1	Co-expression and purification of the RadA recombinase with the		
2	RadB paralog from <i>Haloferax volcanii</i> yields multimeric ring		
3	structures.		
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- 1 Abstract
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The present study demonstrates the successful co-expression of two RecA-like proteins, RadA 3 4 and RadB, from the halophilic eurvarchaeon, Haloferax volcanii (Hvo). A co-expression approach was used to improve the purification of RadA by providing its lone paralog, RadB, as a 5 6 binding partner. At present, structural and biochemical characterization of Hvo RadA is lacking 7 and the physical interaction of Hvo RadA and Hvo RadB has not been demonstrated under either in vitro or in vivo conditions. Here, we describe for the first time co-expression of Hvo RadA 8 with RadB and interaction of these proteins as a complex under in vitro conditions. Purification 9 procedures were performed under conditions of high salt concentration (>1 M sodium chloride) 10 to maintain the solubility of the proteins. Quantitative densitometry analysis of the co-expressed 11 and co-purified RadA-B complex estimated the ratio of RadA to RadB to be 4:1, which suggests 12 that the proteins interact with a specific stoichiometry. Based on a combination of analyses, 13 including size exclusion chromatography, Western blot and electron microscopy observations we 14 suggest that RadA multimerizes into a ring-like structure in the absence of DNA and nucleoside 15 cofactor. 16 17 18 **Key Words:** 19 Homologous recombination, RadA, Haloferax volcanii, Co-expression, Halophilic protein 20 purification. 21 22 23 24

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- 1 Introduction
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Homologous recombination is one of the principal pathways to repair deleterious DNA damage 3 4 such as double-strand breaks. The basic mechanism of homologous recombination, comprising homologous base-pairing and a strand exchange reaction, is similar in all forms of life, however 5 it varies in complexity and enzymology. Various enzymes participate in the recombination 6 7 process, including the key recombinase which belongs to the RecA family of proteins [1]. These enzymes are well conserved in all three domains of life and are termed RecA in bacteria [2], 8 9 RadA in archaea [3], and Rad51 in eukaryotes [4]. The strand exchange protein identified in 10 archaea, RadA, is more similar to the eukaryotic Rad51 protein (~40% amino acid identity) than 11 to its prokaryotic counterpart, RecA in eubacteria (~20% amino acid identity) [3, 5, 6]. 12 Crystallographic structure analysis of archaeal RadA has contributed significantly to our understanding of the structure and function of eukaryotic Rad51 [7, 8]. Various accessory 13 proteins also participate in the strand exchange reaction and these are usually termed 14 15 recombination mediators. Many recombination mediators are encoded by genes that arise as a result of a duplication event within the genome, and are termed paralogs. In yeast, the paralogous 16 proteins Rad55 and Rad57 share some similarity to Rad51 and function as a heterodimeric 17 complex which may stabilize the Rad51-assembled nucleoprotein filament [9]. Five Rad51 18 paralogs, XRCC2, XRCC3, RAD51B/RAD51L1, RAD51C/RAD51L2, and 19 RAD51D/RAD51L3, are found in eukaryotes. These paralogs form various heteromeric 20 complexes, mostly dimers such as RAD51B-RAD51C, RAD51D-XRCC2, RAD51C-XRCC3 or 21 tetramers such as RAD51B-RAD51C-RAD51D-XRCC2 [10]. 22 23 The identification and characterisation of RecA-like proteins in both crenarchaea and euryarchaea shows that, in common with eukaryotic Rad51, archaeal RadA proteins exhibit a 24

high level of diversity. This may reflect that the archaea demonstrate an evolutionary mixture of
 replication, repair and recombination functions in between simple bacterial and more complex
 eukaryotic forms of life.

RadB is the first RadA paralog to be characterised in archaea and its presence is confined to the
Euryarchaeota. RadB lacks the N-terminal domain of RadA [11] and strand exchange activity
[12]. Interaction of RadB with RadA has been demonstrated in *Pyrococcus furiosus* (Pfu) and it
was proposed that RadB functions in a manner analogous to the yeast Rad55-57 proteins in the
strand exchange reaction [12]. Genetic analysis of *radB* from the euryarchaeon *Haloferax volcanii* (Hvo) demonstrates that RadB functions in the homologous recombination pathways in
concert with RadA [13, 14].

Although Hvo RadB has been successfully over-expressed in *E. coli* [15] and both Hvo RadA and
RadB proteins have been partially purified after conditional over-expression of proteins in their
native host [16, 17], complete purification and structural and biochemical characterization of
RadA is still lacking in *H. volcanii*.

