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Ribonucleolytic resection is required for repair of strand displaced NHEJ intermediates

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Abstract

Non homologous end-joining (NHEJ) pathways repair DNA double-strand breaks (DSBs) in eukaryotes and many prokaryotes, although not reported to operate in the third domain of life, archaea. Here, we describe a complete NHEJ complex, consisting of DNA ligase (Lig), polymerase (Pol), phosphoesterase (PE) and Ku from a mesophillic archaeon, *Methanocella paludicola* (*Mpa*). *Mpa* ligase has limited DNA nick sealing activity but is efficient at ligating nicks containing a 3’ ribonucleotide. *Mpa* Pol preferentially incorporates NTPs onto a DNA primer strand, filling DNA gaps in annealed breaks. *Mpa* PE sequentially removes 3’-phosphates and ribonucleotides from primer strands, leaving a ligatable terminal 3’-monoribonucleotide. These proteins, together with the DNA end-binding protein Ku, form a functional NHEJ break repair apparatus that is highly homologous to the bacterial complex. Although the major roles of Pol and Lig in break repair have been reported, PE’s function in NHEJ has remained obscure. We establish that PE is required for ribonucleolytic resection of RNA intermediates at annealed DSBs. Polymerase-catalyzed strand displacement synthesis on DNA gaps can result in the formation of non-ligatable NHEJ intermediates. PE’s function in NHEJ repair is to detect and remove inappropriately incorporated ribonucleotides, or phosphates, from 3’ ends of annealed DSBs to configure the termini for ligation. Thus, PE prevents the accumulation of abortive genotoxic DNA intermediates, arising from strand displacement synthesis, that would otherwise be refractory to repair.
NHEJ is a major DNA break repair pathway in eukaryotes and prokaryotes but assumed to be absent in archaea. This study establishes that a functionally homologous pathway is present in archaea. We have reconstituted archaeal NHEJ repair \textit{in vitro}, demonstrating that it is closely related to the bacterial apparatus and preferentially repairs breaks using RNA intermediates. We identify a role for a functionally unascribed nuclease in preventing the accumulation of genotoxic repair intermediates produced by strand displacement. This study has important implications for our understanding of the mechanisms of DNA break repair by NHEJ and the evolution of end-joining pathways.

\textbf{Introduction}

DNA double-strand breaks (DSBs) are one of the most lethal forms of damage encountered by cells (1). Non homologous end-joining (NHEJ) is the primary pathway for repairing DSBs in higher eukaryotes (2). NHEJ does not require the presence of a sister chromatid, in contrast to homologous recombination (HR), and can therefore operate in quiescent cells. The direct repair and ligation of DSBs by NHEJ is considered to be more error-prone as a consequence of mending breaks without the assistance of an intact DNA template.

Higher eukaryotic NHEJ complex is primarily composed of Ligase IV, XRCC4, XLF (LXX complex), DNA-PKcs, Ku70/80 (1, 3). Ku and DNA-PKcs both assist in bridging the gap between the broken termini by promoting end-synapsis. NHEJ repair of non homologous breaks requires remodelling of the DNA termini by processing enzymes, including polymerases (Pol \(\lambda\) and \(\mu\)) and nucleases (Artemis and Fen-1), to prepare them for ligation (4-7). The LXX complex is recruited by Ku, which enables the ligation of the DNA ends.
Most bacterial organisms also possess a NHEJ complex, composed of a more limited set of proteins, including Ligase D (LigD) and Ku (8, 9), which is required for DSB repair in stationary phase (10). The mycobacterial LigD gene encodes a multifunctional enzyme which, along with Ku, is capable of remodelling and ligating a broad variety of DSBs (8, 11). In many bacteria, LigD is found as a fusion of DNA ligase (Lig), polymerase (Pol) and phosphoesterase (PE) domains, in a variety of configurations. The NHEJ ligase (Lig or LigDom) has a strong propensity for ligating nicks with a ribonucleotide on the 3’ position of the break (12) and the polymerase (Pol or PolDom) preferentially incorporates NTPs, rather than the expected dNTPs (12, 13). Together, these findings establish that NHEJ repair requires polymerase insertion of ribonucleotides at the DNA termini for efficient ligation. The phosphoesterase (PE or NucDom) domain of LigD, originally shown to have 3’ exonuclease activity (8), preferentially removes ribonucleotides and phosphates from the 3’ termini strand, by sequential phosphodiesterase and phosphomonoesterase activities (14). Whilst PE’s biochemical activities are known, the specific role of this nuclease in DSBs repair remains unclear, with the nucleolytic activities of PE appearing to be antagonistic to those of the polymerase. Indeed, no obvious alterations to repair efficiency or fidelity was observed in plasmid-based DSB repair assays that excluded PE (15).

