



Genetic and biochemical identification of a novel single-stranded DNA-binding complex in *Haloferax volcanii*

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Single-stranded DNA (ssDNA)-binding proteins play an essential role in DNA replication and repair. They use oligonucleotide/oligosaccharide-binding (OB)-folds, a five-stranded β -sheet coiled into a closed barrel, to bind to ssDNA thereby protecting and stabilizing the DNA. In eukaryotes the ssDNA-binding protein (SSB) is known as replication protein A (RPA) and consists of three distinct subunits that function as a heterotrimer. The bacterial homolog is termed SSB and functions as a homotetramer. In the archaeon *Haloferax volcanii* there are three genes encoding homologs of RPA. Two of the *rpa* genes (*rpa1* and *rpa3*) exist in operons with a novel gene specific to Euryarchaeota; this gene encodes a protein that we have termed RPA-associated protein (*rpap*). The *rpap* genes encode proteins belonging to COG3390 group and feature OB-folds, suggesting that they might cooperate with RPA in binding to ssDNA. Our genetic analysis showed that *rpa1* and *rpa3* deletion mutants have differing phenotypes; only $\Delta rpa3$ strains are hypersensitive to DNA damaging agents. Deletion of the *rpa3*-associated gene *rpap3* led to similar levels of DNA damage sensitivity, as did deletion of the *rpa3* operon, suggesting that RPA3 and RPAP3 function in the same pathway. Protein pull-downs involving recombinant hexahistidine-tagged RPAs showed that RPA3 co-purifies with RPAP3, and RPA1 co-purifies with RPAP1. This indicates that the RPAs interact only with their respective associated proteins; this was corroborated by the inability to construct *rpa1 rpap3* and *rpa3 rpap1* double mutants. This is the first report investigating the individual function of the archaeal COG3390 RPA-associated proteins (RPAPs). We have shown genetically and biochemically that the RPAPs interact with their respective RPAs, and have uncovered a novel single-stranded DNA-binding complex that is unique to Euryarchaeota.

Keywords: archaea, *Haloferax volcanii*, RPA single-strand DNA-binding protein, COG3390 RPA-associated protein, DNA repair, protein overexpression, Cdc48d

INTRODUCTION

Genomic DNA must be unwound in order to be replicated or repaired, leaving it vulnerable to nuclease and chemical attack as well as open to the possibility of forming secondary structures. Binding of the single-stranded DNA (ssDNA)-binding proteins (SSB) RPA and SSB prevents any of these events from occurring (Lu et al., 2009). The SSB is denominated SSB in bacteria and replication protein A (RPA) in eukaryotes; they bind to ssDNA with high affinity and to dsDNA and RNA with low affinity (Wobbe et al., 1987; Wold et al., 1989; Kim et al., 1992). They play a vital organizational role in the central genome maintenance of the cell, providing docking platforms for a wide range of enzymes to gain access to genomic substrates (Lu and Keck, 2008). The bacteriophage T4 gene 32 monomer was the first SSB to be identified (Alberts and Frey, 1970). RPA was first identified as an essential protein for DNA replication in the eukaryotic simian virus (SV40; Wobbe et al., 1987) by stimulating the T antigen-mediated unwinding of the SV40 origin of replication (Kenny et al., 1989). RPA and SSB have now been established as essential

proteins for DNA metabolism including DNA replication, recombination, and repair in all domains of life (Wobbe et al., 1987; Heyer et al., 1990; Coverley et al., 1991, 1992; Moore et al., 1991; Wold, 1997). The basic architecture of RPA and SSB is based on the oligonucleotide/oligosaccharide-binding (OB)-fold, a five-stranded β -sheet coiled into a closed barrel, but the number of OB-folds present varies from species to species (Bochkarev and Bochkareva, 2004; Fanning et al., 2006).

Unlike in bacteria and eukaryotes, the architecture of SSBs present in archaea is not uniform. There is wide diversity of SSBs in the two main archaeal phyla, Euryarchaea, and Crenarchaea. Crenarchaea possess SSBs similar to those of bacteria, consisting of a single subunit with one OB-fold and an acidic C-terminus tail (Rolfsmeier and Haseltine, 2010). Euryarchaea have RPA-like proteins that show homology with the eukaryotic RPA, but from species to species the architecture of euryarchaeal RPAs varies dramatically from a single polypeptide RPA to an RPA made up of several subunits. Each of these RPAs can contain up to four OB-folds as well as a zinc finger motif (Rolfsmeier and Haseltine, 2010).

Pyrococcus furiosus RPA consists of three subunits RPA41, 14, and 32, denominated RPA1, 2, and 3, respectively, which form a heterotrimer as seen in eukaryotes. Strand exchange and immunoprecipitation assays have shown that *P. furiosus* heterotrimeric RPA stimulates strand exchange, and interacts with the clamp loader RFC and both DNA polymerases B and D (Komori and Ishino, 2001). The heterotrimeric complex seen in *P. furiosus* is also found in *P. abyssi* and *P. horikoshii*. However, in other archaeal species the *rpa* genes have undergone lineage-specific duplications, resulting in differing numbers of SSBs with diverse structures. Unlike the RPA complex found in *Pyrococcus* spp., or eukaryotic RPA, these do not form trimeric complexes (Robbins et al., 2004).

Methanosarcina acetivorans possesses three RPA subunits, MacRPA1, 2, and 3, which are unlikely to form a heterotrimeric complex as seen in *P. furiosus* and in eukaryotes. MacRPA1 contains four DNA-binding domains (DBD) containing OB-folds, MacRPA2 and 3 both have two OB-fold containing DBDs. Each of the three MacRPAs can function as SSBs, and are able to stimulate primer extension by *M. acetivorans* DNA polymerase BI (Robbins et al., 2004, 2005). This demonstrates an element of redundancy between the three MacRPAs, and suggests that the heterotrimeric RPA structure observed in *P. furiosus* is the exception and not the rule. Lin et al. (2008) suggest that intramolecular recombination between RPA homologs may have led to the diversity of RPAs found in euryarchaea, which can function in different pathways or cellular processes.

A similar pattern of lineage-specific gene duplication is seen with the archaeal MCM helicase, where the number and type of MCM subunits that make up the hexameric helicase differ between archaeal species. The genes encoding the MCM subunits fall into distinct phylogenetic clades, but these do not correspond to specific subunits of eukaryotic MCM. Instead they have arisen through lineage-specific gene family expansion (Chia et al., 2010). Such gene duplication might allow different archaeal species to refine the structure and function of MCM (and potentially RPA) for differing conditions and specialized roles.

Sulfolobus solfataricus has a bacterial-like SSB consisting of a small 20 kDa peptide containing one OB-fold and an acidic C-terminus tail (Haseltine and Kowalczykowski, 2002; Rolfsmeier and Haseltine, 2010). The *S. solfataricus* SSB quaternary structure is similar to that of *E. coli* SSB, however the primary structure of the OB-fold shows greater homology to that of the eukaryotic RPA70 DNA-binding domain B (DBDB). This suggests that euryarchaeal SSBs may be structurally similar to bacterial SSB but at a protein sequence level show homology to the eukaryotic RPA (Haseltine and Kowalczykowski, 2002; Kerr et al., 2003). In *S. solfataricus* there is an absence of DNA damage recognition proteins such as homologs of XPA or XPC to initiate NER. The ability of *S. solfataricus* SSB to specifically bind and melt damaged duplex DNA *in vitro* suggests SSB may play a role in the identification and binding of damaged DNA, followed by the subsequent recruitment of NER repair proteins (Cubeddu and White, 2005).

Haloferax volcanii encodes three RPA genes *rpa1*, *rpa2*, and *rpa3* (Hartman et al., 2010). Recent studies have shown RPA2 to be essential while RPA1 and RPA3 are not (Skowrya and MacNeill, 2012). Note that these authors used the nomenclature *rpaA1*, *A2*, *B1*, *B2*, *B3*, and *C* to refer to *rpa3*, *rpap3*, *rpa1*, *rpap1*, *rpe*, and

rpa2, respectively, while we have chosen to maintain the official nomenclature as described in Table 4 of the *H. volcanii* genome paper (Hartman et al., 2010). Both *rpa1* and *rpa3* are in operons with other genes; *rpa1* is in an operon with genes encoding an OB-fold containing protein (hereby designated RPA-associated protein or RPAP) and a calcineurin-like phosphoesterase, while only one OB-fold *rpa*-associated protein (*rpap*) gene is present in the *rpa3* operon (Figure 1). The presence of an *rpap* gene in the same operon as *rpa* can be found in other euryarchaeota, including *Halobacterium marismortui*, *Halobacterium salinarum*, and *Natronomonas pharaonis*, as well as in *M. mazei* and *M. barkeri*. The *rpap* gene has been assigned to the cluster of orthologous groups (COG) 3390 (Berthon et al., 2008).