In addition to RadB, other DNA-associated halophilic proteins, Hvo PCNA and RPA3, have 15 previously been purified to suitable purity and in sufficient quantities for structural and 16 17 biochemical analyses [18, 19]. However when similar approaches were employed with Hvo RadA, the recombinant protein was found to co-purify with DNA when over-expressed alone in 18 19 *Escherichia coli*. The study of recombination proteins in the hyperthermophilic crenarchaeon, 20 Sulfolobus tokodaii (St), has shown that over-expression and characterization of StRad55B, a paralog of StRadA, was possible only when it was co-expressed with StRadA protein [20]. We 21 22 explored co-expression as a suitable strategy for studying protein-protein interactions using a 23 non-halophilic host, E. coli. The present study describes, for the first time, soluble over-

1	expression of the Hvo RadA-RadB complex, overt demonstration of their interaction and robust	
2	methodology to enable their further characterization and study.	
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21	Materials and Methods	
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23	Bacterial strains, plasmids and growth conditions.	
24	<i>E. coli</i> chemically competent strains DH5 α^{TM} (Invitrogen) and Rosetta 2 (DE3) (Novagen) were	
25	used for cloning and gene expression, respectively. The PCR-amplified target gene was cloned	

1 first into the Zero Blunt PCR vector (Invitrogen) and sequenced to exclude amplification errors.

The pET11 expression vector (Novagen) was used when single overexpression of radA or radB 2

was desired. For co-expression and purification of RadA and RadB, the E. coli Rosetta 2 (DE3) 3

4 strain was transformed with pET-Duet-1-based plasmid constructs (Novagen).

Bacterial cultures were grown at 37°C in Luria-Bertani (LB) broth in a shaking incubator or on 5

- LB agar plates supplemented with ampicillin (100 μ g ml⁻¹), chloramphenicol (34 μ g ml⁻¹) or 6
- kanamycin (34 μ g ml⁻¹) as appropriate. 7

Knock out of the arnA gene from E.coli Rosetta2 (DE3) expression strain 8

9 A mutant of the Rosetta2 (DE3) expression strain carrying a kanamycin resistance cassette

(kan^R) in place of the *arnA* gene (arnA736 (del):: kan^R) was generated by P1 phage-mediated 10

transduction [21]. Donor strain, obtained from the E.coli Genetic Stock center (CGSC), was 11

grown at 37°C to an OD₆₅₀ of 0.3-0.4 to add the P1 phage stock ($\sim 10^7$ -10⁸ PFU). After 3-4 h 12

incubation, purified lysate was used to transduce the kan^R marker into recipient Rosetta2 (DE3) 13

cells. Recipient strain was grown at 37°C to OD₆₅₀ of 0.8, pelleted, re-suspended in 0.1 M 14

MgSO₄, 5 mM CaCl₂ and incubated with purified lysate for 25 minutes at 37°C. After addition 15

of sodium citrate to a final concentration of 0.5 M to chelate calcium ions and prevent further 16

17 total lysis of the recipient strain, transductants were plated and subcultured to purity on LB agar containing kanamycin (34 μ g ml⁻¹) and chloramphenicol (34 μ g ml⁻¹).

19 **Molecular methods**

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Hvo radA gene (Accession number: U45311) was PCR-amplified from the H. volcanii wild-type 20 DS2 strain [22]. The forward primer (radA6HisF) contained an NdeI restriction site (underlined) 21 22 and a 6xHis tag

5'GACCTCATATGCATCACCATCACCATCACATGGCAGAAGACGACCTC-3' and the 23

1 reverse primer (*radABamR*) encoded a BamHI restriction site (underlined) 5'-

GCAATGGATCCTTATTACTCGGGCTTGAGACCGGCGTCCTG-3'. The His-tagged radA 2 gene was cloned first into the Zero Blunt PCR vector for sequencing and then sub-cloned for 3 overexpression either into pET11 (using NdeI and BamHI restriction sites) or pET-Duet1 4 plasmid at multiple cloning site 2 (MCS-2) (using the NdeI and EcoRV restriction sites). The 5 6 Hvo radB coding sequence (690bp) was excised from plasmid pCPG42 using the NdeI and BamHI restriction sites [15]. After sequence verification in a sub-cloning vector, radB was 7 cloned into MCS-1 of pET-Duet1 using the NcoI and HindIII restriction sites to yield the 8 9 pBPRAD2 construct. The relevant properties of the strains are listed in Table 1. **Over-expression, purification and analysis of RadA and RadB proteins** 10 Rosetta2 (DE3) or Rosetta2 (DE3) $\Delta arnA E. coli$ transformed with the pBPRAD2 construct were 11 used for over-expression and purification of co-expressed His-tagged RadA and un-tagged RadB 12 proteins. An overnight culture was inoculated into baffle flasks containing 2.4 L LB broth 13 supplemented with antibiotics as appropriate, and grown at 37° C. Over-expression of *radA-B* 14 was induced with 0.2 mM isopropyl β -D-1-thiogalactopyranoside (IPTG) at an OD₆₀₀= 0.4 – 0.6 15 with incubation for 3 further hrs in a shaking incubator at 30° C. The cell pellet was harvested by 16 17 centrifugation and lysed by sonication on ice in buffer A (50 mM HEPES, pH7.0, 10 mM imidazole and 1 M NaCl) containing EDTA-free protease inhibitor cocktail (Roche). The cell 18 lysate was clarified by centrifugation at 16,000 x g for 30 minutes at 4°C to remove insoluble 19 20 debris. Soluble proteins in the supernatant were used for downstream purification of the complex using 21 22 Cobalt-based immobilized metal affinity chromatography (IMAC). A Liquid Chromatography 23 Column (Size: 2.5 cm x 10 cm, bed: 49 mL.-Sigma-Aldrich) containing 5 mL of Talon® metal

