64, 111–114.

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Céline Lafaye,^a Thomas Iwena,^b Jean-Luc Ferrer,^b J. Simon Kroll,^c Mickael Griat^a and Laurence Serre^a*

^aLaboratoire des Protéines Membranaires, Institut de Biologie Structurale, CEA/CNRS/ Université Joseph Fourier, 41 Rue Jules Horowitz, 38027 Grenoble CEDEX 01, France, ^bLaboratoire de Cristallogénèse et Cristallisation des Protéines, Institut de Biologie Structurale, CEA/CNRS/Université Joseph Fourier, 41 Rue Jules Horowitz, 38027 Grenoble CEDEX 01, France, and ^cDepartment of Paediatrics, Imperial College London, St Mary's Hospital Campus, Norfolk Place, London W2 1PG, England

Correspondence e-mail: laurence.serre@ibs.fr

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Preliminary crystallographic data of the three homologues of the thiol-disulfide oxidoreductase DsbA in *Neisseria meningitidis*

Bacterial virulence depends on the correct folding of surface-exposed proteins, a process that is catalyzed by the thiol-disulfide oxidoreductase DsbA, which facilitates the synthesis of disulfide bonds in Gram-negative bacteria. Uniquely among bacteria, the *Neisseria meningitidis* genome possesses three genes encoding active DsbAs: DsbA1, DsbA2 and DsbA3. DsbA1 and DsbA2 have been characterized as lipoproteins involved in natural competence and in host-interactive biology, while the function of DsbA3 remains unknown. In an attempt to shed light on the reason for this multiplicity of *dsbA* genes, the three enzymes from *N. meningitidis* have been purified and crystallized in the presence of high concentrations of ammonium sulfate. The best crystals were obtained using DsbA1 and DsbA3; they belong to the orthorhombic and tetragonal systems and diffract to 1.5 and 2.7 Å resolution, respectively.

1. Introduction

Neisseria meningitidis is an invasive bacterial pathogen that causes life-threatening infection in children worldwide. The host-pathogen interactions and therefore the virulence depend on the correct folding of many surface-exposed proteins, a process that is catalyzed by the thiol-disulfide oxidoreductase DsbA, which performs the synthesis of disulfide bonds in the periplasm of Gram-negative bacteria (Bardwell et al., 1991). This reaction is essential for the folding of virulence components, such as elements of the type IV secretion system, that require the formation of several disulfide bonds (Tinsley et al., 2004) to be functionally active. While Escherichia coli and many other Gram-negative bacteria, including human pathogens such as Shigella and Haemophilus, have a single dsbA gene, N. meningitidis uniquely possesses three genes (named nmb0278, nmb0294 and nmb0407 in the sequenced genome of the serogroup B strain MC58) encoding three active DsbAs (DsbA1, DsbA2 and DsbA3), each of which shows different enzymatic properties (Sinha et al., 2004; Tinsley et al., 2004). DsbA1 and DsbA2 are membraneassociated lipoproteins, while DsbA3 is a soluble periplasmic protein like its well studied E. coli counterpart. Deletion of each of these genes has highlighted that the presence of at least one of the membrane-anchored enzymes is sufficient to restore full neisserial virulence (Tinsley et al., 2004) or to maintain natural competence for DNA uptake at the wild-type level (Sinha et al., 2008). In contrast, the role of the periplasmic enzyme seems to be unrelated to either of these phenotypes. The neisserial enzymes share 21% sequence identity with their E. coli homologue, the crystal structure of which is composed of one thioredoxin domain and one helical domain (Martin, Bardwell et al., 1993). The three proteins are sufficiently similar to be defined unambiguously as DsbAs, but their primary sequences differ markedly, notably around the active-site domain (Fig. 1; Sinha et al., 2004; Tinsley et al., 2004). These differences, with their anticipated structural consequences, may underlie the distinct properties of each protein (Ondo-Mbele et al., 2005). Strikingly, the presence of three dsbA genes is a feature of N. meningitidis alone among neisserial species. N. gonorrhoeae, N. lactamica and some

other commensals only possess homologues of *nmb0278* and *nmb0407*, while other commensals such as *N. flava* possess only one *dsbA*, a homologue of *nmb0294* (Sinha *et al.*, 2004). These differences may reflect the striking differences in host–pathogen interactive biology, which is plausibly related in part to the folding of the substrates of DsbA. In order to begin to understand the specific contributions of each of these enzymes to neisserial biology, we have initiated a structural study of these three enzymes; here, we present preliminary crystallographic data obtained on the three *N. meningi-tidis* DsbAs.

2. Experimental and results

2.1. Cloning

Cloning of each of the *N. meningitidis dsbA* genes was achieved using the ROBIOMOL platform (Institut de Biologie Structurale, Grenoble, France).