To examine if RPA1 and 3, as well as RPAP1 and RPAP3 play a role in DNA repair, as is true for both the bacterial SSB and eukaryotic RPA, DNA damage assays were performed using the single and operon deletion mutants. Cells with deletions of the *rpa1* and *rpa3* operons had previously been examined by Skowrya and MacNeill (2012). However, this is the first report investigating the individual function of the archaeal COG3390 RPAP. We show genetically and biochemically that the RPAPs interact with their respective RPAs, and have thereby uncovered a novel SSB complex that is unique to Euryarchaeota.

MATERIALS AND METHODS

All chemicals were from Sigma and restriction enzymes from New England Biolabs, unless stated otherwise. Standard molecular techniques were used (Sambrook and Russell, 2001).

STRAINS AND PLASMIDS

Haloferax volcanii strains (Table 1) were grown at 45°C on complete (Hv-YPC), casamino acids (Hv-Ca), or minimal (Hv-Min) agar, or in Hv-YPC or Hv-Ca broth as described previously. Isolation of genomic and plasmid DNA, as well as transformation of *H. volcanii* were carried out as described previously (Allers et al., 2004).

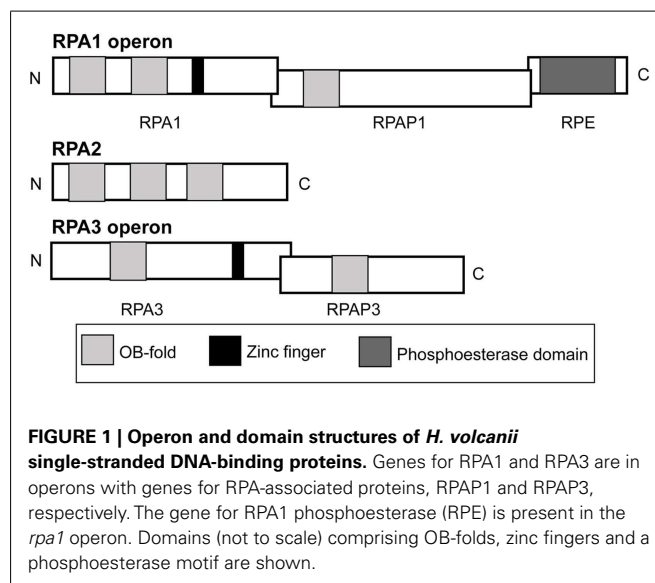


Table 1 | *Haloferax volcanii* strains.

| Strain | Relevant genotype* | Source or reference |
|--------|--|---------------------------------|
| DS2 | Wild-type | Mullakhanbhai and Larsen (1975) |
| H195 | Δ pyrE2 bgaHa-Bb leuB-Ag1 Δ trpA Δ hdrB | Guy et al. (2006) |
| H1209 | Δ pyrE2 Δ hdrB Nph-pitA Δ mrr | Allers et al. (2010) |
| H1216 | Δ pyrE2 bgaHa-Bb leuB-Ag1 Δ trpA Δ hdrB Δ rpa1::trpA+ | H195 pTA1170 |
| H1217 | Δ pyrE2 bgaHa-Bb leuB-Ag1 Δ trpA Δ hdrB Δ rpa1::trpA+ | H195 pTA1166 |
| H1244 | Δ pyrE2 bgaHa-Bb leuB-Ag1 Δ trpA Δ hdrB Δ rpa3::trpA+ | H195 pTA1174 |
| H1246 | Δ pyrE2 bgaHa-Bb leuB-Ag1 Δ trpA Δ hdrB Δ rpa1 operon | H195 pTA1189 |
| H1260 | Δ pyrE2 Δ hdrB bgaHa-Bb Δ rpa3 operon::trpA+ leuB-Ag1 Δ trpA | H195 pTA1207 |
| H1280 | Δ pyrE2 bgaHa-Bb leuB-Ag1 Δ trpA Δ hdrB Δ rpa1 | H1216 pTA1217 |
| H1281 | Δ pyrE2 bgaHa-Bb leuB-Ag1 Δ trpA Δ hdrB Δ rpa1 | H1217 pTA1141 |
| H1282 | Δ pyrE2 bgaHa-Bb leuB-Ag1 Δ trpA Δ hdrB Δ rpa1 operon rpa3 operon::[Δ rpa3 operon::trpA+, pyrE2+] | H1246 pTA1207 pop-in |
| H1326 | Δ pyrE2 bgaHa-Bb leuB-Ag1 Δ trpA Δ hdrB Δ rpa1 rpa3+::[Δ rpa3::trpA+ pyrE2+] | H1280 pTA1174 pop-in |
| H1333 | Δ pyrE2 Δ hdrB Nph-pitA Δ mrr Hsa-cdc48d | H1209 pTA1240 |
| H1390 | Δ pyrE2 bgaHa-Bb leuB-Ag1 Δ trpA Δ hdrB Δ rpa1 operon rpa3 operon::[Δ rpa3 operon::trpA+, pyrE2+] <rpa1 operon+hdrB+pyrE2> | H1282 pTA1265 pop-in |
| H1410 | Δ pyrE2 bgaHa-Bb leuB-Ag1 Δ trpA Δ hdrB Δ rpa3 | H195 pTA1284 |
| H1424 | Δ pyrE2 Δ hdrB Nph-pitA Δ mrr cdc48d-Ct | H1333 pTA1294 |
| H1430 | Δ pyrE2 Δ hdrB Nph-pitA Δ mrr cdc48d-Ct <p.tna::his tag+pyrE2+ hdrB+> | H1424 pTA963 |
| H1473 | Δ pyrE2 bgaHa-Bb leuB-Ag1 Δ trpA Δ hdrB Δ rpa1 rpap3+::[Δ rpap3::trpA+ pyrE2+] | H1281 pTA1284 pop-in |
| H1480 | Δ pyrE2 Δ hdrB Nph-pitA Δ mrr cdc48d-Ct <p.tnaA::his tag-rpa3 rpap3 pyrE2+ hdrB+> | H1424 pTA1280 |
| H1481 | Δ pyrE2 Δ hdrB Nph-pitA Δ mrr cdc48d-Ct <p.tnaA::rpa3 his tag-rpap3 pyrE2+ hdrB+> | H1424 pTA1281 |
| H1482 | Δ pyrE2 Δ hdrB Nph-pitA Δ mrr cdc48d-Ct <p.tnaA::his tag-rpa1 rpap1 pyrE2+ hdrB+> | H1424 pTA1326 |
| H1483 | Δ pyrE2 Δ hdrB Nph-pitA Δ mrr cdc48d-Ct <p.tnaA::rpa1 his tag-rpap1 pyrE2+ hdrB+> | H1424 pTA1327 |

*Genes shown within <> are present on an episomal plasmid, genes shown within [] are present on an integrated plasmid (pop-in).

CONSTRUCTION OF MUTANT STRAINS

Deletion mutants were constructed as described previously (Allers et al., 2004). Plasmids for gene deletion are shown in **Table 2**, and were generated by PCR using oligonucleotides shown in **Table 3**. Template DNA for the PCRs was isolated from genomic DNA.

CONSTRUCTION OF PROTEIN OVEREXPRESSION STRAINS

Protein overexpression strains were constructed by transformation with episomal overexpression plasmids as described previously (Allers et al., 2010). Plasmids for protein expression are shown in **Table 2**, and were generated by PCR using oligonucleotides shown in **Table 3**. Template DNA for the PCRs was isolated from genomic DNA.

UV IRRADIATION ASSAYS

UV irradiation assays were carried out as described previously (Delmas et al., 2009).

MITOMYCIN C ASSAYS

Mitomycin C (MMC) assays were carried out as described previously (Lestini et al., 2010).

PROTEIN OVEREXPRESSION AND PURIFICATION

Protein overexpression was carried out as described previously (Allers et al., 2010) with the following amendments: cultures were incubated at 45°C overnight to an OD₆₅₀ of 0.5, when protein expression was induced by adding 3 mM Trp to the culture

followed by incubation at 45°C, with shaking for a further 1 h until OD₆₅₀ ≈ 0.7.