1	affinity resin (Clontech) was equilibrated with 20 column volumes of buffer A (50 mM HEPES
2	pH 7.0, 1 M NaCl and 30 mM imidazole). The clarified soluble fraction was loaded and
3	incubated with rolling for 10 minutes at room temperature. The flow-through and two 20 mL
4	buffer A washes were retained separately in fresh tubes for SDS-PAGE gel analysis. Bound
5	proteins were eluted with 13 mL of buffer B (50 mM HEPES pH 7.0, 1 M NaCl and 300 mM
6	imidazole) following rolling for 5 minutes at room temperature, prior to collection of the eluate.
7	The eluate was loaded onto a 26/60 Superdex 200 (S200) preparative column (GE Healthcare)
8	using an ÄKTA Prime Plus system. The column was pre-equilibrated and run using size
9	exclusion chromatography (SEC) buffer (50 mM HEPES pH 7.0 and 1 M NaCl). The proteins in
10	each IMAC and SEC fraction were analyzed by SDS-PAGE gel. Samples containing low
11	concentrations of protein were first concentrated using StrataClean resin (5 μ l ml ⁻¹) (Stratagene)
12	as previously described [23] prior to gel loading. All samples were adjusted to maintain
13	equivalent loading in all gels.
14	The identity of RadA and RadB proteins was confirmed using MALDI-TOF-MS analysis of
15	excised bands by the Biopolymer Synthesis and Analysis Unit (BSAU), University of
16	Nottingham. Quantitative reflectance densitometric analysis of Coomassie Brilliant Blue-stained
17	proteins was performed using a BioRad GS-800 calibrated densitometer and Quantity One®
18	Software. Western blots were blocked with 5%, w/v, milk powder in phosphate buffered saline
19	(PBS)-Tween 20, then probed with 10 μL (1/1000 diluted) of alkaline phosphatase conjugated-
20	Mouse Anti-Hexa-His antibodies (Sigma) for 1 h and developed using BCIP/NBT substrate
21	according to manufacturer's instructions (Sigma).

22 Electron microscopy

Co-expressed, purified RadA-B proteins were concentrated to 1 mg ml⁻¹ using a Vivapore 10/20
7500 Da cutoff (Vivascience) and were maintained in 1 M NaCl buffer to preserve the soluble
and native state of the proteins. Protein concentrations were determined by using the Qubit
protein assay kit (Invitrogen). Protein samples were applied to a carbon-Formvar grid (Agar
Scientific) and allowed to settle for 10 minutes. The sample was then stained with either 0.5 or
1%, w/v, phosphotungstic acid (PTA; pH 7.0) for 1 minute. The grids were then imaged using a
JEOL JEM1010 transmission electron microscope at x100, 000 or x200, 000 magnification.

8 Homology Modelling

9 Homology modelling of the Hvo RadA primary sequence was performed using the PyMod 2.0 plugin module for PyMol [24, 25] as a convenient interface to Modeller 9v4 [26]. The Pfu RadA 10 structure (1PZN) chain A was used as the highest scoring template and aligned against the Hvo 11 RadA sequence using Clustal Omega. Modeller parameters were adjusted to accommodate 12 automated building of disulfide bridges and the highest level of optimization and refinement, 13 with additional energy minimisation performed on resulting models. Output models with 14 corresponding low scoring Discrete Optimised Protein Energy (DOPE) profiles, indicative of 15 limited modelling errors were further inspected for agreement with secondary structural elements 16 17 and diversity in loop disposition prior to final model selection.

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20 **Results**

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Conservation of the ATPase domain of RadA from both halophilic and non-halophilic species

24 Structural comparison of RadA from halophilic *P. furiosus* with RadA homologs in other

25 domains of life reveals similarity in the ATPase domain (AD) although differences are apparent

at amino and carboxyl terminal domains of the proteins (Fig. S1a). Conservation of amino acid
residues at the Walker A, Walker B and DNA binding (L1 and L2) motifs is also apparent in a
protein sequence alignment of Hvo RadA with RadA homologs (Fig. S1b) and the Hvo RadB
protein. Aside from this motif organization, Hvo RadA and RadB proteins are otherwise
dissimilar, sharing only 18.5% 34.1% identity/similarity at the amino acid level. Hvo RadB is
also a somewhat smaller protein, due to lack of a RadA-equivalent N-terminal domain (Fig.
S1c).

Along with a characteristic negative charge distribution on the protein surface, a distinct
preference for certain amino acids is also a hallmark of halophilic proteins. This is apparent in
Hvo RadA as a comparative increase in the acidic residue, aspartate, which is the most common
adaptation relative to mesophilic and thermophilic counterparts (Fig. S2). However, whereas a
marked reduction in the lysine content and increases in serine and alanine have been reported in
other halophilic proteins [27] this is not similarly the case for Hvo RadA. This disparity reflects
the lack of any currently known universal form of halophilic adaptation.

