Structural Determination of Functional Units of the Nucleotide Binding Domain (NBD94) of the Reticulocyte Binding Protein Py235 of Plasmodium yoelii

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Abstract

Background: Invasion of the red blood cells (RBC) by the merozoite of malaria parasites involves a large number of receptor ligand interactions. The reticulocyte binding protein homologue family (RH) plays an important role in erythrocyte recognition as well as virulence. Recently, it has been shown that members of RH in addition to receptor binding may also have a role as ATP/ADP sensor. A 94 kDa region named Nucleotide-Binding Domain 94 (NBD94) of Plasmodium yoelii YM, representative of the putative nucleotide binding region of RH, has been demonstrated to bind ATP and ADP selectively. Binding of ATP or ADP induced nucleotide-dependent structural changes in the C-terminal hinge-region of NBD94, and directly impacted on the RBC binding ability of RH.

Methodology/Principals Findings: In order to find the smallest structural unit, able to bind nucleotides, and its coupling module, the hinge region, three truncated domains of NBD94 have been generated, termed NBD94 444–547, NBD94 566–663 and NBD94 674–793, respectively. Using fluorescence correlation spectroscopy NBD94 444–547 has been identified to form the smallest nucleotide binding segment, sensitive for ATP and ADP, which became inhibited by 4-Chloro-7-nitrobenzofurazan. The shape of NBD94 444–547 in solution was calculated from small-angle X-ray scattering data, revealing an elongated shape with three architectural segments. By comparison, the low resolution structure of NBD94 674–793 in solution represents a chair–like molecule, comprised of two globular domains, connected by a spiral segment of about 73.1 Å in length. The high quality of the constructs, forming the hinge-region, NBD94 566–663 and NBD94 674–793 enabled to determine the first crystallographic module, the hinge region, three truncated domains of NBD94 have been generated, termed NBD94 444–547, NBD94 566–663 and NBD94 674–793, respectively. Using fluorescence correlation spectroscopy NBD94 444–547 has been identified to form the smallest nucleotide binding segment, sensitive for ATP and ADP, which became inhibited by 4-Chloro-7-nitrobenzofurazan. The shape of NBD94 444–547 in solution was calculated from small-angle X-ray scattering data, revealing an elongated molecule, comprised of two globular domains, connected by a spiral segment of about 73.1 Å in length. The high quality of the constructs, forming the hinge-region, NBD94 566–663 and NBD94 674–793 enabled to determine the first crystallographic and solution structure, respectively. The crystal structure of NBD94 566–663 consists of two helices with 97.8 Å and 48.6 Å in length, linked by a loop. By comparison, the low resolution structure of NBD94 674–793 in solution represents a chair–like shape with three architectural segments.

Conclusions: These structures give the first insight into how nucleotide binding impacts on the overall structure of RH and demonstrates the potential use of this region as a novel drug target.

Introduction

Malaria continues to be one of the major public health problems for mankind; it is the transmissible disease with the greatest morbidity around the world [1]. The complex life cycle of the protozoan parasite is characterized by distinct invasive forms of the sporozoite and merozoite that invade hepatocytes and erythrocytes in the vertebrate host, respectively, and the ookinetes inside the insect vector that penetrates the mosquito midgut epithelium [1–6]. Invasion of red blood cells (RBC) by the merozoite and the subsequent cyclical replication of the parasite is the cause of malaria associated pathology. Multiple merozoite protein families are implicated in the invasion of RBCs, including the erythrocyte binding proteins (EBPs) and the reticulocyte binding protein homologues (RH), which bind to different RBC membrane receptors [1,6–16]. Little is known about how the large RH transmembrane proteins mediate their function during erythrocyte invasion, but a crucial step appears to be the proteolytic cleavage during the invasion process [14,17]. Members of RH have been identified in all Plasmodium species so far analyzed indicating the conserved function and importance of this protein family to the malaria parasite [1]. The P. yoelii RH protein, termed Py235 (235 kDa in mass), has been shown to be a potential virulence factor that allows the parasite to invade a wider range of host erythrocytes [18–20]. Py235 is also involved in clonal phenotypic variation of merozoites [21], enabling the parasite to evade immune responses and adapt to changes in the host environment during the invasion step [22]. Previous studies in P. vivax has indicated that RH may have an initial sensing role preceding and possibly enabling the subsequent interaction of the
EBP member with its corresponding receptor [23]. Besides multiple receptor ligand interaction between the merozoite and the RBC, the effects of other intra and/or extra cellular molecules like ATP on invasion have been discussed [24–27]. ATP, released by erythrocytes under normal physiological conditions, has recently been shown to serve as a signalling molecule regulating vascular tone [24–26] and ATP receptors have been implicated in a number of signalling pathways [28,29]. Mechanical deformation of erythrocytes as encountered in capillaries forming the microvasculature lead to increased ATP release [30]. Importantly, intracellular ATP is a requirement for merozoite invasion [31–34] with erythrocytes that have been depleted of ATP being refractory to invasion. These findings suggest that it would be advantageous for merozoites to sense the intracellular ATP level of the erythrocyte. In this context a 94 kDa domain of Py235 of *P. yoelii*, which is highly conserved amongst the RBLs and called the nucleotide-binding domain (NBD94), has been found to selectively bind ATP and ADP. The amino acid sequence 488EKLKHYNFDDFVK500 in NBD94 has been identified as nucleotide-analogue binding sequence, together with Py235 to RBCs in the presence of MgATP which is significantly lower in the presence of MgADP or the absence of nucleotides [27]. Based on these traits and the absence of significant ATPase activity of NBD94, this domain was suggested to serve as an ATP/ADP sensor during the invasion process [27]. However, the recombinant NBD94 protein is prone to degradation over a period of time. Whether degradation occurs through autolysis or proteolytic cleavage, as described for a series of proteolytic cleavage events of both the EBP and RH protein families during invasion, resulting in 9 to 13 kDa fragments, is unclear [35,36].

