Molecular Characterization of NAD⁺-Dependent DNA Ligase from Wolbachia Endosymbiont of Lymphatic Filarial Parasite Brugia malayi

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Abstract

The lymphatic filarial parasite, Brugia malayi contains Wolbachia endobacteria that are essential for development, viability and fertility of the parasite. Therefore, wolbachial proteins have been currently seen as the potential antifilarial drug targets. NAD⁺-dependent DNA ligase is characterized as a promising drug target in several organisms due to its crucial, indispensable role in DNA replication, recombination and DNA repair. We report here the cloning, expression and purification of NAD⁺-dependent DNA ligase of Wolbachia endosymbiont of B. malayi (wBm-LigA) for its molecular characterization. wBm-LigA has all the domains that are present in nearly all the eubacterial NAD⁺-dependent DNA ligases such as N-terminal adenylation domain, OB fold, helix-hairpin-helix (HhH) and BRCT domain except zinc-binding tetracysteine domain. The purified recombinant protein (683-amino acid) was found to be biochemically active and was present in its native form as revealed by the circular dichroism and fluorescence spectra. The purified recombinant enzyme was able to catalyze intramolecular strand joining on a nicked DNA as well as intermolecular joining of the cohesive ends of BstEII restricted lambda DNA in an in vitro assay. The enzyme was localized in the various life-stages of B. malayi parasites by immunoblotting and high enzyme expression was observed in Wolbachia within B. malayi microfilariae and female adult parasites along the hypodermal chords and in the gravid portion as evident by the confocal microscopy. Ours is the first report on this enzyme of Wolbachia and these findings would assist in validating the antifilarial drug target potential of wBm-LigA in future studies.

Introduction

More than 1 billion people in 81 countries worldwide are threatened by lymphatic filariasis (LF). The current antifilarial treatments e.g. diethylcarbamazine, ivermectin and albendazole interrupt the cycle of transmission of the causative filarial parasites Wuchereria bancrofti, Brugia malayi and B. timori, by predominantly killing the microfilariae (mf). However, a little adulticidal effect allows adult worms to survive in human hosts for up to decades. The current treatments have to be administered annually on a community wide basis for several years to break transmission of infection and this may instigate emergence of drug resistance. There is an urgent need to develop novel drug target/s particularly pertaining to adult parasites. With the discovery of intracellular endosymbiont alphaproteobacteria, Wolbachia, and the experimental evidence showing its indispensable role in parasite survival, development and embryogenesis [1,2], makes Wolbachia an attractive antifilarial drug target. Furthermore, Wolbachia have been identified as a major contributor of the inflammatory host pathology and severe adverse reactions to antifilarial chemotherapy. Therefore, characterization of novel proteins of Wolbachia is a prerequisite to identify the factors responsible for pathogenesis of the disease, understanding the host-parasite interactions and developing new protein/enzyme targets for antifilarial chemo-thrapy. A few proteins of Wolbachia have been identified and their functions are being investigated such as Wolbachia surface protein (WSP), heat shock protein 60 (HSP60), independent phosphoglyc-erate mutase (iPGM) and the enzymes involved in Heme biosynthetic pathway. The sequenced genome of the Wolbachia endosymbiont of B. malayi (wBm) offers an unprecedented opportunity to identify new Wolbachia drug targets [3]. DNA ligases are indispensable for many fundamental processes in DNA metabolism including the linkage of Okazaki fragments during replication, recombination processes and repair pathways requiring resynthesis of DNA [4,5]. Their crucial function is emphasized by the fact that the eukaryotic cells contain several isoenzymes and that the viruses encode their own ligases [6]. They are grouped into two families depending on their cofactor requirement, one is NAD⁺-dependent DNA ligase requiring NAD⁺ as a cofactor (present in bacteria and eukaryotic viruses) and the other is ATP-dependent DNA ligase requiring ATP as nucleotide cofactor (present in eukaryotes) [7,8]. NAD⁺-dependent DNA ligases have been characterized in several organisms such as Ferroplasma [9], Mycobacterium tuberculosis [10], Escherichia coli [11], Ansera moori entomopoxvirus [12] and Staphylococcus aureus [13]. Due to its important role in nuclear metabolism, this enzyme is being exploited as a potential antibacterial drug target in M. tuberculosis,

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E. coli, Salmonella enterica serovar Typhimurium, Bacillus subtilis and S. aureus [14,15,16]. Therefore, an attempt was made to clone, overexpress and purify NAD⁺-dependent DNA ligase from Wolbachia endosymbiont of Brugia malayi (wBm-LigA) with an aim to functionally characterize it. To the best of our knowledge, this is the first ever report on the characterization of any nuclear metabolic enzyme from an obligate mutualist alphaproteobacterium, Wolbachia of filarial parasite.

Results

~75 kDa recombinant wBm-ligA was over-expressed and purified

wBm-ligA gene was amplified from the genomic DNA of adult B.malayi worms and the amplified product was successfully cloned in pTZ57R/T vector for maintenance. Further, wBm-ligA gene was subcloned in pET28a(+) and confirmed by restriction digestion of recombinant construct of pET28a-ligA. wBm-LigA was expressed as a ~75 kDa protein with (His)₆-tag fused at the N and C-terminus in Rosetta E. coli cells (Figure 1A). The recombinant protein was obtained in the soluble form and purified by affinity chromatography on a Ni-NiNitrilotriacetic agarose column (Ni-NTA). The recombinant protein was eluted at 250 mM imidazole (Figure 1B) and finally confirmed with anti-His antibody in the Western blot (Figure 1C).