It is of particular interest to understand how halophilic proteins have adapted to interact with 15 DNA, given the involvement of basic residues in DNA binding of the phosphate backbone. Guy 16 17 and others (2006) demonstrated that residues in a conserved basic patch (the KHR triplet) were 18 crucial for DNA binding in Hvo RadB. To gain some insight into how preference for particular 19 amino acid usage in Hvo RadA, would likely affect the surface character of Hvo RadA, we built 20 a structural model in Modeller 9v4 for comparative analysis. Superposition of the Hvo RadA model with RadA homologs shows the expected correspondence with the principal structural 21 22 elements in the N-terminal and ATPase domains (Fig. 1a) although, unlike other RadA proteins, 23 an extensive insertion loop found only in the Hvo RadA sequence (159-182) is also apparent

1 (Fig. S1b).

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Fig. 1b shows the conserved elements of RadA in the Hvo RadA homology model in relation to 2 lysine residues within the molecule. The L1 and L2 loops within the ATPase domain are 3 involved in ssDNA binding. Chen and others (2007) identified a number of key positively 4 charged residues in Sso RadA L1 (R217, R223 and R229) that are also conserved in Hvo RadA. 5 6 Conversely, of the two lysine residues in the N-terminal domain of Sso RadA involved in dsDNA binding, only the equivalent of K27 is retained in Hvo RadA, with K60 being substituted 7 with as aspartate residue (D52). However, Hvo RadA possesses an alternative lysine at position 8 9 42. The majority of the lysine residues within Hvo RadA are located on the face of the molecule containing the NTD, L1 and L2 motifs, suggesting their retention may be related to involvement 10 11 in DNA-binding. The large insertion sequence seemingly unique to Hvo RadA is of potential interest (159-182) 12 (Fig. 1 and Fig S1b). Given the lack of homology to other family members the conformation of 13 the modelled loop remains speculative but it can be assumed that it is positioned on the face of 14 the molecule as indicated in Fig.1b. The loop region comprises a large number of acidic residues, 15 both aspartate and glutamate, which, in the context of the uncommonly higher proportion of

17 lysines in Hvo RadA relative to other halophilic proteins, may function to maintain solubility of the molecule. 18

Differential co-expression and purification of soluble RadA-RadB proteins 19

20 His-tagged Hvo radA and un-tagged radB were co-expressed from pBPRAD2 in the Rosetta 2 (DE3) *E.coli* strain in order to minimise issues with codon usage that can occur when expressing 21 22 archaeal proteins in a heterologous host [28]. SDS-PAGE gel analysis of clarified lysates of 23 induced cultures identified overexpressed proteins at the anticipated molecular weight for 6xHis-

1 RadA (~ 40 kDa) and RadB (~25 kDa) (Fig. 2). However, by comparison with the broadly equivalent amounts of RadA and RadB observed when each protein was expressed from pET11 2 individually (Fig 2. lanes 2 & 4 respectively), RadB, when co-expressed with RadA, appeared 3 4 substantially diminished in abundance. This may suggest a possible issue with bicistronic 5 expression, although DUET vectors are engineered for independent promotor control of both 6 multiple cloning sites to avoid such problems. Alternatively, it may be the case that RadA affects RadB expression or stability; a previous analysis of the RadA:RadB ratio in P. furiosus cell 7 extract showed the cellular amount of RadB protein to be ~200 times lower than that of RadA 8 9 [12]. Consequently, the lack of equimolar expression observed may better reflect a more physiological ratio of RadA and RadB proteins and, as such, was not considered to be an 10 impediment to further analysis. 11 H. volcanii requires 2 M sodium chloride to maintain internal osmotic balance in artificial media 12 [22]. Lowering the salt concentration could adversely affect the solubility of both proteins and 13 hence result in aggregation or precipitation. Therefore during purification procedures a high 14 ionic strength was maintained in all buffers (1 M NaCl) to promote the solubility of Hvo RadA 15 and RadB proteins. IMAC was used as a first step for the purification of His-tagged RadA and 16 17 un-tagged RadB using a cobalt-based metal affinity resin. The cobalt-based resin was observed 18 to bind His-tagged halophilic RadA protein with a higher specificity compared to nickel-based 19 resins, reducing non-specific binding of host proteins (data not shown). 20 Co-eluted His-tagged RadA and un-tagged RadB proteins were observed at a similar ratio as before at their expected molecular weight positions by SDS-PAGE analysis (Fig. 2b), indicating 21

22 specific interaction. Eluted fractions were subsequently subjected to size exclusion

chromatography (SEC) for further purification and to provide an initial indication of the

stoichiometry of RadA-RadB interaction. Both proteins eluted together as a complex of ~370
kDa in a broad peak from 120-170 mL (Fig. 3a). No difference in protein ratio was apparent in
any elution fraction suggesting that complex formation may be constrained by a particular
stoichiometric configuration (Fig. 3b).