In many cases a protein fragment obtained by cleavage of the full length protein or by expression of part of the protein can retain a functional domain. This is particularly relevant in drug designing. In order to get insight into the functional regions of NBD94 and to define the smallest segment, still able to bind ATP/ADP, three truncated constructs, called NBD94444–547, including the 8-N3-3'-biotinyl-ATP binding sequence, as well as NBD94566–663 and NBD94674–793, described to belong to the C-terminal hinge region and to couple the nucleotide-binding event [27], have been generated (Fig. 1) and purified to homogeneity. As demonstrated by fluorescence correlation spectroscopy (FCS), NBD94444–547 forms the smallest ATP/ADP-binding segment, which becomes inhibited by the ATPase/-synthase inhibitor 4-Chloro-7-nitrobenzofurazan (NBD-C3). The high quality of all proteins formed the platform to solve the first low resolution structure of NBD94444–547, and NBD94674–793 in solution and the first crystallographic structure of NBD94566–663. The structural features of this ensemble will be discussed in the light of coupling of nucleotide-binding with structural rearrangements in the hinge region, followed by downstream effects, enabling the parasite to continue the invasion process.

**Results**

**Production and Purification of Truncated Forms of NBD94 of Plasmodium yoelii**

In order to understand the events of nucleotide-binding in NBD94 and its concerted structural alteration(s) in the C-terminal hinge region, as well as to determine the smallest segment of NBD94, still able to bind nucleotide, and thereby forming an attractive target for therapeutic agents, the truncated forms NBD94444–547, NBD94566–663 and NBD94674–793 have been generated (Fig. 1), in which predicted α-helical structures have been taken into account (see below). The SDS-PAGE of the produced recombinant NBD94444–547, NBD94566–663 and NBD94674–793 of *Plasmodium yoelii* revealed a prominent band of 13.2 kDa, 12.8 kDa and 15.2 kDa, respectively, which was found entirely within the soluble fraction. ANF94-NTA resin column and an imidazole-gradient were used to separate NBD94444–547, NBD94566–663 and NBD94674–793 of *Plasmodium yoelii* revealed a prominent band of 13.2 kDa, 12.8 kDa and 15.2 kDa, respectively, which was found entirely within the soluble fraction. ANF94-NTA resin column and an imidazole-gradient were used to separate NBD94444–547, NBD94566–663 and NBD94674–793 of *Plasmodium yoelii* revealed a prominent band of 13.2 kDa, 12.8 kDa and 15.2 kDa, respectively, which was found entirely within the soluble fraction. ANF94-NTA resin column and an imidazole-gradient were used to separate NBD94444–547, NBD94566–663 and NBD94674–793, respectively, from the main contaminating proteins. The protein NBD94444–547, NBD94566–663 and NBD94674–793, respectively, eluting at 125–300 mM imidazole were collected, concentrated and subsequently applied to a size exclusion column (Superdex 75 HR 10/30 column).