Conserved domain architecture of wBm-LigA showing phylogenetic divergence from eukaryotic DNA ligases

The Clustal alignment of amino acid sequences of NAD⁺-dependent DNA ligase from an array of prokaryotic species including alpha, beta, gamma proteobacteria and cyanobacteria indicated that the wBm-LigA was 79% identical, 89% similar to DNA ligase of Wolbachia endosymbiont of Drosophila melanogaster; 79% identical, 90% similar to Wolbachia endosymbiont of D. simulans; 78% identical, 88% similar to Wolbachia endosymbiont of Culex quinquefasciatus; 45% identical, 62% similar to DNA ligase of Rickettsia rickettsii; 42% identical, 59% similar to Rhizobium leguminosarum; 40% identical, 54% similar to Bordetella pertussis; 40% identical, 58% similar to E. coli and 37% identical, 55% similar to Pochonococcus marinus (Figure 2). The conserved domain architecture, unique to DNA ligase superfamily, was identified in wBm-LigA. Residues 9 to 304 constitute an adenylation domain (LIGANc) carrying KXDG catalytic motif (114–117amino acid), involved in adenylate reaction. C-terminal region has an OB fold domain (amino acid 307–387), HisH (helix hairpin helix motif) and BRCT domain (amino acid 609–671), all characteristics of the NAD⁺-dependent DNA ligase superfamily (Figure 3A).

Analysis of the phylogenetic tree (Figure 3B) showed that wBm-LigA forms a discrete cluster A with closely related NAD⁺-dependent DNA ligase of all the prokaryotic bacteria which is subdivided into two branches, the first one constituting the subcluster A1 comprised of Nitrosomonas europaea, Bordetella pertussis, Gamma proteobacteria, Pseudomonas aeruginosa, E. coli, M. tuberculosis, Frankia, Thermus thermophilus, Pseudococcus marinus, Mycosphaera mycoides, Bacillus subtilis, Bodo trichophyton, Rhizobium leguminosarum, Agrobacterium vitis and the second one (subcluster A2) comprising wBm-LigA along with the DNA ligase of Wolbachia endosymbiont present in the insects such as Drosophila melanogaster, D. simulans, Culex quinquefasciatus, other alpha proteobacteria such as Wolbachia pipientis, Anaplasma centrale, Ehrlichia chaffensis and Rickettsia rickettsii. All the eukaryotic DNA ligases which are ATP dependent viz. Homo sapiens, Mus musculus, B. malayi, Plasmodium falciparum, P. knowlesi and Saccharomyces cerevisiae are present in a separate cluster B indicating significant divergence between wBm-LigA and B. malayi or H. sapiens.

Structural analysis of recombinant wBm-ligA, circular dichroism and fluorescence spectra

The circular dichroism (CD) spectra revealed 2 negative bands at 207 nm and 220 nm, and marginally positive ellipticity at 196 nm (Figure 4A). K2D2 web server was employed to estimate the ratio of secondary structure elements. K2D2 server accepts a CD spectrum as an input and gives out the estimated secondary structure content of the corresponding protein as an output. The deconvolution of the CD spectra with K2D2 indicated a secondary structure ratio of 36.44% α-helix, 12.07% β strands that were consistent with the predicted values (GOR1V method) of 44.51% α-helix and 14.20% β strands. The 3D structure of wBm-LigA was deduced from 3D-JIGSAW automated web server and is represented as a ribbon diagram in Figure 4B. This model indicates a total of 26 α-helices and 29 β strands including 3 β sheets and 69 turns. The conserved active site residues KXDG

Figure 1. Overexpression, purification and Western blot analysis of recombinant wBm-LigA. (A) Overexpression of wBm-LigA in Rosetta E. coli cells. lane M, standard protein molecular weight marker; lane1, total proteins extracts isolated from the uninduced E. coli cells; lanes 2–4, total proteins extracts isolated from the E. coli cells induced with 0.2 mM, 0.5 mM and 1.0 mM of isopropyl-1-thio-β-D-galactopyranoside (IPTG). ~20 µg of proteins were loaded per lane. (B) Purification of wBm-LigA by affinity chromatography using nickel column. lane 1, soluble E. coli proteins following induction with 1.0 mM IPTG at 25°C for 5 h; lane 2, flow through from the nickel column; lanes 3–6, washes from column prior to elution; lane 7, elution of His-tagged purified recombinant wBm-LigA at 250 mM imidazole concentration. 5 µg of recombinant protein was loaded onto the gel; lane M, standard protein molecular weight marker. (C) Western blot analysis. lane 1, confirmation of His-tagged wBm-LigA expression with reaction of 20 µg of recombinant protein with anti-His antibody; lane 2, reaction of 10 µg of wBm-LigA protein with anti-His antibody; lane M, standard protein molecular weight marker.

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(amino acid residues 114–117) are also displayed in the figure. The fluorescence emission spectrum of the purified wBm-LigA protein in standard buffer at 25°C showed maximum emission at 333 nm when excited at 280 nm. Guanidium hydrochloride induced decrease in the intrinsic fluorescence intensity and a red-shift in the emission maximum wavelength, indicating denaturation of the protein (Figure 4C).