PROTEIN PRECIPITATION

Deoxycholate was added to 0.015%, vortexed, and incubated for 10 min at room temperature. Trichloroacetic acid was added to 7.2% and incubated at room temperature for 5 min. Samples were centrifuged at 14,000 × g at room temperature for 8 min. Supernatant was removed and precipitated protein resuspended in 15 μl resuspension buffer (330 mM Tris-HCl pH 7.2, 2.6% SDS, 17 mM NaOH, 5% glycerol, 0.25 mg/ml bromophenol blue). Samples were heated for 10 min at 94°C and cooled on ice before loading onto an SDS-PAGE gel.

MASS SPECTROMETRY

Mass spectrometry of excised protein bands was carried out as described previously (Allers et al., 2010). Details of protein identification are given in the **Table A1** in Appendix.

RESULTS

RPA3 BUT NOT RPA1 FUNCTIONS IN DNA REPAIR

In eukaryotes, specifically *Saccharomyces cerevisiae*, all three RPA subunits have been shown to be essential for cell survival (Brill and Stillman, 1991). Work by Skowyra and MacNeill (2012) has shown that *H. volcanii* rpa2 is essential, which is in agreement with our fruitless attempts to delete rpa2 (data not shown). To examine if the other rpa genes of *H. volcanii* are also essential,

Table 2 | Plasmids.

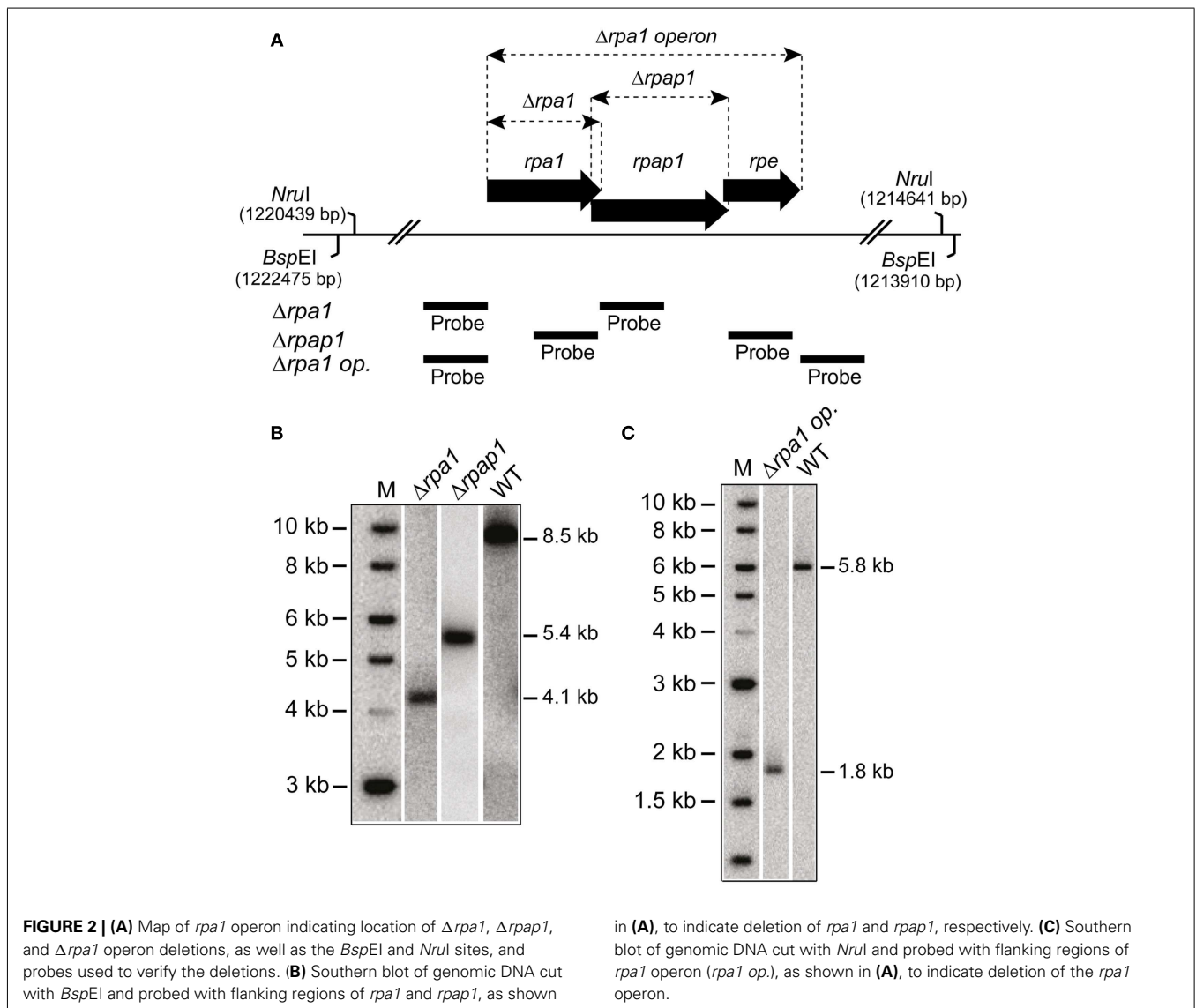
| Plasmid | Relevant properties | Source or reference |
|--------------------|---|----------------------|
| pBluescript II SK+ | Standard cloning vector | Stratagene |
| pTA131 | Integrative vector based on pBluescript II, with <i>pyrE2</i> marker | Allers et al. (2004) |
| pTA409 | Shuttle vector containing ampicillin, <i>pyrE2</i> and <i>hdrB</i> markers, and pHV1/4 replication origin | Delmas et al. (2009) |
| pTA884 | pBluescript II with <i>H. volcanii</i> 5,038-bp <i>EcoRI/NotI</i> genomic fragment containing <i>rpa3</i> operon | This study |
| pTA898 | pBluescript II with <i>H. volcanii</i> 7,335-bp <i>EcoRI/NotI</i> genomic fragment containing <i>rpa2</i> | This study |
| pTA937 | pBluescript II with <i>H. volcanii</i> 8,565-bp <i>BspEI</i> genomic fragment containing <i>rpa1</i> operon | This study |
| pTA963 | Overexpression vector with 6xHis-tag, <i>pyrE2</i> and <i>hdrB</i> markers, and pHV2 origin | Allers et al. (2010) |
| pTA1141 | pTA131 containing <i>rpa1</i> deletion construct inserted at <i>KpnI</i> and <i>XbaI</i> sites, contains an internal <i>NdeI</i> site | This study |
| pTA1142 | pTA131 containing <i>rpa3</i> deletion construct inserted at <i>EcoRI</i> and <i>KpnI</i> sites, contains an internal <i>NdeI</i> site | This study |
| pTA1166 | <i>rpa1</i> deletion construct pTA1141 with <i>trpA</i> marker, amplified from pTA298 introducing <i>NdeI</i> restriction sites to insert at internal <i>NdeI</i> restriction site in pTA1141 | This study |
| pTA1170 | Deletion construct of <i>rpap1</i> containing <i>trpA</i> marker from pTA298 inserted at <i>EcoRI</i> and <i>KpnI</i> sites in pTA131 | This study |
| pTA1174 | <i>rpa3</i> deletion construct containing <i>trpA</i> marker from pTA1166 inserted at <i>NdeI</i> restriction site | This study |
| pTA1180 | pTA131 with <i>cdc48d</i> deletion construct | Allers et al. (2010) |
| pTA1189 | pTA131 with <i>rpa1</i> operon deletion construct inserted at restriction sites <i>XbaI</i> and <i>EcoRI</i> with an internal <i>NdeI</i> site | This study |
| pTA1196 | <i>rpa3</i> operon deletion construct, using <i>NdeI/EcoRI</i> downstream fragment from pTA1282 (<i>rpap3</i> deletion construct) inserted at <i>NdeI/EcoRI</i> sites in pTA1142 (<i>rpa3</i> deletion construct), to replace the downstream fragment of the <i>rpa3</i> deletion construct | This study |
| pTA1207 | Deletion construct of <i>rpa3</i> operon pTA1196 with insertion of the <i>trpA</i> marker from pTA1166 at internal <i>NdeI</i> site | This study |
| pTA1217 | <i>RPAP1</i> deletion construct pTA1170 with upstream and <i>trpA</i> fragment replaced with the upstream fragment amplified from pTA937 by PCR, to introduce compatible <i>SphI</i> sites, generating non- <i>trpA</i> -marked deletion construct | This study |
| pTA1218 | pTA963 with <i>rpa3</i> inserted downstream of His-tag. <i>Asel</i> inserted after <i>rpa3</i> stop codon to allow insertion of His-tagged <i>rpa3</i> upstream of His-tagged <i>rpap3</i> (<i>Asel</i> is <i>NdeI</i> compatible) | This study |
| pTA1222 | pTA963 with <i>rpa1</i> N-terminally His-tagged, has an <i>Asel</i> site downstream of <i>rpa1</i> to allow insertion of <i>rpap1</i> (<i>NdeI</i> compatible) | This study |
| pTA1223 | pTA963 overexpression vector with <i>rpap1</i> N-terminally His-tagged inserted at <i>PsiI</i> and <i>BamHI</i> sites. <i>rpap1</i> was amplified by PCR from pTA937 introducing <i>BspHI</i> and <i>BamHI</i> sites | This study |
| pTA1224 | pTA963 with <i>rpap3</i> N-terminally His-tagged inserted at <i>PsiI</i> and <i>EcoRI</i> sites. <i>RPAP3</i> was amplified by PCR from pTA884 introducing <i>BspHI</i> and <i>EcoRI</i> sites | This study |
| pTA1240 | Gene replacement construct with insertion of 896 bp <i>Hsa-cdc48d</i> gene (amplified from <i>H. salinarum</i> DNA) between upstream and downstream flanking regions of <i>H. volcanii cdc48d</i> deletion construct pTA1180 | This study |
| pTA1265 | pTA409 with insertion of <i>rpa1</i> operon from pTA937 at <i>EcoRV</i> site | This study |
| pTA1280 | pTA1218 with <i>rpap3</i> amplified from pTA884 by PCR and inserted at <i>BstEII</i> and <i>EcoRI</i> sites after the N-terminally His-tagged <i>rpa3</i> , maintaining reading frame | This study |
| pTA1281 | pTA1224 with <i>rpa3</i> amplified from pTA884 by PCR and inserted upstream of N-terminally His-tagged <i>rpap3</i> at <i>NdeI</i> site | This study |
| pTA1282 | <i>rpap3</i> deletion construct with upstream and downstream regions amplified from genomic clone pTA884, introducing external <i>KpnI</i> and <i>EcoRI</i> sites, used to ligate into pTA131, and internal <i>NdeI</i> site | This study |
| pTA1284 | <i>rpap3</i> deletion construct pTA1282 with <i>trpA</i> marker digested from pTA1166 using <i>NdeI</i> and inserted at <i>NdeI</i> site in pTA1282, generating <i>trpA</i> -marked <i>rpap3</i> deletion construct | This study |
| pTA1288 | pBluescript II with <i>H. volcanii</i> 3,299-bp <i>Sall/BspHI</i> genomic fragment containing <i>cdc48d</i> gene | This study |
| pTA1294 | pTA131 with 2,247 bp <i>Hvo-cdc48d-Ct</i> gene replacement construct amplified from pTA1288: 1,797 bp <i>EcoRI-NheI</i> fragment with C-terminally truncated <i>cdc48d</i> plus upstream region, ligated to 485 bp <i>NheI-KpnI</i> fragment with downstream region of <i>cdc48d</i> , inserted at <i>EcoRI</i> and <i>KpnI</i> sites | This study |
| pTA1326 | pTA1222 with <i>rpap1</i> , amplified from pTA937 introducing <i>BstEII</i> and <i>BamHI</i> sites, and inserted downstream of His-tagged <i>rpa1</i> at <i>BstEII</i> and <i>BamHI</i> sites | This study |
| pTA1327 | pTA1223 with <i>rpa1</i> inserted upstream of His-tagged <i>rpap1</i> at <i>NdeI</i> site. <i>rpa1</i> was amplified from pTA937 introducing <i>NdeI</i> and <i>Asel</i> (<i>NdeI</i> compatible) sites | This study |