PCR amplification was achieved with Phusion enzyme (Finnzymes) using a template containing the different recombinant neisserial *dsbA* genes (*nmb0278*, *nmb0294* and *nmb0407*) and specific primers containing the sequence essential for ligation-independent cloning. The PCR products were inserted into the vector pLIM01 (His₆ fusion, AmpR) by annealing to promote the cytoplasmic expression of the protein of interest in fusion with an N-terminal MAHHHHHGHHHQLENLYFQG tag (pLIM01) containing a tobacco etch virus (TEV) protease cleavage site. The ligation products were transformed into *E. coli* and the presence of the gene of interest in the vector was verified by restriction-endonuclease analysis. The nucleotide sequences encoding the periplasmic secretion signal and

the membrane-anchor sequence of the DsbA1 and DsbA2 proteins were excluded (Fig. 1).

2.2. Purification

The three recombinant proteins were expressed and purified at 277 K according to the same protocol, except that all the buffers used to purify DsbA3 were supplemented with 500 m*M* NaCl in order to avoid protein aggregation.

Recombinant plasmids were transformed into E. coli BL21(DE3)-Codon Plus-RIL. 11 LB broth medium containing 1 mM ampicillin was inoculated with 20 ml of an overnight culture at 310 K. At an OD_{600} of approximately 0.6, expression of the recombinant enzymes was induced by the addition of 1 mM IPTG and cells were grown for an additional 3 h. Cells were pelleted by centrifugation (5000g), resuspended in 10 mM HEPES pH 7.0 (buffer A) and disrupted using a French press. The soluble fraction was recovered by centrifugation at 18 000g and mixed with 5 ml of a suspension of Ni-NTA resin (Qiagen) pre-equilibrated with buffer A at 277 K. The resin was then loaded onto a column. To remove unbound proteins, the resin was extensively washed with 50 ml buffer A containing 50 mM imidazole. The proteins were eluted using a linear gradient of buffer A containing 50-400 mM imidazole. The pooled peak fractions were dialysed against buffer A and incubated in TEV buffer (50 mM Tris-HCl pH 8.0, 0.5 mM EDTA, 1 mM DTT) with recombinant His₆tagged TEV protease in a 1:100 molar ratio for 1 h at room temperature followed by 2 h at 277 K. The protein solution was dialysed against buffer A and loaded onto Ni-NTA beads to remove all His-tagged proteins. The purified proteins were then concentrated to approximately 10 mg ml⁻¹ using Centricon devices (Amicon,

DsbA1 DsbA2 DsbA3 Tcpg DsbA	1 MKSRHLALG MKLKTLALT MKLKHLLPL MKKLFALVA MKKIWLALA	O VAALFALAACS SLTLAIAACS LLSAVISAQA. TLMLSVSAYA. GLVLAFSASA.	30 SKVQTSVPAL KQAETSVPAL	9 40 DSAPAASAA.AAPAC DSAQSSSSAPAAPAE 	50 LVECQNYTVLANE LNECVNYTVLST LTEGEDYLVLCK FKECEHYQVLKT YEDCKQYTTLEKE	60 IPQQQAGKVEVLE IPQQQAGKIEVLE IPQEQSGKIEVLE ASSSPVVE AAGAPQVLE
DsbA1 DsbA2 DsbA3 Tcpg DsbA	70 FFGYFCPHC FFGYFCPHC FFGYFCVHC FFSFYCPHC FFSFCPHC	80 AHLEPVLS AHLEPVLS HHFDPLLL NTFEPIIA YQFEEVLHISD	90 KHAKSFKDDN KHIKTFKDD KLGKALPSDA QLKQQLPEGA NVKKKLPEGV	100 110 MYLRTE HVWQKEMI TYMRREHVWGDEMK AYLRTEHVWWQPEMI AKFQKNHVSFMGGNM /KMTKYHVNFMGGDI	120 T LARLAAAVON P LARLAAAVON G. LARMAAAVNI IGQAMSKAYATN GKDLTQAWAVAMA	130 MAAADSKDVANSHI MAGESDKANSHI SGL.KYQANPAV MIALEVEDKMVPVM ALG.VEDKVTVPL
DsbA1 DsbA2 DsbA3 Tcpg DsbA	140 EDAMVNQKI EDAMVNQKI EKAVYEQKI ENRIHTLRK EEGVQKTQT	150 KLQNPEVLKKW NLADTDTLKKW RLENRSVAGKW PPKDEQELRQI IRSASDIRDVF	160 LGEQTAF DG K LSEQTAF DG K ALSQKGF DG K FLDE.GI D AP INAGI <u>K</u> GE	170, 1 KKVLAAYESPESQAR KKVLAAFEAPESQAR KKIMRAYDSPEAAAA KKFDAAYNGFAVDSM SEYDAAWNSFVVKSI	80, 190, ADKMQELTETFQI AAQMEELTNKFQI ALKMQKLTEQYRI VRRFDKQFQDSGI VAQQEKAAADVQI	200 UGTPTVIVGGKYK SGTPTVIVGGKYQ DSTPTVIVGGKYQ TGVPAVVVNRYL RGVPAMFVNGKYQ
DsbA1 DsbA2 DsbA3 Tcpg	210 Vefad.w Vefkd.w VIfnngf Vo.gos.v	220 ESGMNTIDLLA QSGMTTIDQLV DGGVHTIKELV KSLD EYF	230 DKVREEQKA DKVREEQKKE AKVREERKRQ	AQ 2Q 2TPAVQK		