**Figure 2. Clustal alignment of amino acid sequence of wBm-LigA with other prokaryotic bacteria.** Amino acid sequence of wBm-LigA (GenBank:AAW71136.1) was aligned with amino acid sequences of DNA ligases of other prokaryotic bacteria including alphaproteobacteria, *Rhizobium leguminosarum* (R.leg, reference sequence YP_002282076.1), *Rickettsia rickettsii* (R.ric, GenBank:ABY73051.1), *Wolbachia* endosymbiont of *Drosophila melanogaster* (WDm, reference sequence NP_966531.1), *Drosophila simulans* (WDs, reference sequence ZP_00372215.1), *Culex quinquefasciatus* (WCq, reference sequence accession YP_001975247.1); Betaproteobacteria, *Bordetella pertussis* (B.per, reference sequence NP_862073.1); Gamma proteobacteria, *Escherichia coli* (E.col, GenBank:AA81395.1) and cyanobacteria *Prochlorococcus marinus* (P.mar, reference sequence NP_876232.1). Residues that are identical are highlighted in black, and the conserved amino acid changes are outlined in gray rectangular boxes.

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**(wBm-LigA demonstrates the nick closing activity**

A concentration-response curve of wBm-LigA nick activity was generated to show the proportionality of the enzyme concentration to the nicked substrate. Incubation of an equimolar ratio of enzyme and substrate caused rapid product formation and quick saturation. Decreasing the enzyme/substrate ratio to 0.04 or 0.02, produced slower product formation (Figure 5A). Using varying
concentration of the nicked substrate, a Km of 0.193 μM was obtained for wBm-LigA. The very first step in ligation reaction is the adenylation of the ligase enzyme requiring AMP moiety that is provided by the NAD⁺ cofactor. All the prokaryotic NAD⁺-dependent DNA ligase bacteria formed a discrete cluster A which is divided into two branches, the first one comprising the subcluster A1 constituting Nitrosomonas europaea (reference sequence NP_841783.1), Bordetella pertussis (reference sequence NP_882073.1), Gamma proteobacteria (reference sequence ZP_05126247.1), Pseudomonas aeruginosa (reference sequence YP_191721.1), Escherichia coli (GenBank:AA81395.1), Mycobacterium tuberculosis (reference sequence ZP_7441467.1), Frankia (reference sequence ZP_06413452.1), Thermoanaerobacterium (GenBank:AAE71598.1), Pseudomonas aeruginosa (reference sequence NG_017030.1), Mycoplasma mycoides (reference sequence NP_388544.1), Bradyrhizobium japonicum (GenBank:AA51685.1), Rhiobium leguminosarum (NCBI reference sequence YP_002812076.1), Agrobacterium vitis (reference sequence ZP_0022650080.1) and the second one (subcluster A2) comprising the DNA ligase of Wolbachia endosymbiont present in Brugia malayi (GenBank:AAW71136.1) and insects as Drosophila melanogaster (reference sequence NP_966331.1), Drosophila simulans (reference sequence ZP_00372215.1) and Culex quinquefasciatus (reference sequence YP_001975247.1), Wolbachia pipientis (GenBank:AY81980.1), Anaplasma centrale (reference sequence NP_00328904.1), Ehrlichia chaffeensis (NCBI reference sequence YP_507122.1) and Rickettsia rickettsii (GenBank:ABY73051.1). All the euukaryotic DNA ligase viz. Homo sapiens (reference sequence NP_002225.1), Mus musculus (reference sequence NP_001186239.1), Brugia malayi (NCBI reference sequence YP_001896804.1), Plasmodium falciparum (GenBank:AN6156.1), Plasmodium knowlesi (reference sequence XP_002261933.1) and Saccharomyces cerevisiae (GenBank:CA91982.1) are present in discrete cluster B indicating Wolbachia endosymbiont are distantly related to the ATP DNA ligase of either its host parasite B. malayi or H.sapiens.

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Maximal activity occurred at 5 mM NaCl, however, further increase in the concentration completely inhibited this activity viz. no activity could be observed at 100 mM NaCl. In presence of KCl, the maximum activity was achieved at 25 mM concentration showing a sharp decline thereafter. The activity increased with the increasing concentrations of NH4Cl up to 10 mM, beyond that the activity decreased (Figure 5F). The enzyme activity increased in the presence of low concentrations of EDTA, however at higher concentration beyond 30 mM, the DNA ligase activity was inhibited. (Figure 5G).

**wBm-LigA demonstrates the cohesive end ligation activity**

wBm-LigA was able to catalyze the ligation of cohesive ends of lambda DNA digested with BstEII enzyme. Optimum temperature for cohesive end ligation was 25°C similar to the nick closing activity (Figure 6A). Optimal pH range was 7.0–8.0 with maximum activity at pH 7.5 (Figure 6B). With increasing enzyme concentration, the product formation also increased initially and then it became constant (Figure 6C). The monovalent as well as divalent cations were required for the cohesive end ligation activity. The maximum enzymatic activity was obtained in the presence of 10 mM MgCl2 whereas 10 mM concentrations of MnCl2 and CaCl2 resulted into 90% and 30% product formation respectively. The ligation activity was not observed in presence of Ni2+ and Co2+ (Figure 7A). The ligation activity was maximum in the presence of 25 mM NaCl, 100 mM KCl or 50 mM NH4Cl (Figures 7B, 7C and 7D). The increasing concentrations of NAD+ showed a rise in wBm-LigA cohesive end ligation activity up to 20 μM concentration (Figure 7E). Time course of ligation of BstEII digested lambda DNA fragments into the product revealed that product quantity was maximum by 45 min and longer incubations periods did not result in an increase in product amount (Figure 7F).