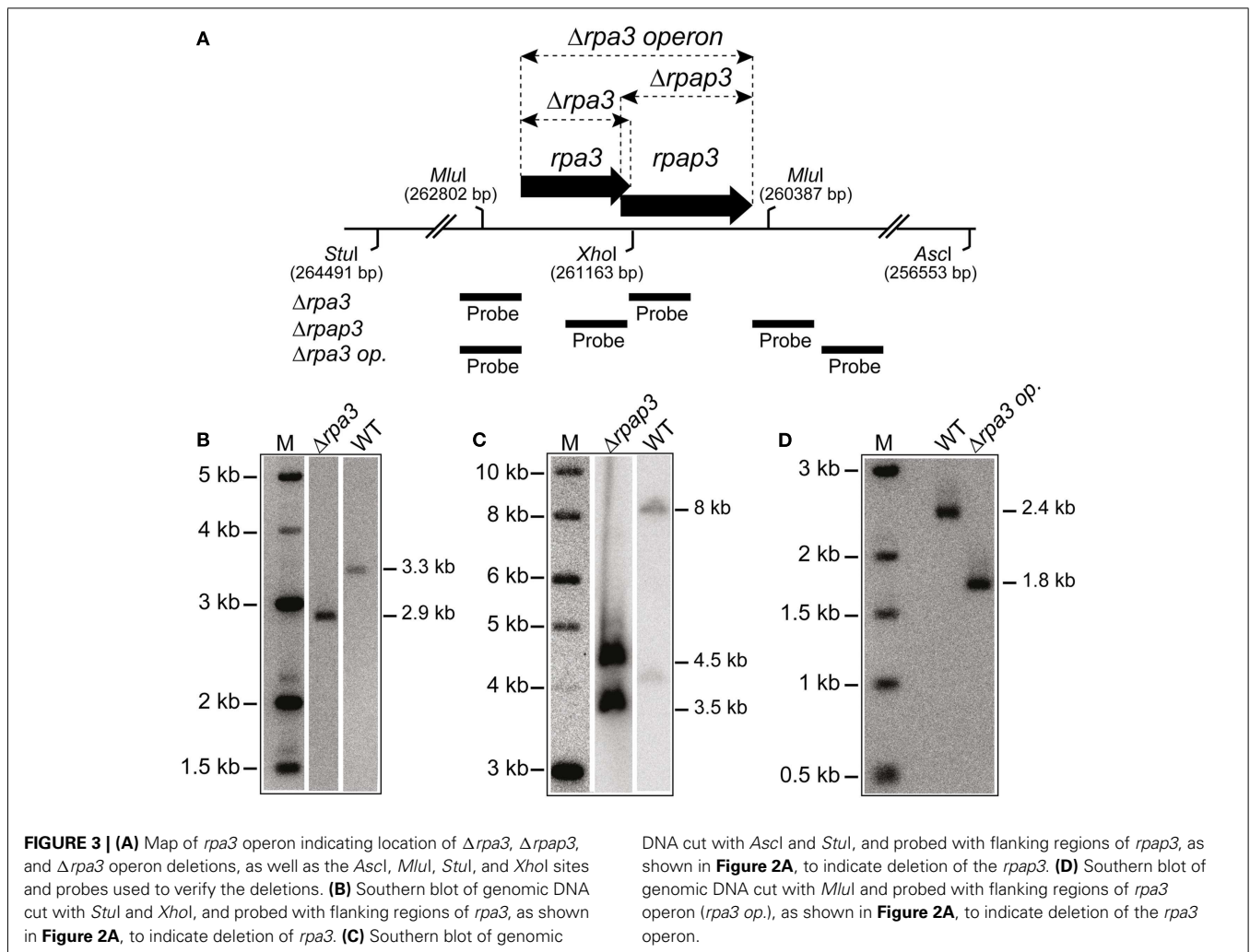
Table 3 | Oligonucleotides.