![Figure 1. Domain features of Py235 of *P. yoelii*. Three truncated constructs, called NBD94444–547 and NBD94566–663 as well as NBD94674–793, both described to belong to the C-terminal hinge region [27], have been generated. NBD94444–547 includes the peptide 488EKLKHYNFDDFVK500, which has been shown to covalently bind the nucleotide-analogue 8-N3-3'-biotinyl-ATP [27], highlighted as a star.](https://www.plosone.org/doi/abs/10.1371/journal.pone.0009146.g001)
Analysis of the isolated protein by SDS-PAGE revealed the high purity of NBD94_{444-547}, NBD94_{566-663} and NBD94_{674-793} (Fig. 2A–C).

The secondary structures of these proteins were determined from circular dichroism spectra, measured between 185–260 nm (Fig. 2A–C). The CD-spectra of NBD94_{444-547}, NBD94_{566-663} and NBD94_{674-793} show that all proteins are mainly α-helical, as reflected by its minima at 208 and 222 nm and as predicted from its amino acid sequence. The α-helical content of NBD94_{444-547}, NBD94_{566-663} and NBD94_{674-793} is determined to be 83%, 86% and 69%, respectively, congruent to the values determined for the entire NBD94 and indicating the correct secondary structure of the recombinant truncated proteins generated, as well as reflecting a proper selection of stable constructs. The molar ellipticity values at 208 nm and at 222 nm of NBD94_{444-547}, NBD94_{566-663} and NBD94_{674-793} are in a ratio of 0.94, 0.99 and 0.97, respectively.

Shape Determination of NBD94_{444-547} and NBD94_{674-793} in Solution

The high purity allowed small-angle X-ray scattering (SAXS) experiments to be performed, with the aim to determine the first low resolution structures of the nucleotide-binding segment NBD94_{444-547} and NBD94_{674-793} in solution. SAXS patterns from solutions of both proteins were recorded as described in Materials and Methods to yield the final composite scattering curves in figure 3A and 4A, respectively, that both proteins are monodispersed in solution. The radius of gyration $R_g$ of NBD94_{444-547} is 33.4±1 Å and the maximum dimension $D_{max}$ of the protein is 134±2 Å (Fig. 3B). The gross structure of NBD94_{444-547} was restored ab initio from the scattering pattern in figure 3A using the dummy residues modeling program GASBOR [37], which fitted well to the experimental data in the entire scattering range (a typical fit displayed in figure 3A, curve 2, has the discrepancy $\chi = 1.28$). Ten independent reconstructions yielded reproducible models and the average model and the most probable model are displayed in figure 3C. NBD94_{444-547} appears as an elongated molecule with a length of 134 Å, composed of two more globular domains and a spiral-like segment of about 73.1 Å in length between both domains.

By comparison, the radius of gyration $R_g$ of NBD94_{674-793} is 22.2±1 Å nm and the maximum dimension $D_{max}$ of the protein is 118±2 Å (Fig. 4B), respectively, suggesting that the protein is rather elongated. The shoulders at larger intraparticle distances of about 70 Å and 100 Å indicate that the molecule consists of three distinct domains. Like for NBD94_{444-547} the shape of NBD94_{674-793} was determined ab initio from the scattering pattern in figure 4A using the program GASBOR [37]. The experimental data fitted well with the calculated data, reflected by an $\chi$ value of 1.03 (Fig. 4A, curve (red)). The ab initio modeling produced a chair–like shape (Fig. 4C) with three domains of 96 Å, 35 Å and 20 Å in length. When rotating the molecule by 90°, the longer domain has a spiral feature and the lower 20 Å long domain turns away from the middle one by 104.5°.
Nucleotide-Binding Determined by Fluorescence Correlation Spectroscopy

The proper structural folding and structure formation of NBD94\textsubscript{444-547} enabled us to study the ability of this protein to bind nucleotides by fluorescence correlation spectroscopy using fluorescent ATP and ADP derivatives ATP ATTO-647N and ADP ATTO-647N, respectively. As a reference, the mean count rate per Cyanine 5 (Cy5) fluorophore was determined to be 32.5 ± 0.4 kHz. Compared to Cy5, the value of ATP ATTO-647N was determined to be 27.9 ± 0.8 kHz and 51.3 ± 3.5 kHz for ADP ATTO-647N. Fitting the autocorrelation functions resulted in characteristic times of diffusion $t_D = 50 ± 1.1$ μs for Cy5, $t_D = 70.2 ± 1.3$ μs for ATP ATTO-647N and $t_D = 68.9 ± 2.2$ μs for ADP ATTO-647N. The autocorrelation curves of the fluorescent ATP analogue for ATP ATTO-647N and ADP ATTO-647N in the absence and presence of increased concentrations of NBD94\textsubscript{444-547} are shown in figure 5A and in supplementary figure S1A, respectively. The increase of the mean diffusion time $t_D$ was due to the increase in the mass of the diffusing particle, when fluorescently labelled nucleotide bound to NBD94\textsubscript{444-547}, which is apparent in the displayed autocorrelation curves with increased protein concentrations from left to right. A binding constant ($K_d$) of 228 ± 2.3 μM for MgATP ATTO-647N and 331 ± 1.8 μM for MgADP ATTO-647N bound to NBD94\textsubscript{444-547} was determined (Fig. 5B, C). By contrast no nucleotide-binding could be observed in the constructs NBD94\textsubscript{566-663} and NBD94\textsubscript{674-793}.