**wBm-LigA is present in all the major life-cycle stages of *B. malayi***

wBm-LigA antiserum generated in BALB/c mice reacted in the blot with endogenous wBm-LigA in the crude extracts prepared from the three major life-stages of *B. malayi* viz. infective larve (L3), adult worms and mf, showing a characteristic band at ~73 kDa. The purified recombinant wBm-LigA which acted as positive control also showed reaction with anti-wBm-LigA antibody giving a characteristic band at ~75 kDa including His-tag at C and N terminus (Figure 8A).

The confocal microscopy localized the endogenous wBm-LigA in adult *B. malayi* and mf

The polyclonal antibody raised in BALB/c mice against wBm-LigA reacted with the endogenous wBm-LigA protein inside the adult female parasite and mf in confocal microscopy. In the case of adult female worm, high level of wBm-LigA expression could be detected inside the uteri in the developing embryos, stretched mf as well as hypodermal lateral chords indicated by green fluorescence (fluorescein isothiocyanate, FITC) (Figures 8C, 9A, 9B and 9C). In contrast, adult worm treated with

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Figure 4. CD Spectra, 3D-JIGSAW model and fluorescence spectra. (A) CD spectra of recombinant wBm-LigA, recorded in far UV region. CD spectra gave 2 negative bands at 207 nm and 220 nm, and slight positive ellipticity at 196 nm. (B) Ribbon diagram of 3D structure model of wBm-LigA made by 3D-JIGSAW and visualized by Rasmol. α Helices is represented in green; β strand in blue and turns in brown. Model indicated a total number of 26 α-helices and 29 β-strands including 3 β sheets and 69 turns. Conserved active site residues KXDG (amino acid residues 114–117) are also indicated. (C) Fluorescence emission spectrum of wBm-LigA recorded between 300 nm and 450 nm wavelength after excitation at a 280 nm. Maximum peak was generated at 333 nm indicating folded form of wBm-LigA that was denatured by Gu-HCl resulting in red shift in spectra and decrease in the fluorescence intensity.

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preimmune BALB/c serum did not reveal any green signal for wBm-LigA presence in adult worms (Figure 8B). 4',6'-diamidino-2-phenylindole (DAPI, blue) stained Wolbachia and B. malayi DNA (Figures 8B, 8C; 9A, 9B and 9C). wBm-LigA was also conspicuous as green fluorescence in isolated mf (Figure 10B and 10B') in contrast to mf, treated with preimmune serum (Figure 10F and 10F').

**Discussion**

The wolbachial growth and multiplication is essential for the survival and fertility of its host, B. malayi. DNA ligases are essential actors in DNA replication by virtue of their ability to join the Okazaki fragments together to form a continuous lagging strand. Therefore, DNA ligase of Wolbachia plays an indispensable role in the intracellular bacterial division inside the B. malayi parasite. It is also involved in the maintenance of genetic stability during DNA damage and in bacterial recombination. The characterization of wBm-LigA is therefore an important step towards understanding the reactions involved in DNA repair and replication at molecular level in this obligate mutualist (wBm) and conclusion drawn from the following study would further assist in validating its drug target potential.

The indispensable nature of this enzyme is attributed to the fact that it is highly conserved across the different bacterial species as evident by the protein sequence alignment that indicated considerable sequence similarity between wBm-LigA and other bacterial DNA ligases including alpha, beta, gamma bacteria as well as blue-green bacteria. DNA ligases of alphaproteobacteria formed a distinct clade in phylogenetic tree which was also strongly supported by bootstrap scores, leads us to speculate that DNA ligase of alpha proteobacteria are evolutionary closely related to each other and they may have followed a different evolutionary path from rest of other bacteria. The possible reason...
could be the lifestyles in which alphaproteobacteria have to survive and divide. It is already well established that the alphaproteobacteria constitute an excellent model system to explore how the bacterial genomes have evolved and how the genomic features are related to environmental adaptation [17]. Earlier studies also demonstrated that $\alpha$-proteobacteria can be clearly distinguished from all other bacteria based upon a large number of molecular characteristic and therefore should be recognized as a main group or phylum within bacteria, rather than as a subdivision or class of the proteobacteria [18]. A significant divergence of $\omega$Bm-LigA from eukaryotic DNA ligases of $H$. sapiens, $M$. musculus, $B$. malayi, $S$. cerevisae and $P$. species which are ATP dependent, indicates the potential of $\omega$Bm-LigA as an attractive antifilarial drug target.