5 Optimisation of the heterologous expression system

Whereas previous experiments demonstrated successful purification of a stable RadA-RadB 6 complex, subsequent analysis was hampered by a limitation of the purification protocol to 7 remove all non-specific host proteins. One major persistent contaminant (~64 kDa) that could not 8 9 be removed by standard optimisation approaches without significant concomitant decrease in yield of the RadA-RadB complex was identified by MALDI analysis to be the product of the 10 arnA gene (bi-functional polymyxin resistance protein, ArnA). The primary sequence of ArnA 11 demonstrates an abundance of histidine residues [27] likely accounting for its capture by IMAC 12 resins. Similarly, the ~74 kDa ArnA protein is reported to form a hexameric structure consisting 13 of a dimer of trimers [29, 30] with a predicted molecular weight of ~440 kDa, close to the 14 observed ~370 kDa RadA-B complex. 15

In view of this, bacteriophage-mediated P1 transduction was performed to delete the *arnA* gene from the Rosetta 2 (DE3) expression strain. The resulting Rosetta 2 (DE3) $\Delta arnA$ mutant strain was found to be equivalent to the parent strain in respect of growth characteristics and overexpression of RadA-RadB. Subsequently, RadA-RadB complex expressed from the modified strain was purified to ~95% homogeneity following the previous two-step purification protocol (data not shown).

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23 RadA oligomerizes into ring like structures

24

1	Recombinase proteins characteristically oligomerise into ring structures or helical nucleoprotein
2	filaments. Electron microscopy of P. furiosus RadA revealed that the protein forms dimers of
3	heptameric ring structures in solution in the absence of DNA [8] whereas, in S. solfataricus,
4	RadA forms an octameric ring bound to DNA [31]. Since Hvo RadA-RadB is consistently
5	purified as a large ~370 kDa stable complex, some form of RadA oligomerization is clearly
6	indicated. Preliminary densitometric analysis of co-purified RadA and RadB protein bands
7	estimates the ratio of RadA co-purified with RadB to be 4:1. Therefore the observed ~370 kDa
8	RadA-RadB SEC complex would most closely correspond to eight molecules of RadA in
9	complex with two molecules of RadB although this estimation is limited by the resolution of the
10	S200 SEC column employed in this study.
11	Subsequent electron microscopy observation of our purified Hvo RadA-RadB protein complex
12	confirms that Hvo RadA-RadB complex also possesses a tendency to self-associate into
13	multimeric structures in the absence of DNA (Fig. 4). In order to distinguish protein structures
14	from EM artifacts resulting as a consequence of high salt levels, several control grids were
15	prepared and imaged with either buffer only or buffer plus stain at various magnifications. The
16	images captured showed ring-like structures throughout the field which suggests that Hvo RadA-
17	RadB complex in its native state similarly exists as a ring-like structure (Fig. 4).

Discussion

The expression and purification of halophilic proteins, particularly from heterologous host
overexpression systems presents several challenges, principally due to the particular
physicochemical properties of these proteins which constrain conventional approaches.
Halophilic proteins are adapted to maintain solubility and function in hyper saline (1-4 M)

1 conditions [32]. The presence of high salt in halophilic organisms affects the gross molecular conformation of both proteins [33] and DNA [34]. Mesophilic DNA-binding proteins typically 2 make heavy use of positively charged residues to mediate electrostatic interactions with DNA 3 4 [15, 35, 36]. In the majority of organisms the presence of higher concentrations of salt interferes 5 with DNA-binding and leads to aggregate formation in mesophilic proteins. The situation is 6 reversed in halophilic counterparts. Halophilic proteins are generally decorated with negatively charged amino acid residues which allow the binding of surplus water and salt to build up a 7 hydrated solvent network on the surface of proteins. Structural studies have shown that binding 8 9 of hydrated cations (provided by excess salt and water molecules) around the negatively charged residues on the protein surface reduces the electrostatic repulsive forces between polyanionic 10 DNA and protein molecules [18, 37]. At lower concentrations of salt, the protective effect is lost 11 and the repulsive forces between the acidic residues lead to unfolding and inactivation of the 12 13 protein [27].

14 Consistently, H. volcanii RadA has an overall abundance of acidic amino acids, particularly aspartate which, based on homology modelling of RadA, contributes to the overall predicted 15 negative surface charge of the protein. Accounting for this, we successfully adapted a protocol 16 17 for purification of soluble RadA under moderate salt conditions. Further protocol modifications 18 were introduced to enable purification from persistent contaminants including both DNA and particular host proteins. Most notably, we overcame the previously observed deleterious effect of 19 20 RadA over-expression in E.coli by co-expression of RadA with RadB. These adaptions enabled RadA purification as a DNA-free complex with RadB using a relatively facile two-step 21 22 purification protocol. Based on our observations we speculate that RadA interacts with a small 23 proportion of RadB in a particular stoichiometry to stabilize it as a binding partner.