Archaea make up the third major domain of life and, although these organisms are one of the most abundant in the biosphere, relatively little is known about how they repair their DNA. Many archaeal species survive in some of the most extreme environments on the planet and therefore it is of importance to investigate
how they maintain genome stability to understand how life evolved in the hostile conditions present on the primordial earth and how they continue to enable organisms to thrive in extreme terrestrial niches. Although some NHEJ-like genes have been identified in archaea (9, 16-18), a complete DNA break repair apparatus has not been reported. Therefore, it is unclear if a related DSB repair pathway operates in archaea. Here, we describe the discovery and characterization of a complete NHEJ complex, composed of Lig, Pol, PE and Ku, in the archaeon, Methanocella paludicola (Mpa). This study establishes, for the first time, that a functional NHEJ apparatus exists in archaea that is highly homologous to the bacterial break repair complex. Furthermore, we have exploited the biochemically tractability of this NHEJ apparatus to identify and characterize an unexpected function of the PE nuclease activity in co-ordinating end-joining. We demonstrate that the phosphoesterase is required for ribonucleolytic resection of polymerase-incorporated RNA intermediates, resulting from strand displacement synthesis, at annealed DSBs. This unusual 3’ resection activity configures nicked NHEJ intermediates for optimal end-joining thus preventing the accumulation of potentially lethal non-ligatable intermediates that can arise from strand displacement synthesis by NHEJ polymerases.
RESULTS AND DISCUSSION

Identification of a putative NHEJ repair complex in archaea

NHEJ pathways operate in both eukaryotes and prokaryotes but the existence of a related DSB repair system exists in the third domain of life, archaea, is unclear. Although a few NHEJ-like genes have been identified in archaea, a complete repertoire of factors has not been discovered in any one species, suggestive that a functional end-joining repair pathway may not exist in this kingdom. To address this question, mycobacterial NHEJ protein sequences (Ku and LigD) were used to iteratively search for related sequences in archaeal genomes. We identified that the genome of a mesophilic archaean, *Methanocella paludicola* (Mpa) (19), possesses an apparently intact NHEJ apparatus. *Mpa* encodes NHEJ ligase (Mcp 2126), polymerase (Mcp 2125) and phosphoesterase (Mcp 2127) genes, located in an operonic arrangement, that share significant homology with the three distinct catalytic domains of mycobacterial LigD (Fig. 1A; Fig. S1). However, unlike LigD, the different putative activities are encoded on individual genes rather than fused together into a single LigD-like gene. *Mpa* also encodes the DSB end-binding protein, Ku (Mcp 0581), a “hallmark” gene for the presence of a *bona fide* NHEJ pathway. Together, these *Mpa* repair protein orthologues represent the first example of a complete NHEJ complex identified in an archaeal organism.

*Mpa* NHEJ proteins possess equivalent activities of prokaryotic orthologues

To allow biochemical characterization of the putative *Mpa* NHEJ proteins, the genes were individually over-expressed in *E. coli*, fused to a N-terminal hexa-histidine tag. The proteins were purified to homogeneity, in high yields, using immobilized metal
affinity chromatography (IMAC), ion exchange chromatography and gel filtration chromatography (see methods).

\textit{Mpa} Ku gene encodes a 252 amino acid polypeptide that shares significant sequence homology with the prokaryotic and viral Ku proteins. Following purification, \textit{Mpa} Ku was observed to migrate at \(~30\)kD by SDS-PAGE (Fig. S2) and eluted as an apparent homodimer from gel filtration columns, consistent with other Ku proteins (9, 20). Increasing titration of \textit{Mpa} Ku reduced the electrophoretic mobility of double-stranded (ds) DNA (Fig. 1B), confirming that it binds to ds DNA, consistent with other Ku proteins (13).

\textit{Mpa} DNA polymerase (\textit{Mpa} Pol) gene encodes a 295 aa protein (~34kD; Fig. S2) containing the three \textit{Archaeo-Eukaryotic Primase} (AEP) catalytic motifs present in other LigD-associated primase-polymerases (Fig. S3). In primer-template extension assays, \textit{Mpa} Pol incorporated either dNTPs or NTPs, in the presence of manganese, opposite a DNA template of a 5' overhang (Fig. 1C). Similarly to other bacterial NHEJ polymerases (13), \textit{Mpa} Pol displayed a marked preference for insertion of NTPs over dNTPs, with optimal activity observed in the presence of manganese ions, rather than magnesium (Fig. S2).