Most recently, we observed that 4-Chloro-7-nitrobenzofurazan (NBD-Cl) is a potent inhibitor of nucleotide-binding in the entire NBD94 [27]. We tested, whether MgATP ATTO-647N-binding of the minimal nucleotide-core domain, NBD94\textsubscript{444-547}, becomes inhibited by NBD-Cl. The plotted autocorrelation functions in

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**Figure 3. Small-angle X-ray scattering data of NBD94\textsubscript{444-547} of *P. yoelii*. (A) Experimental scattering data (o) and the fitting curves (---; green: experimental, red: calculated from *ab initio* model) for NBD94\textsubscript{444-547} of *P. yoelii*. (B) The distance distribution function of the same protein. (C) Low resolution structure of NBD94\textsubscript{444-547} in solution determined from SAXS data. The two shapes are rotated clockwise by around 90° along the Y-axis. doi:10.1371/journal.pone.0009146.g003**
supplementary figure S1B show a change of the diffusion time due to an increase of the concentration of NBD-Cl from right to left for MgATP ATTO-647N. The calculated bound fraction for increasing inhibitor concentrations were plotted to determine the effect of nucleotide binding. The interaction of MgATP ATTO-647N to NBD94–547 showed an IC₅₀ value of 400 ± 3 μM (Fig. 5D). A similar effect of NBD-Cl-inhibition has been observed for NBD94–547-ADP-ATTO-647N formation (data not shown).

Crystallographic Structure of NBD94_566–663

The second domain, NBD94_566–663, covering the amino acids close to the nucleotide binding segment NBD94_444–547, has been crystallized by vapor diffusion using 2-methyl-2,4-pentanediol as a precipitant in a form suitable for X-ray diffraction analysis. This success was due to controlling the rate of vapor diffusion via the introduction of an oil barrier over the reservoir of a vapor-diffusion trial. The two different oils, paraffin and silicone, were applied as barriers, separately and also as mixtures of these two oils. Paraffin oil enabled the increase of crystal size [Supplementary Figure S2A and S2B] as well as improvement in resolution. The native crystal diffracted to 4.1 Å while the SeMet crystals diffracted to 3.9 Å at peak wavelength. A MAD dataset was collected which was indexed, integrated and scaled using the HKL-2000 [38] package. The data collection statistics are given in Table 1. Both the native and the SeMet derivative, crystallizes in the hexagonal space group P6₃22 with three molecules in the asymmetrical unit and a solvent content and Vₐ value of 35.44% and 1.9 Å³ per Dalton, respectively [39]. The data collection statistics are given in Table 1.

Both MAD and SAD techniques for structure solution were tried. Various programs like, SHELX [40], SOLVE [41], CNS [42] and CCP4 [43] were used to identify the heavy atom sites (Se). Reasonable statistics and good initial map could be achieved with the SAD technique and using the help of SHELX program [40]. The resolution cut-off used for Se site identification is 4.4 Å and the correlation co-efficient stood at 39.51% with the Pseudo-
free CC of 71.47%. Peak wavelength (0.97898 Å) data was used for the SAD technique in which 12 out of 18 Se sites (6 SeMet in each molecule) could be identified by SHELXD [44] that were further refined by SHELXE [45]. These 12 Se-sites were used to phase the structure factor with SHELXE and the resulting electron densities were improved by solvent flattening with SOLOMON and density modification by DM from the CCP4 package and RESOLVE [46]. The helical region could be readily identified from the initial map and the model of the NBD94566–663 was built manually using the program COOT [47]. Several cycles of manual building and fitting were carried out by COOT in combination with restrained refinement using REFMAC5 [48] of the CCP4 suite, keeping the temperature factor at overall. Each time a simulated annealing omit map was calculated and was used for model building (Supplementary Figure S3). The electron density map for the main chain atoms is excellent showing the typical sausage like features for the helical regions (Fig. 6A), whereas for the side chain atoms no visible densities could be identified. Se positions from the anomalous map were used to trace the chains. The final refined model has an R-factor of 33.9% and a R-free of 37.32% with good stereochemistry as can be inferred from the Ramachandran plot statistics given by PROCHECK [49]. The detailed summary of the refinement statistics is given in Table 1. The final electron density map for the structure shows good density for most of the backbone residues except for the residues 1–6 and 73–75 in chain A (Fig. 6A), 1–12 and 73–75 in chain B and 1–13, 31–33 and 73–75 in chain C, respectively (Fig. 6B).