$\omega$Bm-LigA has a modular structure typical of NAD$^+$-dependent DNA ligases and each step of ligation pathway depends upon a different subset of these modules. However, tetracysteine zinc finger domain which is important in the formation of stable complex with DNA substrate in ligation reaction is not present in $\omega$Bm-LigA. This feature is in contrast to that of $R$. rickettsii as well as $E$. coli DNA ligase which possess C4 type Zn domain for DNA binding. It is possible that being an endosymbiont in nature, $W$oblachia has retained only the essential domains and deleted those, whose function could be provided in trans by some other factors conceivably by undefined region of $\omega$Bm-LigA or might be HhH motif and BRCT domain could be solely responsible for stable DNA binding. The conserved KXDG catalytic site in the N terminal domain of $\omega$Bm-LigA is formed by a single $\beta$ strand and the random coils while the C-terminal region of $\omega$Bm-LigA is predominantly composed of the alpha helices as evident by the homology model. Published reports implicate C-terminal domain of NAD$^+$-dependent DNA ligase in recognition of DNA substrate principally through the protein-protein interaction [19,20]. Besides, the BRCT domain also acts as a signal transducer transmitting the signal from DNA damage sensors to the other components of the DNA damage-responsive checkpoint machinery via specific protein–protein interaction in DNA repair system [21]. Thus, homology model structure clearly confirms that C-terminal region acts through protein-protein interaction since alpha helices are known to play a prime role in protein–protein interaction as well as in DNA binding. Thus, the presence of all the characteristic domains of DNA ligase superfamily in DNA ligase of endosymbiont $W$oblachia demonstrates the remarkable ability of $W$oblachia for self perpetuance in its host.

Immunoblotting studies with crude extract of adult worm, L3 and mf indicated the presence of $\omega$Bm-LigA in all the major life-

Figure 6. Cohesive end ligation activity of $\omega$Bm-DNA ligase at various temperature, pH and enzyme concentration. (A) Temperature dependence of cohesive end ligation activity of $\omega$Bm-LigA. Lane 1, 2, 3, 4, 5, 6 showed ligation at 10°C, 25°C, 30°C, 35°C, 40°C and 45°C temperature res.; lane C, control in which no enzyme was added. Enzyme showed maximum ligation at 25°C. Quantitation of the data is shown below as a bar diagram. (B) pH dependence on the cohesive end ligation. Lane 1, 2, 3, 4, 5 represents ligation at 6.5, 7.0, 7.5, 8.0, 8.5 pH res.; lane C, control. Maximum ligation activity was observed at 7.5 pH. Quantitation of the data is shown below as a bar diagram. (C) (Lanes 1–12), effect of enzyme concentration (0–25nM) on cohesive end ligation; Lane C, control. Quantitation of the data is shown to the right as a bar diagram. Ligation of cohesive ends of fragment 1 and fragment 4 of lambda DNA digested with BstEII into 14 kb ligated product was analyzed on a 1.0% agarose gel. Activity values were calculated by measuring the net intensity of the product band (indicated as 1+4 in the figures) relative to a non-substrate fragment (fragment 7).

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stages of *B. malayi*. In *vivo* localization of *wBm-LigA* by immunofluorescent microscopy using anti-*wBm-LigA* antibody demonstrated the presence of enzyme in the hypodermal chord of female adult worm as well as inside the uteri of female parasite i.e. in the developing embryos and mf. The enzyme showed its presence also in isolated mf. The ligA gene expression in all the major life-stages of *B. malayi* further indicated the essentiality of this enzyme for the parasite. In several organisms, DNA ligases have been reported to be essential [13,15,16], and therefore there is great likelihood that it is also essential in *Wolbachia* thereby suggesting its significance as an antifilarial drug target.

The full-length *wBm-LigA* expressed as the recombinant protein in *E. coli* was enzymatically active when quantitated in an *in vitro* ligation assay using a fluorescent labeled substrate in nick closing assay and BstEII digested lambda DNA as a substrate in the cohesive end ligation assay. The biochemical characterization established that the *wBm-LigA* requires an optimal temperature 25°C, optimal pH range 7.5–8.5 for nick closing activity and pH 7.0–8.0 for cohesive end ligation activity displaying its strict pH dependent nature. The data reveal that *wBm-LigA* fulfills the requirement of a divalent metal cofactor by magnesium cations which are most often associated with ligation activity. It is worth mentioning that *wBm-LigA* was able to catalyze nick sealing as well as cohesive end ligation even in the presence of *Mn*²⁺ as well *Ca*²⁺, since *Ca*²⁺ has a larger ionic radius than *Mg*²⁺ or *Mn*²⁺ this suggests more flexibility within the “catalytic-core” of *wBm-LigA* which is evident from 3D homology model where a large portion of catalytic site is formed by the random coils. *wBm-LigA* possesses high affinity for the nicked DNA substrate as reflected by its lower Km (0.193 mM) than that of *E. coli* DNA ligase (0.702 mM) [22]. Lower Km value for the NAD⁺ cofactor could be attributed to the fact that *wBm* lacks complete pathways for de novo biosynthesis of cofactor NAD⁺ which makes *wBm-LigA*, dependent on the host for the supply of the precursors [3]. Reactions carried out at increasing concentrations of EDTA and fixed concentration of *Mg*²⁺ indicated increased enzymatic activity with concentrations below 20 mM of EDTA. The reason could be that at these concentrations of EDTA, the *Mg*²⁺ that is available to *wBm-LigA* is optimal for its activity. The other point is that enzymatic activity remain high even when EDTA is at equimolar concentration of *Mg*²⁺. This could be hypothesized that EDTA although chelates *Mg*²⁺, but due to high affinity of *wBm-LigA* for *Mg*⁺, available *Mg*²⁺ to *wBm-LigA* is sufficient for the increased activity. However, EDTA at 30 mM or higher concentration could efficiently compete and strip the *Mg*²⁺ resulting into inhibition of *wBm-LigA* activity. There are reports...
where small amount of EDTA are being used in the ligation reaction of DNA ligases \cite{13, 23, 24} but their effect on ligation activity is not demonstrated. However, inhibition of DNA ligase by employing high concentration of EDTA (2 or more times than the concentration of Mg\(^{2+}\)) is reported for many DNA ligases \cite{25–27}.