| Oligonucleotide | Sequence (5'–3') | Relevant properties | Use (plasmid generated) |
|-----------------|--------------------------------------|---|------------------------------------|
| Rpa1CF DS | GTTTCGAGGTACCGTTCGGGGAGC | $\Delta rpa1$ external downstream primer, <i>KpnI</i> site | pTA1141 |
| Rpa1CR DS | AGGTGCGCATATGAGCGCCTCGC | $\Delta rpa1$ internal downstream primer, <i>NdeI</i> site | pTA1141 |
| Rpa1 CR US | TACTACGTCTAGACGGACCTGTTCG | $\Delta rpa1$ external upstream primer, <i>XbaI</i> site | pTA1141 |
| Rpa1 CF US | GGTCGAGTTCATATGGTCGGGATTCCGC | $\Delta rpa1$ internal upstream primer, <i>NdeI</i> site | pTA1141 |
| Rpa3KpnI F | GCCGGTGGTACCACAGCCTC | $\Delta rpa3$ external upstream primer, <i>KpnI</i> site | pTA1142 |
| Rpa3NdeI R | GCAAATCAGTCATATGTACTCTCGCC | $\Delta rpa3$ internal upstream primer, <i>NdeI</i> site | pTA1142 |
| Rpa3EcoRI R | GACGGTGGAAATTCGGCCGTCC | $\Delta rpa3$ external downstream primer, <i>EcoRI</i> site | pTA1142 |
| Rpa3NdeI FC | GCGAGGTTCATATGATGAGTTCCAACG | $\Delta rpa3$ internal downstream primer, <i>NdeI</i> site | pTA1142 |
| trpANdeIF | CTCTGCACATATGTCGCTCGAAGACGC | <i>trpA</i> forward primer containing <i>NdeI</i> site | pTA1166 |
| trpANdeIR | TGCATGCCATATGCGTTATGTGCG | <i>trpA</i> reverse primer containing <i>NdeI</i> | pTA1166 |
| RPAP11kpnIus | CCGCGAGTGGTACCGCAAGCCCG | $\Delta rpa1$ external upstream primer, <i>KpnI</i> site | pTA1170 |
| RPAP11nsilIus | CGACGACCGGCGATGCATTATGCGCGC | $\Delta rpa1$ internal upstream primer, <i>NsiI</i> site | pTA1170 |
| RPAP11sphIus | GCTGAAGGGCATGCGAGGCCGTGC | $\Delta rpa1$ internal downstream primer, <i>SphI</i> site | pTA1170 |
| RPAP11ecoRIus | CGGCGAGAGAATTCCTGCCCGGG | $\Delta rpa1$ external downstream primer, <i>EcoRI</i> site | pTA1170 |
| PEcorI F | GCCCGAATTCCTGTCTGATTG | $\Delta rpa1$ operon external downstream primer, <i>EcoRI</i> site | pTA1189 |
| Rpa1CR US | TACTACGTCTAGACGGACCTGTTCG | $\Delta rpa1$ operon external upstream primer, <i>XbaI</i> site | pTA1189 |
| RPEndel R DS | CTACCGGAACATATGACTCGGGTCCG | $\Delta rpa1$ operon internal downstream primer, <i>NdeI</i> site | pTA1189 |
| Rpa1ndel F2 | GTTGGACCCATATGTCGAACGACG | $\Delta rpa1$ operon internal upstream primer, <i>NdeI</i> site | pTA1189 |
| RPAP11SphI US | GCGATTTCCCGCATGCGGACGACCG | $\Delta rpa1$ internal upstream primer, <i>SphI</i> site | pTA1217 |
| RPAP11 kpnI us | CCGCGAGTGGTACCGCAAGCCCG | $\Delta rpa1$ external upstream primer, <i>KpnI</i> site | pTA1217 |
| Rpa3BspHI F | AGGTAGATCATGACTGATTTGC | <i>rpa3</i> forward primer, <i>BspHI</i> site | pTA1218 |
| Rpa3 RAsel | CGAGTGGGGAATTCGTTGGAATTAATTACATC | <i>rpa3</i> reverse primer, <i>Asel</i> site | pTA1218 |
| Rpa1F NcoI | CCCGACTCCATGGAACCTCGACC | <i>rpa1</i> forward primer, <i>NcoI</i> site | pTA1222 |
| Rpa1Asel/EcoRI | CGGCGGCGAATTCGCGGTAGCGGATTAATCGCGTGC | <i>rpa1</i> reverse primer, <i>Asel</i> and <i>EcoRI</i> sites | pTA1222 |
| | pTA1327 | | |
| RPAP1F BspHI | GGTGCCTCATGAGCGCCTCG | <i>rpa1</i> forward primer, <i>BspHI</i> site | pTA1223 |
| RPAP1BamHI | CGTTCGGGGATCCGCGCCTGC | <i>rpa1</i> reverse primer, <i>BamHI</i> site | pTA1223 |
| | pTA1326 | | |
| RPAP3BspHI F | GTCGATGTTTCATGAGTTCCAACG | <i>rpa3</i> forward primer, <i>BspHI</i> site | pTA1224 |
| RPAP3EcoRI R | CGGTTCGGAATTCAGCCGAC | <i>rpa3</i> reverse primer, <i>EcoRI</i> site | pTA1224 |
| | pTA1280 | | |
| HsaCdc48F | GTTCTTGGCATATGACCGAGGCTCTC | Forward primer for <i>Hsa-cdc48d</i> , <i>NdeI</i> site | pTA1240, Probe Figure 5B |
| HsaCdc48R | CTGACAGATCTCGCAGTCACAGC | Reverse primer for <i>Hsa-cdc48d</i> , <i>BglIII</i> site | pTA1240, Probe Figure 5B |
| Rpa3BstEII | GATGCGCGGTGACCTCGTGG | <i>rpa3</i> forward primer, native <i>BstEII</i> site | pTA1280 |
| Rpa3NdeI | CGAGGTAGCATATGACTGATTTGCG | <i>rpa3</i> forward primer, <i>NdeI</i> site | pTA1281 |
| RPAP3 gitF | CTCCCAATGGGTACCAAGGTGGAGGC | $\Delta rpa3$ internal upstream primer, <i>NdeI</i> site | pTA1282 |
| RPAP3 gitR | TCGTTGGACATATGTTACATCGACCTCGC | $\Delta rpa3$ external upstream primer, <i>KpnI</i> site | pTA1282 |
| RPAP3 F DS | CTCGCTGAATTCGGTGGGTGC | $\Delta rpa3$ external downstream primer, <i>EcoRI</i> site | pTA1282 |
| RPAP3 R DS | CTGAGCGCATATGCGGGCGTCTCG | $\Delta rpa3$ internal downstream primer, <i>NdeI</i> site | pTA1282 |
| cdc48dUF | ACGGGTACCACGTTGCTGG | <i>Hvo-cdc48d</i> external upstream primer, <i>KpnI</i> site | pTA1294 |
| cdc48dDR | GCCGAATTCGAGCCGAGGTGG | <i>Hvo-cdc48d</i> external downstream primer, <i>EcoRI</i> site | pTA1294 |
| cdc48d-CtrR | CGGCGCGCTAGCCGACCGGTTACGC | Internal reverse primer to generate C-terminally truncated <i>Hvo-cdc48d</i> , <i>NheI</i> site at <i>cdc48d</i> stop codon | pTA1294 |
| cdc48d-CtrF | CTGTGGTGTAGCCGTCGTCGACCCCG | Internal forward primer to generate C-terminal truncated <i>Hvo-cdc48d</i> , <i>NheI</i> site at <i>cdc48d</i> stop codon | pTA1294 |
| cdc48dSeqF | GGAAAAAGGGGCAGATGGTG | Forward primer to downstream flanking region of <i>Hvo-cdc48d</i> | PCR Figure 5C |
| cdc48dHvSeqR | CGACGACATCTCGCTGATTCCG | Reverse primer to <i>Hvo-cdc48d</i> gene | PCR Figure 5C |
| cdc48dHsalSeqR | GGTCAACACGCTGCTGAAGTCC | Reverse primer to <i>Hsa-cdc48d</i> gene | PCR Figure 5C |
| Rpa1BstEII | CCGGACGCGTACCGCCATCC | <i>rpa1</i> forward primer, native <i>BstEII</i> site | pTA1326 |
| Rpa1NdeI | CCCGACCATATGGAACCTCGACC | <i>rpa1</i> forward primer, <i>NdeI</i> site | pTA1327 |

genomic deletions of *rpa1* and *rpa3* were generated using the counter selective pop-in/pop-out method (Allers et al., 2004). To generate the deletion constructs by PCR, *rpa1* and *rpa3* operons were first isolated from wild-type (WT) *H. volcanii* using native *BspEI* and *EcoRI/NotI* restriction sites, respectively, to generate genomic libraries. These were then screened for the presence of the *rpa1* and *rpa3* operons, individually, using colony hybridization. The isolated plasmids, pTA937 (*rpa1* operon) and pTA884 (*rpa3* operon) were confirmed by DNA sequencing. Deletion constructs for *rpa1* and *rpa3* were designed to avoid polar effects on the expression of the downstream *rpap* genes by maintaining the reading frame. Genomic deletions of both *rpa1* and *rpa3* (*trpA*-marked) were successful, generating strains H1217 and H1244, respectively (Figures 2 and 3, respectively). The ability to delete both *rpa1* and *rpa3* with relative ease, but not *rpa2*, indicates that the cellular requirement for each RPA is not equal, making it unlikely that they function collectively.