The structure of the NBD94566–663 molecule consists of two helices that are linked by a loop, which is not visible in the electron density map. The length of the long and short helix is 97.8 Å and 48.6 Å, which are formed by 65 and 30 residues, respectively (Fig. 6B). Amino acids Gly61 and Ser85 of the N- and C-terminal helix, respectively, are in a very close proximity of 9.85 Å. Due to the kinks from residues Tyr60-Lys62 and Ser85-Glu87 both helices spread apart by an angle of 40.2°, thereby forming a hinge-like feature. The total number of amino acid residues is 107 that includes the N-terminal His-tag residues. The longer helix of chain

Figure 5. Binding traits of NBD94444–547 to fluorescently labeled MgATP ATTO-647N. (A) Normalized autocorrelation functions of MgATP ATTO-647N (B) obtained by increasing the quantity of NBD94444–547 (increased protein concentration from left to right). (C) Binding of NBD94444–547 to MgADP ATTO-647N. The nucleotide analogue is displayed as relative bound fraction versus protein concentration. The best fit to titration curve A and B are shown as a non-linear, logistic curve fit. (D) Influence of NBD-Cl to MgATP ATTO-647N binding traits of NBD94444–547. The best fits at titration curves of supplementary figure 1B are shown as a pharmacological dose-response curve with variable Hill slope.

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C in the asymmetric unit is highly distorted when compared to the other molecules (Fig. 6B). Slight kinks could be noted in the long helix from residues Val20-Lys22 and Tyr60-Lys62 in chain A. Whereby in chain B, the residues Asp24-Lys26 show slight helical distortion but for residues Leu37-Lys47 the helix is deviated more. The shorter helices of all the three molecules do show some amount of distortion but is less when compared to the long helices, wherein residues Ser85-Glu87 in chain A, Glu80-Met82 in chain B and Tyr89-Lys91 in chain C show notable deviations.

### Discussion

The mechanism by which a merozoite recognizes a suitable host cell is mediated by a cascade of receptor ligand interactions. In addition to the availability of the appropriate receptors, intracellular ATP plays an important role in determining if erythrocytes are suitable for merozoite invasion [24–26]. In erythrocyte binding assays (EBA) it has been demonstrated that Py235 of *P. yoelii* binds strongly to erythrocytes in the presence of ATP, whereby weak interactions have been found in the presence of ADP or the absence of nucleotides [27]. The ATP/ADP modulation of Py235-receptor binding suggested a nucleotide-dependent rearrangement, making the binding domain of Py235 more accessible. Such a nucleotide induced change has been observed in the 94 kDa nucleotide-binding domain, NBD94, of Py235, in which ATP-binding causes alterations in the C-terminal hinge region [27].

### Table 1. Statistics of crystallographic data collection for NBD94_{566–663}.

| Se-derivative | Peak | Edge | Remote |
|---------------|------|------|--------|
| **Data collection statistics** | | | |
| Wavelength (Å) | 0.97898 | 0.97936 | 0.96412 |
| Space group | P6,22 | P6,22 | P6,22 |
| **Unit cell parameters** | | | |
| a = b = 70.22 Å | 193.22 Å | 90° | 120° |
| c | 193.22 Å | 90° | 120° |
| Resolution range (Å) | 30–3.9 | 30–4.0 | 30–4.15 |
| Number of unique reflections | 2936 | 2762 | 2789 |
| Completeness (%) | 95.6 (97.1) | 85.6 (86.9) | 81.6 (76.9) |
| R merged (%) | 10.2 (39.3) | 6.8 (42.9) | 7.0 (34.8) |
| Multiplicity | 15.4 (12.4) | 15.0 (12.1) | 14.4 (11.6) |
| **Refinement statistics** | | | |
| Resolution range (Å) | 30–4.0 |
| R factor | 33.90 |
| R free | 37.32 |
| Ramachandran statistics | | | |
| Most favored (%) | 81.1 |
| Additionally allowed (%) | 17.3 |
| Generously (%) | 1.6 |
| R.M.S. deviations | | | |
| Bond lengths (Å) | 0.004 |
| Bond angles (°) | 0.839 |
| Mean atomic B values | Overall | 64.57 |
| | Wilson plot | 81.82 |

*Values in parentheses refer to the corresponding values of the highest resolution shell; for peak (4.04–3.9), edge (4.14–4.0) and remote (4.14–4.0).*

* R factor = Σ|Fobs|−|Fcalc|/Σ|Fobs|, where Fobs and Fcalc are measured and calculated structure factors, respectively.