1 H. volcanii RadA is similar to other characterised RadA homologues. Consistent with its function as a recombinase, ATP and DNA binding motifs are apparent which have a well-2 conserved sequence composition despite the common skews in overall amino acid usage 3 typically observed in halophilic proteins (Fig. S1 and Fig. 1). The abundance of acidic amino 4 5 acids such as aspartate in Hvo RadA could potentially result in electrostatic repulsion with the 6 negatively charged DNA backbone under in vitro conditions (Fig. 1). The crystal structure analysis of Hvo PCNA indicates that this protein compensates for the reduction in positively 7 charged surface residues by employing cation binding [18]. However, the interaction in this 8 9 instance is largely topological in nature rather than via interaction of specific residues. A conserved basic patch on the surface of RadB has been shown to be crucial in DNA binding [15] 10 and number of similarly conserved positively charged residues are retained in Hvo RadA. The 11 presence of a highly negatively charged loop uniquely present in Hvo RadA may compensate for 12 the retention of lysine residues that presumably are maintained to enable direct DNA-binding 13 14 (Fig. 1a).

The failure to observe DNA binding activity in Hvo RadA across a range of methods (refer to 15 supplementary text and Table S1) suggests that, for this protein, the protein-DNA interaction 16 17 may require additional, as yet unidentified, factors. It is also possible that under the conditions of 18 our assays, RadA and RadB interaction occluded the DNA binding site or prevented an active 19 conformational state of the proteins required for DNA binding and ATPase activity. It is also 20 possible that additional co-factors are required for enzymatic activity of the complex. Recombinases have been generally observed to self-assemble into ring structures (six to eight 21 22 protomers) in their native state and monomerize to interact with DNA to form the helical 23 nucleoprotein filaments. So far the precise function of the ring structures is not clear in the

1	process of homologous base pairing. The toroidal form in recombinases has been suggested to	
2	function as an inactive storage form of protein, which is likely utilized to occlude the	
3	polymerization motif, preventing unwanted interaction of the protein with DNA. EM analysis of	
4	the purified protein complex showed that RadA-RadB also multimerizes into ring-like structures,	
5	which we consider may be of the order of a heptamer or octomer (Fig. 5), similar to its	
6	RadA/Rad51 homologs. On the basis of our collective observations, further studies to	
7	understand the dynamics of halophilic protein-protein and protein-DNA interactions in H.	
8	volcanii, including gross structural determination and robust biophysical characterization can	
9	now be anticipated.	
10		
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14 15	Conflicts of interest: none.	
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1 **References**

- 2
- Kowalczykowski SC, Eggleston AK. Homologous pairing and DNA strand-exchange proteins. Annu Rev
 Biochem. 1994;63:991-1043.
- 5 2. **Story RM, Weber IT, Steitz TA**. The structure of the E. coli recA protein monomer and polymer.
- 6 Nature. 1992;355(6358):318-25.
- 7 3. Sandler SJ, Satin LH, Samra HS, Clark AJ. recA-like genes from three archaean species with putative
- protein products similar to Rad51 and Dmc1 proteins of the yeast Saccharomyces cerevisiae. Nucleic
 Acids Res. 1996;24(11):2125-32.
- 10 4. Ogawa T, Shinohara A, Nabetani A, Ikeya T, Yu X, et al. RecA-like recombination proteins in
- eukaryotes: functions and structures of RAD51 genes. Cold Spring Harb Symp Quant Biol. 1993;58:567 76.
- 13 5. Shinohara A, Ogawa H, Ogawa T. Rad51 protein involved in repair and recombination in S. cerevisiae
- 14 is a RecA-like protein. Cell. 1992;69(3):457-70.
- 15 6. Komori K, Miyata T, Daiyasu H, Toh H, Shinagawa H, et al. Domain analysis of an archaeal RadA
- protein for the strand exchange activity. J Biol Chem. 2000;275(43):33791-7.
- 17 7. Ariza A, Richard DJ, White MF, Bond CS. Conformational flexibility revealed by the crystal structure of
- 18 a crenarchaeal RadA. Nucleic Acids Res. 2005;33(5):1465-73.
- 19 8. Shin DS, Pellegrini L, Daniels DS, Yelent B, Craig L, et al. Full-length archaeal Rad51 structure and
- 20 mutants: mechanisms for RAD51 assembly and control by BRCA2. EMBO J. 2003;22(17):4566-76.
- 21 9. Sung P, Klein H. Mechanism of homologous recombination: mediators and helicases take on
- regulatory functions. Nat Rev Mol Cell Biol. 2006;7(10):739-50.
- 23 10. Symington LS. Role of RAD52 epistasis group genes in homologous recombination and double-strand
- 24 break repair. Microbiol Mol Biol Rev. 2002;66(4):630-70, table of contents.
- 25 11. Sandler SJ, Hugenholtz P, Schleper C, DeLong EF, Pace NR, et al. Diversity of radA genes from
- 26 cultured and uncultured archaea: comparative analysis of putative RadA proteins and their use as a
- 27 phylogenetic marker. J Bacteriol. 1999;181(3):907-15.
- 28 12. Komori K, Miyata T, DiRuggiero J, Holley-Shanks R, Hayashi I, et al. Both RadA and RadB are
- involved in homologous recombination in Pyrococcus furiosus. J Biol Chem. 2000;275(43):33782-90.
- 13. Allers T, Ngo HP. Genetic analysis of homologous recombination in Archaea: Haloferax volcanii as a
 model organism. Biochem Soc Trans. 2003;31(Pt 3):706-10.
- 14. **Haldenby S, White MF, Allers T.** RecA family proteins in archaea: RadA and its cousins. Biochem Soc Trans. 2009;37(Pt 1):102-7.
- 15. Guy CP, Haldenby S, Brindley A, Walsh DA, Briggs GS, et al. Interactions of RadB, a DNA repair
- protein in archaea, with DNA and ATP. J Mol Biol. 2006;358(1):46-56.
- 36 16. Allers T. Overexpression and purification of halophilic proteins in Haloferax volcanii. Bioeng Bugs.
 37 2010;1(4):288-90.
- 17. Allers T, Barak S, Liddell S, Wardell K, Mevarech M. Improved strains and plasmid vectors for
- 39 conditional overexpression of His-tagged proteins in Haloferax volcanii. Appl Environ Microbiol.
- 40 2010;76(6):1759-69.
- 41 18. Winter JA, Christofi P, Morroll S, Bunting KA. The crystal structure of Haloferax volcanii proliferating
- 42 cell nuclear antigen reveals unique surface charge characteristics due to halophilic adaptation. BMC
 43 Struct Biol. 2009;9:55.
- 19. Winter JA, Patoli B, Bunting KA. DNA binding in high salt: analysing the salt dependence of
- 45 replication protein A3 from the halophile Haloferax volcanii. Archaea. 2012;2012:719092.