Both eukaryotic and bacterial SSB are involved in DNA repair. To examine if *H. volcanii* RPA1 and RPA3 function in DNA repair, the effects of DNA damage on cell survival of H1217 and H1244 were examined. UV irradiation results in the formation of cyclobutane pyrimidine dimers and 6-4 pyrimidine-pyrimidone dimer photoproducts, as well as ssDNA nicks that indirectly generate double-stranded DNA breaks (DSBs). The latter require repair by homologous recombination (HR) or single-strand DNA annealing (Fousteri and Mullenders, 2008; Rouillon and White, 2011). MMC is a chemotherapeutic agent that reacts with DNA generating covalent interstrand cross-links, requiring removal by nucleotide excision repair (NER) and HR (Tomasz et al., 1987). The $\Delta rpa1$ mutant H1217 was no more sensitive than the WT to UV and MMC-induced DNA damage, however the $\Delta rpa3$ mutant H1244 exhibited moderate sensitivity to both UV and MMC-induced DNA damage (Figure 4).





RPAP3 BUT NOT RPAP1 FUNCTIONS IN DNA REPAIR

Analysis of predicted protein domains indicates that RPA1 and RPA3 both possess zinc finger domains, and that RPA1 has three OB-folds compared to the single OB-fold present in RPA3 (**Figure 1**). Both COG3390 RPAs RPAP1 and RPAP3 possess a single OB-fold suggesting a possible role in DNA binding. The RPA1 phosphoesterase (RPE) has a calcineurin-like phosphoesterase domain, and was not investigated individually. However, our results and those of Skowryra and MacNeill (2012) show that *rpe* is a non-essential gene.

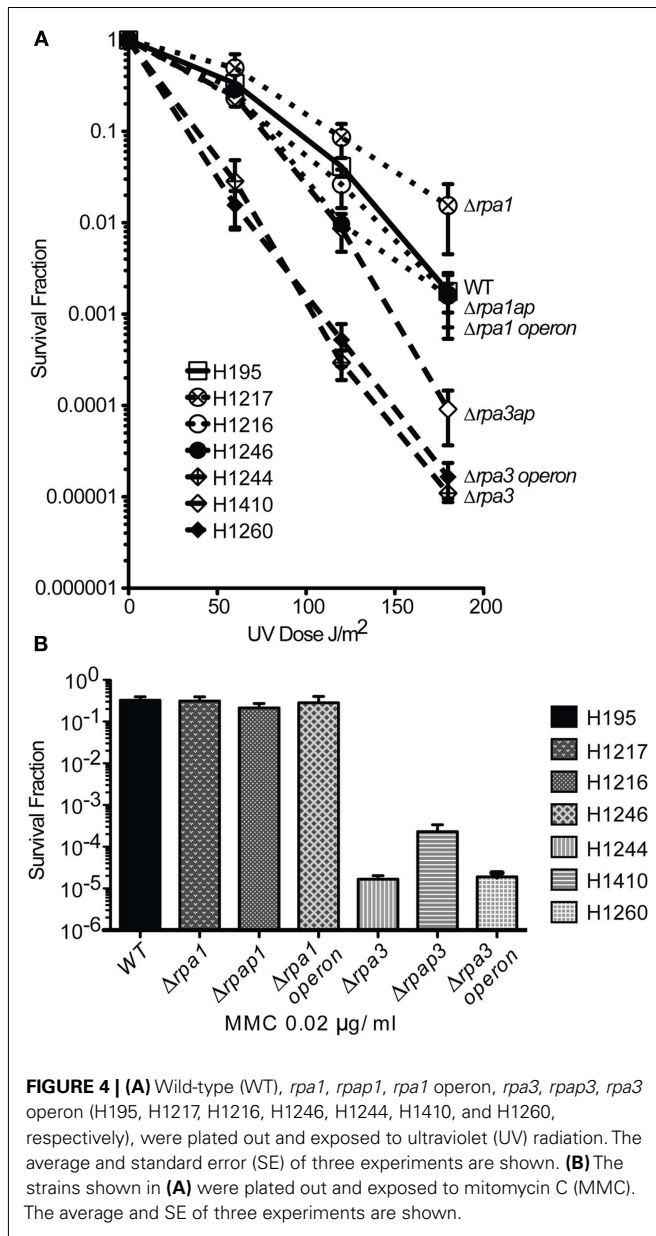
To study the roles of RPAP1 and RPAP3 in DNA repair, $\Delta rpap1$ (H1216) and $\Delta rpap3$ (H1410) mutants were generated, both using *trpA*-marked deletion constructs (**Figures 2 and 3**, respectively). As with $\Delta rpa1$ strain H1217, the $\Delta rpap1$ mutant H1216 showed no increased sensitivity to UV irradiation or to MMC-induced DNA damage. However, the $\Delta rpap3$ deletion mutant H1410 was hypersensitive to both types of DNA damage, and the level of sensitivity was similar to that exhibited by the $\Delta rpa3$ mutant H1244.

We examined whether the absence of both RPA and RPAP results in a synergistic deficiency in DNA repair. Genomic deletions of the *rpa1* and *rpa3* operons were generated in strains H1246 and H1260, respectively, with only the latter being a *trpA*-marked

deletion (**Figures 2 and 3**, respectively); deletions of the *rpa1* and *rpa3* operons have previously been reported by Skowryra and MacNeill (2012). The $\Delta rpa1$ operon mutant showed no increased sensitivity to UV irradiation or to MMC-induced DNA damage. However the $\Delta rpa3$ operon deletion mutant was hypersensitive to both types of DNA damage, and the level of sensitivity was similar to that exhibited by the single $\Delta rpa3$ and $\Delta rpap3$ mutants H1244 and H1410, respectively (**Figure 4**). This result suggests that RPA3 and RPAP3 function in the same pathway(s) of DNA repair.

REDUNDANCY BETWEEN RPA1 AND RPA3 OPERONS

In order to test for redundancy between the two RPAs, an attempt was made to generate a double $\Delta rpa1$ operon $\Delta rpa3$ operon deletion. This involved constructing the strain H1282, which contained the pop-in of a *trpA*-marked $\Delta rpa3$ operon construct (pTA1207) in an unmarked $\Delta rpa1$ operon background (H1246). An episomal plasmid (pTA1265), marked with *pyrE2* and providing in *trans* expression of the *rpa1* operon was used for complementation during the pop-out step (note that this episomal plasmid is lost during counter-selection with 5-FOA). Neither of the two pop-outs generated from this strain (H1390) yielded the desired $\Delta rpa1$ operon



$\Delta rpa3$ operon mutant (see **Figure A1** in Appendix). This indicates that the cell requires either RPA1 or RPA3 (and/or their respective RPAPs) for survival.

Next we attempted to generate $\Delta rpa1 \Delta rpa3$ and $\Delta rpa3 \Delta rpa1$ deletion mutants. This would test whether the RPAPs can complement each other, or whether they are instead specific for their respective RPAs. The *trpA*-marked $\Delta rpa3$ construct (pTA1284) was used in an unmarked $\Delta rpa1$ background (H1280), and the *trpA*-marked $\Delta rpa3$ construct (pTA1207) was used in an unmarked $\Delta rpa1$ background (H1281). In both cases two pop-outs were generated but none proved to be the desired deletions (see **Figure 1** in Appendix). This suggests that the putative RPA:RPAP complex is dependent upon specific RPA:RPAP interactions for functionality.