The CD spectra of \(wBm\)-LigA in far-UV range showed the presence of a typical \(\alpha/\beta\) type secondary structure. The alpha helix gives negative band at 208 and 222 nm, while the presence of \(\beta\) sheet gives negative band at 218 nm causing a shift in the wavelength at 222 nm of the helix structure. This was further confirmed through the CD spectra deconvolution with K2D2. K2D2 is a webserver that estimates the secondary structure content of a given protein from its CD spectra. The fluorescence spectra clearly indicated that the purified recombinant enzyme was properly folded and was present in its native form. A peak at 333 nm which is red shifted after adding guanidine HCl demonstrated that the three tryptophan residues were buried in hydrophobic environment of the protein and become exposed to more polar environment once the native 3D structure is disrupted via denaturation that led to the decreased intrinsic fluorescence intensity \cite{28}.

Ours is the first report on the molecular characterization of NAD\(^{+}\)-dependent DNA ligase of \(Wolbachia\) which is essential for DNA replication, repair and recombination. The presence of this enzyme in all the major life cycle stages of \(B. malayi\), high level of \textit{in vivo} endogenous expression and conserved domain organization indicated its imperative role in the biology of filarial endosymbiont. Further, profound divergence from the eukaryotic DNA ligases was also found. Since \(Wolbachia\) play an important role in the development, viability, fertility of filarial nematodes, and they are also responsible for filarial pathology, \(wBm\)-LigA presents an attractive antifilarial drug target. The specific inhibitors of this enzyme are known and new chemical structures may be synthesized which will help in validating \(wBm\)-LigA as a candidate drug target. The expression and characterization of this protein represents a critical step towards understanding the molecular trappings underlying \(Wolbachia\) maintenance in filarial parasite thereby providing more insight into mutualistic interaction between \(Wolbachia\) and its host, \(B. malayi\).

\textbf{Materials and Methods}

\textbf{Ethics statement}

The animals used in the study were maintained at Laboratory Animal Division of Central Drug Research Institute (CDRI), Lucknow, under pathogen-free conditions. All the animals,
Cloning of full length NAD+ -dependent DNA ligase gene

The full length open reading frame of zm-ligA gene (2052 bp long) was amplified from the genomic DNA isolated from B. malayi adult worms [29] by PCR using gene specific forward primer 5'-GAATTCCATGGCTGATTTGGAAAAGATG-3' and reverse primer 5'-AAGCTTTTCGGACATTCCGAATTTAT-3'. The forward primer contained EcoRI site (underlined) and the reverse primer contained HindIII site (underlined). The amplification of ligA gene was carried out using 1 μM of each primers (Sigma, USA), 200 μM of each dNTPs (Invitrogen, USA), 0.5 unit Taq DNA polymerase (Invitrogen, USA), 1xPCR buffer (Invitrogen, USA), and 1.5 μM of MgCl2 (Invitrogen, USA) under the conditions of initial denaturation at 94°C for 10 min, followed by 29 cycles at 94°C for 45 sec, 65°C for 1 min, 72°C for 2 min, and 1 cycle at 72°C for 10 min. The adult worms were obtained from the peritoneal cavity of jirds (Meriones unguiculatus) infected 3–4 months earlier by inoculation of 150–200 L3 intraperitoneally. The amplified gene product was cloned in pTZ57R/T vector as per manufacturer’s instructions (Invitrogen, USA) and transformed into competent DH5α E. coli cells. The gene constructs were screened for the presence of recombinant plasmid carrying zm-ligA insert by gene-specific PCR, followed by double digestion of the recombinant plasmid with EcoRI and HindIII (Fermentas, USA) for subcloning in pET 28a(+) vector. The plasmid construct in pET28a(+) was confirmed by further restriction digestion.

Protein expression, purification and localization of the recombinant protein (zm-ligA) in Western blot

The construct of pET28a(+) -zm-ligA was transformed into various E. coli strains viz. BL21(DE3), BL21(DE3)pLysS, Rosetta and C41 (Novagen, USA), however, the recombinant protein was expressed only in Rosetta cells. A single colony of Rosetta cells containing pET construct was inoculated in 10 ml of Lysogeny Broth (LB) medium fortified with 50 μg/ml kanamycin and the cells were grown overnight (O/N) at 37°C. 5 ml of this O/N grown culture was inoculated in 500 ml of LB medium containing kanamycin (50 μg/ml). The culture was grown in a shaker incubator with constant stirring till A600 reached ~0.6. The culture was induced with 1.0 mM isopropyl-β-D-thiogalactopyranoside (Sigma, USA) and incubated at 25°C for 5 h. The bacterial cells were pelleted at 3000xg for 10 min at 4°C and resuspended in Buffer A (20 mM Tris-Cl pH 8.0, 250 mM NaCl, 10 mM imidazole). The cells were lysed in the presence of lysozyme (1 mg/ml) and Triton X-100 (0.1%) and the lysate was sonicated to reduce viscosity. The soluble fraction obtained after centrifugation was applied over a Ni-NTA agarose column (Qiagen, USA) that was pre-equilibrated with Buffer A. The column was washed with Buffer B (20 mM Tris-Cl pH 8.0, 250 mM NaCl, 50 mM imidazole) and the recombinant protein was eluted with Buffer C (20 mM Tris-Cl pH 8.0, 250 mM NaCl, 250 mM imidazole). The recombinant zm-ligA protein appeared as a single band on a 10% sodium dodecyl sulfate (SDS) polyacrylamide gel electrophoresis (SDS-PAGE). The protein was dialyzed against Buffer B (20 mM Tris-Cl pH 8.0, 250 mM NaCl, imidazole 50 mM) and subsequently
against Buffer D (Tris-Cl 20 mM, 250 mM NaCl). The protein content was estimated with Bradford reagent using Bovine serum albumin (BSA) as standard. Approximately, 2 mg was recovered from 1 liter of the bacterial culture. For the Western blot analysis, purified recombinant protein from the SDS-PAGE gel was transferred to a nitrocellulose membrane. The membrane was blocked with 3% BSA for 1 h and incubated at room temperature (RT) with 1:2500 dilution of mouse anti-His antibody (Sigma, USA), re-incubated with 1:10,000 dilution of goat anti-mouse IgG-HRP antibody (Sigma, USA) for 1 h at RT and the blot was developed with the substrate 3,3'-diaminobenzidine tetrahydrochloride (DAB, Sigma, USA) [30].