Figure 1

Sequence alignment of the DsbA homologues in *N. meningitidis* (DsbA1, DsbA2 and DsbA3), *E. coli* (DsbA) and *V. cholerae* (Tcpg). Identities (black shading) and highly conserved residues are highlighted. The arrow indicates the N-terminal residue of the three neisserial enzymes studied in this work. The conserved active-site tetrapeptide, Cys-X-X-Cys, is underlined.

LNPQGMD.TSNMDVFVQQYADTVKYLSEK.K....

DsbA

10 000 Da cutoff) and loaded onto a size-exclusion chromatography column (Hiload Superdex 75, 16/60, GE Healthcare) for the final step of the purification procedure. The column was equilibrated with buffer A and the pooled peak fractions were concentrated to 12 mg ml⁻¹. The purity of the protein solutions was confirmed by SDS–PAGE (Fig. 2). The molecular weights obtained by mass spectrometry were 21 477, 21 461 and 21 836 Da for DsbA1, DsbA2 and DsbA3, respectively. The final protein concentrations were determined by UV spectrometry assuming molar absorption coefficients at 280 nm of 24 075, 22 585 and 24 535 M^{-1} cm⁻¹ for DsbA1, DsbA2 and DsbA3, respectively. The final yield of each enzyme was 50 mg pure protein per litre of LB medium.

2.3. Oxidoreductase activity of the recombinant enzymes

To check the oxidoreductase activity of the three purified recombinant proteins, we tested their ability to catalyze the reduction of insulin interchain disulfide bonds by dithiothreitol (Holmgren, 1979). This reduction leads to insulin B-chain precipitation, which was monitored by the change in absorbance at 650 nm. Bovine insulin was concentrated to 1.7 m*M* in 25 m*M* HEPES and the final pH of the solution was adjusted to 8.0. A typical reaction mixture (0.2 ml) contained 150 μ *M* insulin in 100 m*M* potassium phosphate pH 7.0, 2 m*M* EDTA and 1 m*M* DTT. The precipitation of insulin was measured after the addition of 5 μ *M* of each enzyme and compared with a control that did not contain any protein. Fig. 3 shows that for all the DsbAs the reaction lag time is shorter than with the control, indicating that the three purified proteins exhibit thiol-disulfide oxidoreductase activity.

2.4. Crystallization

For crystallization, each protein was concentrated to 12 mg ml^{-1} in buffer A. As in the purification steps, buffer A was supplemented with 500 mM NaCl for the crystallization of DsbA3. Initial trials were performed using the sitting-drop method and commercial crystallization screening kits from Qiagen and Hampton Research. For the initial trials, we used a Tecan Genesis robot coupled with 96-well plates. 1 µl protein solution was mixed with 1 µl reservoir solution and the resulting drops were equilibrated against 100 µl reservoir solution. After optimization of the conditions in larger crystallization boxes (NeXTaL, 1 ml reservoir), the best DsbA1 crystals grew at



Figure 2

Analysis of the purified protein samples after gel-filtration chromatography. The purified samples were loaded onto a Coomassie Blue-stained 12% SDS–PAGE gel. Lanes M, A, B and C contain molecular-weight markers (kDa), recombinant DsbA2, DsbA1 and DsbA3, respectively.

room temperature in 2.4 *M* ammonium sulfate, 0.1 *M* bicine pH 9.0 or in 1.6 *M* trisodium citrate as thick plates or tetragonal bipyramids (Fig. 4*a*). DsbA2 crystals appeared in 2.4 *M* ammonium sulfate, 0.1 *M* MES pH 5.0 as hexagonal bipyramids at room temperature (Fig. 4*b*). DsbA3 crystallized as small rectangular prisms in 2.2 *M* ammonium sulfate, 0.25 *M* sodium malonate, 0.1 *M* MES pH 5.0 at 281 K (Fig. 4*c*). These crystals were routinely reproduced using the hanging-drop method of crystallization, mixing 2 µl protein solution with 2 µl reservoir solution.