CONSTRUCTION OF PROTEIN OVEREXPRESSION STRAIN WITH C-TERMINAL TRUNCATION OF CDC48D

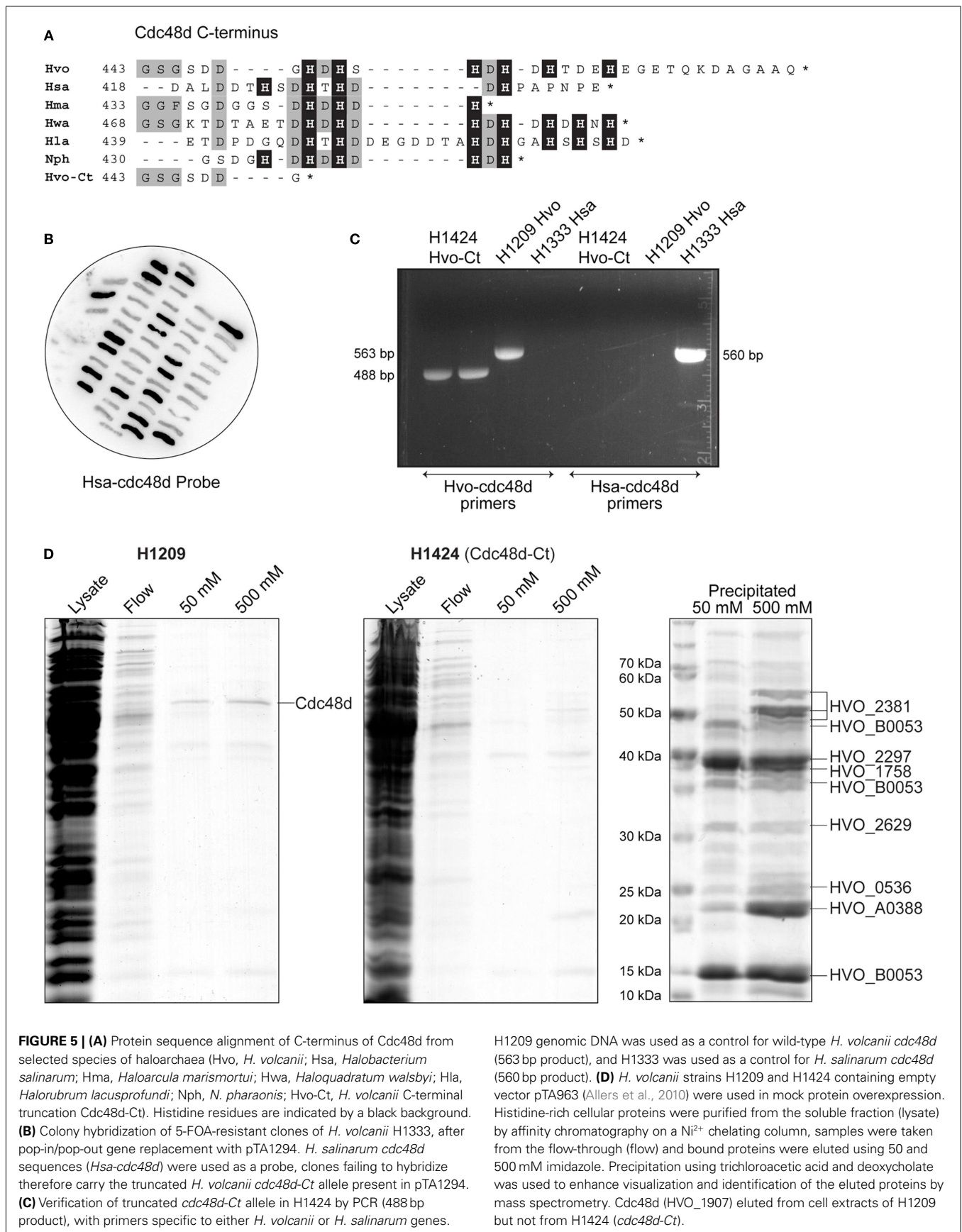
In a previous publication (Allers et al., 2010), we constructed a strain of *H. volcanii* where the histidine-rich *pitA* gene is replaced by the ortholog from *N. pharaonis*. The latter protein lacks the histidine-rich linker region found in *H. volcanii* PitA and does not co-purify with His-tagged recombinant proteins. The absence of Hvo-PitA revealed an additional co-purifying protein, which we identified as Cdc48d (HVO_1907) and features a histidine-rich C-terminus (**Figure 5A**). We were unable to delete *cdc48d*, indicating that this gene is essential (Allers et al., 2010). The presence of this contaminating protein was problematic for purification of His-tagged RPA1 and RPAP1, due to similar molecular weights (Cdc48d, 53 kDa; RPA1, 46 kDa; RPAP1, 65 kDa).

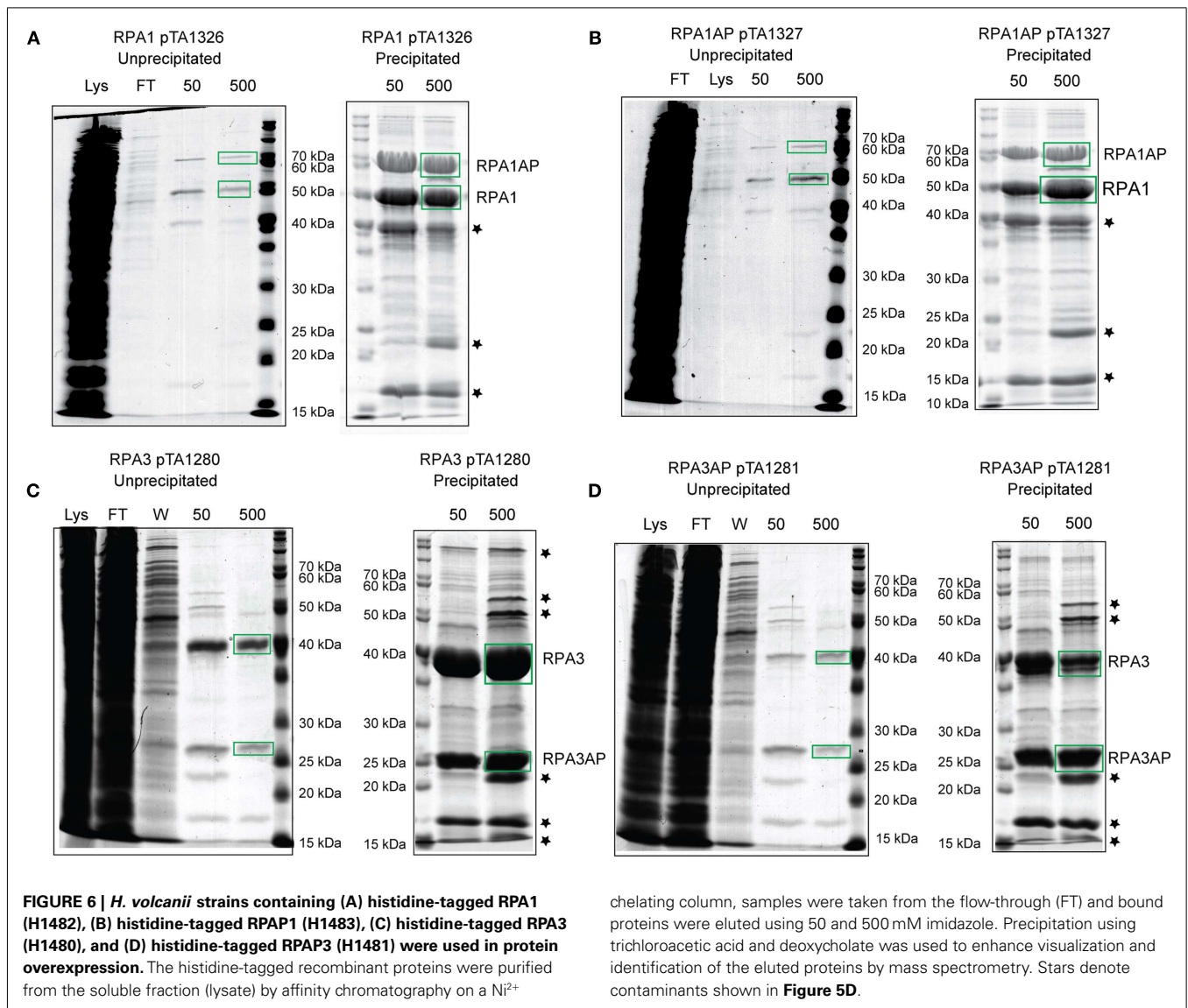
All orthologs of Cdc48d from haloarchaea feature a histidine-rich C-terminus, however Cdc48d from *Haloarcula marismortui* and *H. salinarum* have only three and four histidines, respectively, compared to six in *H. volcanii* (**Figure 5A**). Therefore, we replaced the *H. volcanii cdc48d* gene in H1209 (Allers et al., 2010) with orthologous genes from *H. marismortui* and *H. salinarum*, generating *H. volcanii* strains H1405 and H1333, respectively. Unfortunately these strains grew poorly and were not suitable for recombinant protein overexpression. Instead, we generated a truncated allele of *H. volcanii cdc48d*, encoding a Cdc48d protein lacking the histidine-rich C-terminus (Cdc48d-Ct; **Figure 5A**). The *cdc48d-Ct* allele was used to replace the *H. salinarum cdc48d* gene in *H. volcanii* H1333, generating H1424 (**Figures 5B,C**). This strain exhibits normal cell growth and the Cdc48d-Ct protein no longer co-purifies with His-tagged recombinant proteins (**Figure 5D**). A number of minor histidine-rich contaminants are now apparent, which have been identified by mass spectrometry.