* R free = Σ|Fobs|−|Fcalc|/Σ|Fcalc|, calculated from 10% of the reflections selected randomly and omitted from the refinement process.

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C-terminal region of NBD94 may mediate the nucleotide-binding with further downstream events [27]. The NBD94<sub>566–663</sub> molecule with its long N-terminal and shorter C-terminal α helices, linked by a flexible loop, shows a hinge-like arrangement (Fig. 6B). In the crystallographic structure, the residues Gly61 and Ser85 of the N- and C-terminal helix, respectively, are in a very close proximity of 9.8 Å and the very N- and C-terminal amino acids of NBD94<sub>566–663</sub> spread apart by an angle of 40.2°. This structural feature of NBD94<sub>566–663</sub> enables this protein to move either like a hinge or like a rattle up and down, thereby transmitting ATP-ADP modulation in NBD94<sub>444–547</sub> with up- and down movements in NBD94<sub>566–663</sub>, which have to be coupled to the α-helical and chair-like NBD94<sub>674–793</sub> fragment of the C-terminal region of NBD94, representing an elongated protein with three subdomains of 96 Å, 35 Å and 20 Å in length. Different orientation of the protein ensemble relative to each other have been speculated, in which the N-terminal segment NBD94<sub>444–547</sub> is linked to the N-terminal helix of NBD94<sub>566–663</sub> and the C-terminal segment NBD94<sub>674–793</sub> is oriented to the C-terminal helix of NBD94<sub>566–663</sub> (Fig. 7A–C). In all hypothesized models, the structure of the NBD94<sub>566–663</sub> molecule forms the middle element, able to mediate the sensing of nucleotide signaling and Py235 binding to the blood cell. An alternative prediction is presented in figure 7D revealing the possibility of a direct NBD94<sub>444–547</sub> and NBD94<sub>674–793</sub> contact, in which NBD94<sub>566–663</sub> presents a linking element. The increased nucleotide affinity in presence of the C-terminal domain of NBD94 determined recently [27] may hint at such an assembly.

In summary, the data presented demonstrate that NBD94 can be divided into a nucleotide-binding segment, NBD94<sub>444–547</sub>, existing in solution as an elongated molecule. The first crystallographic and solution structure of NBD94<sub>566–663</sub> and the C-terminal domain NBD94<sub>674–793</sub> of the C-terminal hinge region in NBD94 provide the structural basis towards a better understanding of a concerted interaction of the protein ensemble in NBD94, which is triggered by ATP, a requirement for merozoite invasion [31–34]. Furthermore, this work will provide the foundation for future studies to identify new compounds that directly interfere with the invasion process.

Materials and Methods

Biochemicals

Pfu DNA polymerase and restriction enzymes were purchased from New England Biolabs (Ipswich, MA, USA). Ni<sup>2+</sup>-Sepharose<sup>TM</sup> High Performance chromatography resin was obtained from GE Healthcare Bio-Sciences AB (Uppsala, Sweden). Chemicals for gel electrophoresis were received from Biorad (Hercules, USA). The ATP- and ADP-analogues EDA-ATP ATTO-647N and EDA-ADP ATTO-647N were received from ATTO-TEC (Siegen, Germany). All other chemicals were at least of analytical grade and purchased from Sigma-Aldrich (Deisenhofen, Germany), Amersham Bioscience (Buckinghamshire, UK), BIOMOL (Hamburg, Germany) and Merck (Darmstadt, Germany).