- 1 20. Sheng D, Li M, Jiao J, Ni J, Shen Y. Co-expression with RadA and the characterization of stRad55B, a
- 2 RadA paralog from the hyperthermophilic crenarchaea Sulfolobus tokodaii. Sci China C Life Sci.
- 3 2008;51(1):60-5.
- 4 21. Sternberg N. Bacteriophage P1 cloning system for the isolation, amplification, and recovery of DNA
- 5 fragments as large as 100 kilobase pairs. Proc Natl Acad Sci U S A. 1990;87(1):103-7.
- 6 22. Mullakhanbhai MF, Larsen H. Halobacterium volcanii spec. nov., a Dead Sea halobacterium with a
- 7 moderate salt requirement. Arch Microbiol. 1975;104(3):207-14.
- 8 23. Ziegler J, Vogt T, Miersch O, Strack D. Concentration of dilute protein solutions prior to sodium
- 9 dodecyl sulfate-polyacrylamide gel electrophoresis. Anal Biochem. 1997;250(2):257-60.
- 10 24. Janson G, Zhang C, Prado MG, Paiardini A. PyMod 2.0: improvements in protein sequence-structure
- analysis and homology modeling within PyMOL. Bioinformatics. 2016.
- 25. DeLano WL. The PyMol Molecular Graphics System. DeLano Scientific LLC, Palo Alto, Calif, USA.
 2008.
- 14 26. Webb B, Sali A. Comparative Protein Structure Modeling Using MODELLER. Curr Protoc
- 15 Bioinformatics. 2014;47:5 6 1-32.
- 16 27. Lanyi JK. Salt-dependent properties of proteins from extremely halophilic bacteria. Bacteriol Rev.
- 17 1974;38(3):272-90.
- 18 28. Poidevin L, MacNeill SA. Biochemical characterisation of LigN, an NAD+-dependent DNA ligase from
- the halophilic euryarchaeon Haloferax volcanii that displays maximal in vitro activity at high saltconcentrations. BMC Mol Biol. 2006;7:44.
- 29. Gatzeva-Topalova PZ, May AP, Sousa MC. Crystal structure and mechanism of the Escherichia coli
- 22 ArnA (PmrI) transformylase domain. An enzyme for lipid A modification with 4-amino-4-deoxy-L-
- arabinose and polymyxin resistance. Biochemistry. 2005;44(14):5328-38.
- 24 30. Breazeale SD, Ribeiro AA, McClerren AL, Raetz CR. A formyltransferase required for polymyxin
- resistance in Escherichia coli and the modification of lipid A with 4-Amino-4-deoxy-L-arabinose.
- Identification and function oF UDP-4-deoxy-4-formamido-L-arabinose. J Biol Chem. 2005;280(14):1415467.
- 31. Yang S, Yu X, Seitz EM, Kowalczykowski SC, Egelman EH. Archaeal RadA protein binds DNA as both
 helical filaments and octameric rings. J Mol Biol. 2001;314(5):1077-85.
- 30 32. **Dym O, Mevarech M, Sussman JL.** Structural features that stabilize halophilic malate dehydrogenase 31 from an archaebacterium. Science. 1995;267(5202):1344-6.
- 32 33. Madern D, Ebel C, Zaccai G. Halophilic adaptation of enzymes. Extremophiles. 2000;4(2):91-8.
- 33 34. Schlick T, Li B, Olson WK. The influence of salt on the structure and energetics of supercoiled DNA.
- Biophys J. 1994;67(6):2146-66.
- 35. Chen LT, Ko TP, Chang YW, Lin KA, Wang AH, et al. Structural and functional analyses of five
- 36 conserved positively charged residues in the L1 and N-terminal DNA binding motifs of archaeal RADA
 37 protein. PLoS One. 2007;2(9):e858.
- 36. Lee CD, Wang TF. The N-terminal domain of Escherichia coli RecA have multiple functions in
- 39 promoting homologous recombination. J Biomed Sci. 2009;16:37.
- 40 37. Bergqvist S, Williams MA, O'Brien R, Ladbury JE. Halophilic adaptation of protein-DNA interactions.
- 41 Biochem Soc Trans. 2003;31(Pt 3):677-80.
- 42 38. Hartman AL, Norais C, Badger JH, Delmas S, Haldenby S, et al. The complete genome sequence of
- 43 Haloferax volcanii DS2, a model archaeon. PLoS One. 2010;5(3):e9605.
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Table