DIRECT RPAP INTERACTION WITH RESPECTIVE RPA

The genetic analysis of *rpa1* and *rpa3* and their respective *rpa* genes indicates not only that RPA3 and RPAP3 function in the same DNA repair pathway(s), but also that they function together as a specific RPA:RPAP complex. To establish whether this is achieved via a direct RPA:RPAP interaction, affinity pull-downs were employed (Allers et al., 2010). The *rpa1* and *rpa3* operons were cloned under control of the tryptophanase promoter in plasmid pTA963, where either the RPA or the RPAP was tagged with a hexahistidine tag.

Histidine-tagged RPA1 and RPA3 pulled down their respective RPAPs, and histidine-tagged RPAP1 and RPAP3 pulled down their respective RPAs (**Figure 6**). However, histidine-tagged RPA1 did not pull down RPAP3, and vice versa. This confirms that the RPAs interact specifically with their respective RPAPs, supporting our conclusions based on the failure to generate $\Delta rpa1 \Delta rpa3$ and $\Delta rpa3 \Delta rpa1$ deletion mutants. Neither RPA1 nor RPA3 pulled down RPA2, and histidine-tagged RPA2 did not pull down RPA1 or 3, or either of the RPAPs (data not shown). This supports the suggestion that the three RPAs of *H. volcanii* do not form a heterotrimer as observed in eukaryotes and *P. furiosus*, but instead form three separate ssDNA-binding factors.





DISCUSSION

There is a unifying theme in archaea of a great variety in the number and type of proteins involved in DNA replication, and repair, whose counterparts in eukaryotes are much more uniform. This has been shown to be the case for RPA, where eukaryotes possess three subunits that all form unified clades in a phylogenetic analysis, but in archaea the number and structure of subunits varies widely. Some euryarchaea possess differing numbers of RPA subunits and some possess differing numbers of RPAs and RPAPs. Crenarchaea also possess varying numbers of SSB, however in both euryarchaea and crenarchaea none of the RPAs, RPAPs, or SSBs fall into unified clades. Again this is seen in the case of MCM, where eukaryotes possess six MCM subunits that each form unified clades, but in archaea there is a vast range in the number of MCM subunits, differing between individual species, and none of which fall into uniform clades (Chia et al., 2010). Characterizing the RPA-RPAP complexes of *H. volcanii* will shed light on how the RPAs and RPAPs function together in binding and stabilizing

ssDNA. This in turn will provide insight for other RPAs and RPAPs in archaea, but also offer reasoning behind the driving force of such non-uniform evolution of archaea.

The genetic and biochemical analysis presented here indicates the three RPAs of *H. volcanii* do not form a heterotrimeric complex as in *P. furiosus* and eukaryotes. Instead, RPA1 and RPA3 form complexes with their respective RPAPs. Unlike *rpa2*, both *rpa1* and *rpa3* genomic deletions were generated with relative ease, showing that the latter are not essential for cell survival and supporting the hypothesis that the three RPAs do not form a heterotrimeric complex.

The ease at which the *rpa1*, *rpap1*, and *rpa1* operon deletion mutants were made, coupled with a lack DNA damage sensitivity, signifies the *rpa1* operon does not play a major role in DNA replication or repair. The moderate DNA damage sensitivity shown by the individual *rpa3*, *rpap3*, and the *rpa3* operon mutants indicates that the efficient repair of UV and MMC-induced DNA damage requires the products of the *rpa3* operon but not the

rpa1 operon. However, it proved impossible to generate a double $\Delta rpa1$ operon $\Delta rpa3$ operon deletion, showing that cellular growth requires either RPA1 or RPA3.

Both single $\Delta rpa3$ and $\Delta rpa3$ mutants showed a similar DNA damage sensitivity to each other, and to the *rpa3* operon mutant, providing genetic evidence that RPA3 and RPAP3 act in the same DNA repair pathway. Furthermore, we were unable to generate $\Delta rpa1 \Delta rpa3$ and $\Delta rpa3 \Delta rpa1$ deletion mutants, indicating that RPAP1 could not substitute for RPAP3 (and vice versa), and suggesting that the RPA3 interacts specifically with RPAP3 (and likewise for RPA1 and RPAP1). However it is unclear what role the associated proteins play, since the presence of an OB-fold does not necessarily indicate direct ssDNA binding. Instead, the RPAPs may provide a platform for protein:protein interactions. This is seen in eukaryotes, where the RPA 14 kDa subunit possesses a single OB-fold, this subunit is essential for formation of the RPA heterotrimer by facilitating protein:protein interactions (Fanning et al., 2006).

The co-purification of histidine-tagged RPA1 and RPA3 with their respective untagged RPAPs (and vice versa) supports our hypothesis that *H. volcanii* RPA1 and RPA3 form complexes with their respective RPAPs. This observation, and the differing outcomes of *rpa1*, *rpa2*, and *rpa3* deletions, indicates that the three RPAs of *H. volcanii* do not function as a heterotrimer. Similar

results have been obtained in *M. acetivorans*, where the three RPAs are able to bind ssDNA individually, in addition to stimulating primer extension by *M. acetivorans* DNA polymerase BI *in vitro*. (Robbins et al., 2004).

This study has shown genetically and biochemically that RPAPs interact with RPAs, and that this interaction is RPA-specific. This is the first report investigating the function of the archaeal COG3390 RPA-associated proteins (RPAPs), thus providing an important insight of the structure and function of *H. volcanii* single-strand DNA-binding proteins.

AUTHOR CONTRIBUTIONS

Amy Stroud and Thorsten Allers wrote the paper; Amy Stroud and Thorsten Allers designed the experiments; Amy Stroud and Thorsten Allers performed the microbiological and biochemical experiments; Susan Liddell carried out the mass spectrometry; Amy Stroud, Susan Liddell, and Thorsten Allers analyzed the data.

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APPENDIX

Table A1 | Identification of proteins present in cellular soluble fraction after purification by affinity chromatography on a Ni²⁺ chelating column.

| Prot accession | Protein name | HVO_# | Predicted MW | Observed MW | MASCOT score | Number of peptides | % coverage | Peptide sequences |
|----------------|---------------------------|-------|--------------|---------------|--------------|--------------------|------------|-------------------|
| gi 292655491] | RPAP1 | 1337 | 64,829 | 57,781 | 671 | 12 | 18 | 8 |
| gi 292655492] | RPA1 | 1338 | 45,954 | 36,960 | 940 | 15 | 40 | 11 |
| gi 292655491] | RPAP1 | 1337 | 64,829 | 62,780 | 578 | 10 | 21 | 9 |
| gi 292655492] | RPA1 | 1338 | 45,954 | 45,686 | 808 | 15 | 40 | 12 |
| gi 292654471] | RPAP3 | 0291 | 21,979 | 15,074 | 435 | 10 | 53 | 7 |
| gi 292654471] | RPAP3 | 0291 | 21,979 | 16,217 | 484 | 10 | 59 | 8 |
| gi 292654472] | RPA3 | 0292 | 34,562 | 28,262 | 796 | 13 | 43 | 10 |
| gi 292654472] | RPA3 | 0292 | 34,562 | 31,741 | 735 | 13 | 48 | 11 |
| gi 292656508] | Hypothetical protein | 2381 | 52,319 | 55,300/50,200 | 240 | 5 | 11 | 5 |
| gi 292493992] | Hypothetical protein | B0053 | 13,897 | 4,73,600 | 259 | 4 | 33 | 3 |
| gi 292656425] | Deoxyhypusine synthase | 2297 | 38,616 | 39,500 | 451 | 7 | 25 | 4 |
| gi 292655899] | Thioredoxin reductase | 1758 | 36,505 | 38,000 | 312 | 6 | 21 | 6 |
| gi 292493992] | Hypothetical protein | B0053 | 13,897 | 36,300 | 208 | 6 | 40 | 4 |
| gi 292656748] | Htr-like protein | 2629 | 30,266 | 31,700 | 93 | 4 | 17 | 4 |
| gi 292654704] | Ferritin | 0536 | 19,892 | 24,900 | 66 | 1 | 6 | 1 |
| gi 292653937] | Transcriptional regulator | A0388 | 20,201 | 22,300 | 586 | 15 | 54 | 8 |
| gi 292493992] | Hypothetical protein | B0053 | 13,897 | 13,500 | 170 | 3 | 33 | 3 |

Prot Accession, the database entry, e.g., gi|292655491; predicted MW, predicted molecular weight (Da) of the protein sequence identified by MASCOT; observed MW, molecular weight estimated from migration on SDS-PAGE; MASCOT score, MASCOT score associated with protein identification; number of peptides, no. of peptides associated with protein identification by MASCOT; % coverage, % of the database sequence entry that is covered by the peptides matched to it in the MASCOT data. Peptide sequences, the number of distinct peptide sequences associated with the protein identified by MASCOT.