Cloning, Expression and Protein Purification

In order to obtain the truncated domains NBD94<sub>444–547</sub>, NBD94<sub>566–663</sub> and NBD94<sub>674–793</sub> of NBD94, the following primers were designed: forward primer 5'-AAC ATC GAA CCA TGG TAA TTC CAT-3' and reverse primer 5'-CGG TTA TTT GGA GCT CTT ACG TTT-3' for NBD94<sub>444–547</sub> forward primer 5'-GAT CCA CCA TGG TAA AGG AAA-3' and reverse primer 5'-AAC ATC GAA CCA TGG TAA TTC CAT-3' and reverse primer 5'-CGG TTA TTT GGA GCT CTT ACG TTT-3' for NBD94<sub>444–547</sub> forward primer 5'-GAT CCA CCA TGG TAA AGG AAA-3' and reverse
primer 5'-GAT ATA TTA AAG AGC TCT TAT GTG TTC ATT T'-3' for NBD94_444-547, and finally forward primer 5'-ATA AAG ATC CAT GGT ACA TTA TAT TAC TAG-3' and reverse primer 5'-TTG ATT CAG GCT CTT ATA TTT TCG ATT-3' for NBD94_674-793. The genomic P. yoelii YM DNA was used as the template. In all the constructs the restriction sites NcoI and SacI were incorporated. Following digestion with NcoI and SacI, the PCR products were ligated into the pET9d1-His3 vector [54]. The pET9d-His3 vector, containing the respective gene, was then transformed into E. coli cells (strain BL21 (DE3)) and grown on 30 mg/ml kanamycin-containing Luria-Bertani (LB) agar-plates. To express NBD94_444-547, NBD94_566-663 and NBD94_674-793, liquid cultures were shaken in LB medium containing kanamycin (30 µg/ml) for about 20 h at 37°C until an optical density of 0.6–0.7 (OD 600 nm) was reached. To induce production of the recombinant proteins, the cultures were supplemented with isopropyl (thio)-β-D-galactoside (IPTG) to a final concentration of 1 mM. Following incubation for another 4 h at 37°C, the cells were harvested at 6 000×g for 20 min, 4°C. Selenomethionine containing protein was expressed in the same way except that methionine biosynthesis was inhibited by the growth conditions [55]. A 1:1000 preculture in LB medium was used to inoculate 300 ml of M9 minimal medium supplemented with 0.4% glucose, 2 mM MgSO4, 30 µg/ml kanamycin, vitamins and trace elements. After overnight growth, the culture was diluted 1:100 into 1 l of minimal medium. After an OD600 of 0.6 was reached, 100 mg/l DL-selenomethionine (Sigma), 100 mg/l lysine, threonine and phenylalanine and 50 mg/l leucine, isoleucine and valine were added as solids. IPTG (1 mM) was added after 15 min, and cells were grown for 3 h and harvested as described above. Cells were lysed on ice by sonication for 3×1 min in buffer A (50 mM Tris/HCl, pH 7.5, 200 mM NaCl, and 2 mM PMSF) for NBD94_444-547, buffer B (50 mM Tris/HCl, pH 9.0, 200 mM NaCl, and 2 mM PMSF) for NBD94_566-663 and buffer C (50 mM Tris/HCl, pH 7.5, 200 mM NaCl, 2 mM PMSF and 0.5 mM DTT) for NBD94_674-793. The lysate was centrifuged at 10 000×g for 35 min. The supernatant was filtered (0.45 µm; Millipore) and passed over a Ni2+-NTA resin column to isolate NBD94_444-547, NBD94_566-663 and NBD94_674-793, respectively. The His-tagged protein was allowed to bind to the matrix for 1.5 h at 4°C and eluted with an imidazole-gradient (25–300 mM) in buffer A, B and C, respectively. Fractions containing His-NBD94_444-547, NBD94_566-663 and NBD94_674-793, were identified by SDS-PAGE [56], pooled and concentrated using Centricon YM-3 (3 kDa molecular mass cut off) spin concentrators (Millipore). The concentrated samples were applied on a size-exclusion column (Superdex™ 75 10/300 GL, GE Healthcare) using their respective buffers with additional 10 mM EDTA. The purity of all protein samples were analyzed by SDS-PAGE [56] and the gels were stained with Coomassie Brilliant Blue G250. Protein concentrations were determined by the bicinchoninic acid assay (BCA; Pierce, Rockford, IL., USA).

Circular Dichroism (CD) Spectroscopy

Measurements of steady state CD spectra were carried out in the far UV-light (185–260 nm) using a CHIRASCAN

Figure 7. Hypothesized arrangements of the three NBD94 segments. (A–C) A gallery of possible arrangements of the nucleotide-binding region NBD94_444-547 (green), the crystallographic NBD94_566-663 structure (dark green) and the very C-terminal structure of NBD94_674-793 (olive), in which the N-terminal helix NBD94_566-663 is linked to the N-terminal NBD94_444-547 segment, while the C-terminal helix of NBD94_566-663 is linked to the very C-terminal NBD94_674-793. (D) An arrangement in which NBD94_444-547 and NBD94_674-793 are in close proximity.