2.5. Crystal parameters and X-ray diffraction

Prior to diffraction, the crystals were soaked in their respective crystallization solutions containing 30% glycerol as cryoprotectant. The crystals were then flash-cooled directly in liquid nitrogen. X-ray diffraction experiments were performed at the European Synchrotron Radiation Facility (Grenoble, France). Data were collected at 100 K on BM30 for DsbA1 crystals and on ID14-EH3 for DsbA2 and DsbA3. Diffraction data sets were processed using *MOSFLM* (Leslie, 1999) and intensities were scaled and reduced with *SCALA* (Evans, 1993) from the *CCP*4 program suite (Collaborative Computational Project, Number 4, 1994). Of the three proteins, only the DsbA2 crystals exhibited a poor diffraction pattern that did not allow us to determine the crystal parameters. The crystal parameters of DsbA1 and DsbA3 and their corresponding data-collection statistics are summarized in Table 1.

3. Conclusion

In this work, we cloned and purified the three DsbA enzymes from the pathogenic bacterium *N. meningitidis*. All the purified proteins were expressed at high levels in soluble form, lacking N-terminal signal sequences, in the cytoplasm of *E. coli*. The insulin-precipitation assay commonly used to test DsbA activity (Holmgren, 1979) shows that the three neisserial enzymes exhibit similar oxidoreductase activity and that the removal of the membrane anchor of the two lipoproteins does not abrogate this. The neisserial DsbAs are highly



Figure 3

Assay of the disulfide oxidoreductase activity of meningococcal DsbAs. The precipitation of insulin was measured after the addition of $5 \,\mu M$ DsbA1 (white triangles), DsbA2 (white circles) or DsbA3 (crosses). The absorbance owing to background precipitation of insulin by non-enzymatic reduction was measured in control tubes containing DTT but no added protein catalyst (white squares). The lag times measured for the control and the three meningococcal DsbAs are around 22 and 17 min, respectively.

crystallization communications





(b)

Figure 4

Crystals of the three *N. meningitidis* DsbA enzymes. All the enzymes crystallized in the presence of ammonium sulfate. Detailed crystallization conditions are described in the text. (*a*) DsbA1, (*b*) DsbA2, (*c*) DsbA3.

(c)

homologous; in particular, the two anchored membrane proteins are more than 70% identical. Accordingly, all three proteins crystallized under similar conditions requiring a high concentration of ammonium sulfate. This requirement is markedly different from the conditions employed for the successful crystallization of the *Vibrio cholerae* or *E. coli* DsbAs in polyethylene glycol (Martin, Waksman *et al.*, 1993; Hu *et al.*, 1997). The neisserial enzymes share 21% sequence identity with their *E. coli* and *V. cholerae* counterparts, the crystal structures of which are known (Martin, Bardwell *et al.*, 1993; Hu *et al.*, 1997). While this identity level might appear sufficient to solve the neisserial enzyme structures by molecular replacement either with the *E. coli* or *V. cholerae* models, no obvious solution has been obtained so far for DsbA1 or DsbA3 using this method of phasing.

Table 1

Crystal parameters and data-collection statistics calculated using SCALA (Evans, 1993).

Values in parentheses are for the last resolution shell.

Protein	DsbA1	DsbA2	DsbA3
N. meningitidis B serogroup gene	nmb0278	nmb0294	nmb0407
Space group	$P2_{1}2_{1}2_{1}$	Not determined	I4 ₁ 22
Unit-cell parameters (Å)	a = 44.08, b = 46.42, c = 79.93		a = b = 160.0, c = 76.80
Molecules per ASU [†]	1		2
$V_{\rm M}$ (Å ³ Da ⁻¹)	1.9		2.8
Solvent content (%)	35		56
Resolution (Å)	20.5-1.5 (1.58-1.5)		20-2.8 (2.95-2.8)
ESRF beamline	BM30A		ID14-EH3
Wavelength (Å)	0.979		0.931
R _{merge} ‡	0.064 (0.241)		0.102 (0.282)
$\langle I/\sigma(I) \rangle$	6.8 (3.1)		6.1(2.7)
Total observations	274425		162296
Unique reflections	26907		12695
Multiplicity	10.2		12.9
Data completeness (%)	99.5		100

† Estimated from the Matthews coefficient value (Matthews, 1968). $\ddagger R_{\text{merge}} = \sum_{hkl} \sum_i |I_i(hkl) - \overline{I(hkl)}| / \sum_{hkl} \sum_i \frac{I_i(hkl)}{I(hkl)}$, where $I_i(hkl)$ is the *i*th measurement of the intensity of the *hkl* reflection and $\overline{I(hkl)}$ is the average intensity obtained with these *i* measurements.

While it is clear on the basis of the sequence that the three neisserial proteins exhibit the DsbA general fold, it seems highly likely that their three-dimensional structures include some significant differences from the *E. coli* and *V. cholerae* enzyme structures. Heavy-atom derivative research is currently in progress in order to solve these crystal structures.

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