**Phylogenetic analysis**

Protein sequences of DNA ligases from the different organisms available on NCBI database were aligned using ClustalW software. The phylogenetic trees were constructed based on the protein sequence alignments with Maximum Likelihood (ML) Method using PROML programs of PHYLIP package version 3.69 with global rearrangements and randomized input order options. Bootstrap re-sampling was done with 100 replicates using the Seqboot program. Finally, the results from random datasets were summarized by consense that makes consensus trees by the Majority-rule consensus tree method. The phylogenetic trees were visualized using the Tree Drawing Tool-Drawgram.

**Structural Analysis**

The CD spectra were recorded on JASCO J810 spectropolarimeter equipped with pelier temperature controller system in a 0.2 cm cell at 25°C. The measurements were carried out in 50 mM sodium phosphate buffer, pH 8.0 at a final protein concentration of 10 μM in the far-UV range (195 nm–240 nm). The ellipticity was reported as the mean residual ellipticity \([\theta](10^3 \text{deg.cm}^2\text{dmol}^{-1})\). K2D2 web server (http://www.ogic.ca/projects/k2d2/) was employed to estimate the ratio of secondary structure elements in the CD spectra [31]. On the other hand, the 2D structure prediction was also carried out using the GOR1V method (Garnier–Osguthorpe–Robson) available online at http://abs.cit.nih.gov/gor/. The 3D model of \(\omega\)Bm-LigA was predicted by amino acid homology modeling using 3D-JIGSAW automated web server (http://bmm.cancerresearchuk.org/3djigsaw/) [32] and viewed by Rasmol (http://www.openrasmol.org/) [33].

**Fluorescence spectroscopy**

The fluorescence spectra were recorded on a fluorescence spectrometer (Perkin Elmer LS50b) using an excitation wavelength of 280 nm and an emission wavelength in the range of 300–450 nm. The path length of quartz cell was 5 mm. The measurements were carried out at a final protein concentration of 1 μM in 50 mM sodium phosphate buffer, pH 8.0 at 25°C and \(\omega\)Bm-LigA was denatured with 6 M guanidine hydrochloride which served as control. For guanidine hydrochloride denaturation, the recombi-
nant protein was dissolved in sodium phosphate buffer (50 mM, pH 8.0), incubated at 25 °C O/N before measurements were made. This incubation time was sufficient for the reaction to achieve equilibrium under all denaturant concentrations.