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s spectrophotometer (Applied Photophysics, UK). The CD spectroscopy measurements of NBD94444–547, NBD94674–793 and NBD94674–793 were performed in their corresponding buffers (see above) using a 60 μl quartz cell (Hellma) at 18°C with 1 nm step points and a protein concentration of 2 mg/ml for each recombinant protein. The spectrum for the buffer was subtracted from the spectrum of the protein. The ellipticity values were calculated by the average of triple determinations for each sample with a bandwidth of 1 nm from 185 to 260 nm (1 nm step points). The CD data were converted to mean residue ellipticity (θ) in units of deg x dmol⁻¹ x nm⁻¹ using the software Chirascan version 1.2.1 (Applied Photophysics). Baseline corrected spectra were used as input for computer methods to obtain predictions of secondary structure. In order to analyze the CD spectrum, the following algorithms were used: Varselec [57], Selcons [58], Contin [59], K2D [60]; all methods as incorporated into the program Dicroprot [61] and NeuralNet [62].

Fluorescence Correlation Spectroscopy (FCS)

Fluorescence correlation spectroscopy was performed at 25°C on a LSM 510 Meta/ConfoCor 3 (Zeiss, Jena, Germany) using the ATP-analogue EDA-ATP ATTO-647N (ATTO-TIC, Siegen, Germany). The temperature was adjusted to 25°C in an incubation chamber (Zeiss). The 635 nm laser line of a HeNe633 laser was attenuated to 6 mW and focused into the aqueous solution by a water immersion objective (40×/0.9 W Corr UL-VIS-IR, Zeiss). The proteins were dissolved in the buffer, containing 50 mM Tris, pH 7.5, and 200 mM NaCl. FCS was measured in 15 μl droplets of the diluted fluorescent derivatives of ATP, which were placed on Nunc 8 well chambered cover glass. Before usage, the cover glasses were treated according to Hunke et al. (2007) [63]. Solutions of Cyanine 5 (Cy5) in pure water (Fluka) were used as references for the calibration of the confocal microscope. The following filter sets were used: MBS: HFT 514/633, EF1: BP 655–710, EF: None, DBS: None. Out-of-focus fluorescence was rejected by a 90° beam splitter. The confocal detection volume was calculated to be approximately 0.32 fl at λ = 655 nm at a numerical aperture NA of 1.2. Variable concentrations of NBD94444–547 and NBD94 674–793 were measured at 8.0 mg/ml, respectively. The protein samples were prepared in 50 mM Tris/HCl (pH 7.5), 200 mM NaCl and 1.25 mM DTT as radical quencher. The two different oils, paraffin and silicone, were applied as barriers, separately and also as mixtures of these two oils.

Structure Determination

Crystals of selenomethionine (SeMet) substituted NBD94674–663 were prepared for MAD phasing. Initially, crystals were screened at beamline BL12B2 at SPring-8 using a Q4R detector. Native dataset and a complete MAD dataset from single crystals of SeMet incorporated NBD94674–663 were collected at 100 K on beamline BL26B2 at SPring-8 using a Mar CCD 225 detector. All the data were indexed, integrated and scaled using the program HKL2000 [38]. Selenium sites were identified and refined by the program SHELX [40], and density modification of the experimental map was performed with RESOLVE [46]. Manual model building and refinement of the structure was carried out in iterative cycles using COOT [47] and REFMAC5 [48]. The atomic coordinates and structure factors for the NBD94674–663 structure have been deposited in the Protein Data Bank under accession code 3HGF.

Supporting Information

Figure S1 Fluorescence correlation spectroscopy studies of NBD94444–547. (A) Normalized autocorrelation functions of MgADP ATTO-647N obtained by increasing the quantity of NBD94444–547 (increased protein concentration from left to right).
Effect of increased NBD-Cl concentration of MgATP ATTO-647N bound to NBD94566–663 shown as normalized autocorrelation functions (increased effector concentration from right to left). Found at: doi:10.1371/journal.pone.0009146.s003 (0.46 MB DOC)

Figure S2 Crystal forms of NBD94566–663. Crystals of selenomethionine substituted NBD94566–663 (10 mg/ml) grown by vapor diffusion using 35% (v/v) 2-methyl-2,4-pentanediol as precipitant, acetate pH 4.3 and 1 mM tris-2-carboxyethyl-phosphine (A). The crystal size and quality has been improved by controlling the rate of vapour diffusion by the introduction of an oil barrier over the reservoir of a vapour-diffusion trial (B). Found at: doi:10.1371/journal.pone.0009146.s002 (3.00 MB DOC)

Figure S3 Simulated annealing omit map for chain A of the NBD94566–663 structure.

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Author Contributions

Conceived and designed the experiments: AG PRP GG. Performed the experiments: AG MSSM AMB CH. Analyzed the data: AG MSSM CH. Contributed reagents/materials/analysis tools: AG. Wrote the paper: AG MSSM PRP GG. X-ray data collection of protein crystal: JJ.
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