4 Table 1. List of strains and plasmids.

Strains	Genotype/Description	Source
DH5a	F- ϕ 80lacZ Δ M15 Δ (lacZYA-argF) U169 recA1 endA1 hsdR17 (rK-, mK+) phoA supE44 λ - thi-1 gyrA96 relA1	Novagen
Rosetta 2 (DE3)	F- ompT hsdSB(rB- mB-) gal dcm (DE3) pRARE23 (CamR)	Novagen
Rosetta 2 (DE3)∆arnA	Rosetta 2 (DE3) with arnA inactivating mutation	This study
DS2	H. volcanii wild-type strain	(38)
		(22)
Plasmids		
pZero Blunt	Cloning vector	Invitrogen
pCPG42	Hvo radB encoding plasmid	(15)
pET11- <i>radA</i>	radA cloned into NdeI/BamHI sites for overexpression of His-	This study
pETDUET-radA	radA cloned into NdeI/EcoRV sites of MCS2 for overexpression	This study
pBPRAD2	<i>radB</i> cloned into <i>NcoI/Hind</i> III sites of pETDUET- <i>radA</i> MCS1 for co-expression of His-tagged RadA and untagged RadB	This study

Figure legends

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Fig. 1. a. Superposition of the Hvo RadA homology model with the Pfu RadA crystal

5 structure. The Hvo RadA homology model is shown in grey and the Pfu RadA structure

6 (1PZN:Chain A) in red (cartoon representation). Good correspondence is apparent between

7 major structural elements, although Hvo RadA can be distinguished by an additional large loop

8 insertion. b. Cartoon representation of the backbone of the modelled Hvo RadA stucture.

9 The Walker A motif (green), Walker B motif (red) and DNA binding loops (blue) are indicated.

10 The C α position of the lysine residues in Hvo RadA are indicated with orange spheres. The C α

11 position of acidic residues located in an insertion loop in Hvo RadA relative to related sequences

12 (see supplementary S1) are indicated with red spheres. The N-terminal domain (NTD), ATPase

13 domain (AD) and the insertion loop are labelled.

14

15 Fig. 2. a. Coomassie Brilliant-Blue stained SDS-PAGE gel demonstrating co-over-

expression of Hvo RadA and RadB in *E. coli*. Broadly equivalent levels of recombinant RadA
and RadB were observed in whole cell lysates when expressed independently (lanes 3 & 4,

18 respectively) compared with reduced levels of RadB observed when co-expressed with RadA

19 (lane 2, arrow). Lane 1 shows the pre-induction sample. RadA and RadB positions are indicated

20 in (B). Molecular weight markers are indicated (kDa). b. Purification of co-expressed RadA-

RadB proteins by cobalt-based immobilized metal affinity chromatography. Untagged RadB
is observed to co-purify with His-tagged RadA.

23

Fig. 3. a. Purification of RadA-RadB complex by Size Exclusion Chromatography (SEC). Chromatogram monitoring UV absorbance at 280 nm indicating the elution of the RadA-RadB complex at 130-170mL volume (arrow). Fractions as collected are indicated on the *x*-axis. The relative elution peaks for a series of molecular weight standards are shown with arrows. b. Coomassie Brilliant-Blue stained SDS-PAGE gel representing the purified proteins after

SEC. Lane 1 showing 10 μ l Talon purified load prior to purification on a S200 column. All S200

30 purified fractions corresponding to the UV peak at 100-180 mL (lanes 2-9) show RadA and

- 1 RadB co-purification at their estimated MW markers. An ~64kDa contaminating host protein
- 2 (ArnA) consistently co-eluted with the RadA-B complex as indicated in lanes 5-8.
- 3

4	Fig. 4. Transmission Electron Microscopy (TEM) images of co-expressed RadA-B complex.	
5	TEM images of co-expressed RadA-B complex at 100,000X (A and B) or 200,000X (C and D)	
6	magnifications. All of the protein samples were negatively stained with either 1% or 0.5%	
7	phosphotungstic acid (PTA). Ring structures are indicated with arrows. a. 100,000x (1% PTA	
8	negative stain), b. 100,000x (0.5% PTA negative stain), c. 200,000x (1% PTA negative stain), d.	
9	200,000x (0.5% PTA negative stain), e. enlarged images of ring-like RadA-B complexes.	
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Fig. 1







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Fig. 3



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- **Fig. 4**



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