Nick closing activity and biochemical characterization of wBm-LigA

Nick closing activity of αBm-LigA was determined by the sealed duplex assay as described earlier [24] with minor modifications. This ligation assay format was based on ligation of a fluorescein labeled oligonucleotide to a biotinylated oligonucleotide within a double strand duplex hybridized on a streptavidin-coated microtitre plate. The amount of fluorescence captured following denaturation of DNA duplex quantifies a successful ligation. Briefly, the three oligonucleotides viz. 5'-fluorescein AAAATGACCCC C-3', 5'-CCAGACACCGTCG-biotin-3' and 5'-GACGTTGCTGGGGGGGTCAATT'TT-3' (Eurogentec, Germany) were suspended in Buffer H [5xSSC (0.75 M sodium chloride, 0.075 M sodium citrate solution), 0.1% BSA, 0.1% Triton X-100] and hybridized on Strepta Well HighBind plates (Thermo-scientific, USA) for 90 min at 20 °C to produce nicked substrate. Wells were washed with 400 μl of ligase buffer (20 mM Tris-HCl, pH 8.0, 10 mM (NH4)2SO4, 10 mM MgCl2, 1.2 mM EDTA, and 1 mM dithioerythritol) and the ligation was performed for 60 min at 25 °C in 200 μl of ligase buffer to which 5 nM of αBm-LigA and 26 μM of NAD+ was added. This was followed by denaturation in 0.5% SDS, 0.5 M NaOH for 1 min. The fluorescence was measured at an excitation wavelength of 485 nm and an emission wavelength of 525 nm after washing the wells twice with 400 μl of TBS-T (Tris-buffer saline containing 0.05% Tween 20, and 0.1% BSA. Substrate concentration dependent nick activity curve of αBm-LigA was generated for determining the steady-state Km for nicked substrate by measuring the formation of fluorescein labeled product at an increasing substrate concentration. To investigate the correlation between the concentration of enzyme and the substrate, different concentration of enzyme were mixed with a fixed amount of substrate such that enzyme, substrate concentration was in the ratio of 1, 0.04 or 0.02. The pH optima of αBm-LigA was determined in Tris–HCl buffer at various pH range between 6.5 and 9.0 while the effect of temperature on the enzymatic activity was determined in Tris–HCl buffer at various pH range between 6.5 and 8.5, 100 mM KCl, 10 mM MgCl2, 26 μM NAD+, 10 mM DTT, 500 ng of BstEI digested lambda DNA and the reaction was performed at 25 °C for 60 min. The reaction was stopped by adding 4 μl of 50 mM EDTA followed by heating at 70 °C for 5 min and subsequent chilling on ice for 5 min. The mixture was run on 1.0% agarose gel prepared in 1xTris acetic EDTA (TAE) buffer at 100 V for 90 min. The activity values were calculated by measuring the net intensity of the product band ([+4] relative to a non-substrate fragment [−7]). The band intensities were determined by measuring UV fluorescence of the bands using the volume rectangle tool of the Quantity One version 4 software package of ChemiDoc system (BioRad, USA), after capturing the digital images of the agarose gel [34]. The activity was observed at varying temperature (10°C–45°C), pH (6.5–8.5), enzyme concentration (0–25 nM) and NAD+ cofactor. The effect of different salts viz. MgCl2, MnCl2, CaCl2, NiCl2 and CoCl2 (as the source of divalent cations); and NaCl, KCl and NH4Cl (as the source of monovalent cations) was also determined on cohesive end ligation activity of αBm-LigA. The time course of ligation of BstEII digested lambda DNA fragments into the product was observed by incubating the reaction mixture at different time intervals.

Generation of polyclonal antibody to wBm-LigA in BALB/c mice

Five eight-week-old male BALB/c mice were administered subcutaneously in the back region with 20 μg of αBm-LigA protein mixed with Freund’s complete adjuvant (FCA, Sigma, USA) in 100 μl volume. The animals further received two more booster doses of same amount of protein in Freund’s incomplete adjuvant (FIA, Sigma, USA) on day 15 and day 21 post first immunization. The blood was collected from the animals just before immunization on day 0 for isolation of preimmune serum and on day 30 to isolate anti-αBm-LigA serum.

Presence of endogenous wBm-LigA in adult worm, L3 and mf stages by immunoblotting

To find out the presence of αBm-LigA in various life-stages of B. malayi, the crude extracts derived from adult worms, mf and L3 were run on 10% SDS-PAGE followed by Western blotting. The target protein was recognized by immunoblotting using polyclonal antibody to αBm-LigA. For preparing the crude somatic extracts, ten adult worms (male-female), 1000 L3 and 1500 mf were homogenized in phosphate buffer saline (PBS) containing protease inhibitor cocktail (Sigma, USA), sonicated and supernatant was obtained after centrifugation at 12,000g for 30 min. The supernatant was mixed with an equal volume of 2x sample buffer [Bromophenol Blue (10 mg/ml), SDS (4.4%), Tris-Cl (0.5 M), β-mercaptoethanol (300 mM)] and heated at 95°C for 5 min followed by separation onto the 10% SDS-PAGE. The separated proteins were then transferred to nitrocellulose membrane. The remaining steps were same as described above for localization of His tagged αBm-LigA in Western blot except that the anti-αBm-LigA antibody was used as a primary antibody (1:500 dilution) instead of anti-His antibody. The recombinant αBm-LigA purified through Ni-NTA column served as a positive control.

Immunolocalization of endogenous wBm-LigA in B. malayi by confocal microscopy

To determine αBm-LigA distribution in the parasites by immunolocalisation, the adult female worm and mf were fixed overnight in 4% paraformaldehyde in M9 buffer (42 mM Na2HPO4, 22 mM KH2PO4, 86 mM NaCl, 1 mM MgSO4.7H2O, pH 7.0) at 4°C and processed as described earlier [35]. Anti-αBm-LigA antibody was used as a primary antibody (1:700) while FITC conjugated IgG (1:200 in 0.5% BSA) was used as a secondary antibody. The worms and mf were incubated with 4',6'-diamidino-2-phenylindole (DAPI, 100 ng/ml) for 3 min before final washing with M9 buffer for staining the DNA. Parasites were mounted on a glass slide in 90% glycerol and 10% PLOS ONE | www.plosone.org 12 July 2012 | Volume 7 | Issue 7 | e41113
p-phenylenediamine (Sigma, USA) in the PBS. The parasites were observed and the images were captured with a Zeiss LSM 510 META (Zeiss, Jena, Germany) confocal laser scanning microscope equipped with a plan-apochromat 63X oil objective and 40X objective. For excitation of FITC-conjugated secondary antibody, argon laser lines of 488 nm were used. DAPI was excited at 405 nm. As a negative control, the same procedure was executed after treating parasites with BALB/c preimmune serum.

Statistical analysis
Data were analysed with the help of statistical software GraphPad PRISM 3.0.

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Author Contributions
Conceived and designed the experiments: NS SMB. Performed the experiments: NS JKN. Analyzed the data: NS JKN. Wrote the paper: NS SMB.