\textit{Mpa} phosphoesterase (PE) gene encodes a 198 aa protein, migrating \(~24\)kD in SDS-PAGE (Fig. S2). \textit{Mpa} PE represents a full-length, stand-alone, orthologue of the LigD PE domain. \textit{Mpa} PE cleaved 3'-ribonucleotides and 3'-phosphates from a recessed DNA-RNA primer strand, leaving a single 3' ribonucleotide containing a 3'
hydroxyl group (Fig. 1E), in common with the orthologous ribonucleolytic activity reported for other 3'-phosphoesterases (14, 18). As previously reported, this ability to remove all but the last ribonucleotide from a primer strand indicates the necessity for the phosphoesterase to recognise a 2'-hydroxyl ribose moiety on the penultimate base. *Mpa* PE required either manganese or cobalt as a cofactor for efficient catalytic activities. Copper, cadmium and magnesium all elicited moderate ribonuclease and phosphatase activities (Fig. S4).

*Mpa* DNA ligase (*Mpa* Lig) gene encodes a 334 aa protein, which migrated at ~40kD in SDS-PAGE (Fig. S2). *Mpa* Lig possesses all of the conserved motifs found in members of the ligase nucleotidyltransferase superfamily (21, 22) (Fig. S5). Notably, *Mpa* Lig had limited ligation activity on nicked DNA substrates (Fig. S6). In contrast, it preferentially catalyzed the ligation of nicked DNA containing at least one ribonucleotide on the 3'-hydroxyl (3'-OH) side of a nick (Fig. 1C). Ligation was optimal in the presence of manganese, with only limited end-joining in the presence of magnesium (Fig. S6). *Mpa* Lig was able to catalyze the ligation of nicks with a 5'-phosphate and a 3'-monoribonucleotide in the absence of ATP, suggesting that some of the recombinant protein was pre-adenylated, as observed previously (23).

Together, these data verify that the *Mpa* NHEJ proteins possess similar biochemical activities to their bacterial counterparts *in vitro* and suggests that they perform equivalent roles in NHEJ-mediated DSB repair processes in this species, implying that this DSB repair pathway also operates in archaea. This NHEJ apparatus provides the first tractable *in trans* model system to study how
components of the archaeo-prokaryotic (AP) NHEJ apparatus co-ordinate the repair of DSBs in vitro.

**Mpa Ku is required for DNA end-joining in vitro**

Ku’s proposed cellular function is to promote end-synapsis and recruit other NHEJ repair factors to the break. We therefore assayed the necessity for Mpa Ku to facilitate end-joining by Mpa Lig. The initial assay was performed on ds DNA containing a 3'-monoribonucleotide and a four base 5' DNA overhang with a self-annealing sequence to form a microhomology thus creating a doubly nicked duplex (Fig. 2A). Mpa Lig alone was not sufficient to ligate these breaks (Fig. 2A) however, incubation with Ku allowed successful ligation. The amount of product was proportional to the concentration of Ku added to the reactions. Mpa Lig's end-joining activity was also assayed using blunt-end ds DNA, containing a 3'-monoribonucleotide and a 5'-phosphate (Fig. 2B). Again, Mpa Lig alone was unable to ligate this substrate, however Ku stimulated its end-joining activity. Ligation of the blunt-ended DNA was not as efficient as that of the “synapsed” DNA, which is expected given the increased stability imparted by the presence of a region of microhomology.

Whilst it had been determined that Mpa Pol was not necessary for the physical act of connecting DNA ends to achieve ligation, a terminal ribonucleotide remained essential for ligation (Fig. S6). To ascertain whether Mpa Pol could efficiently fill a gap following break annealing, to promote ligation, we designed a DNA terminus with a 5 base 3' DNA overhang with a self-annealing sequence of four
bases (Fig. 2C). The non-complementary base requires gap-filling, in a template-dependent manner (S11A, S11C), subsequent to break annealing, before ligation can occur. These results indicate nucleotide incorporation by Mpa Pol and subsequent ligation of RNA/DNA nicks. The successful ligation of this substrate was again dependent on the presence of Ku for end-joining. Together, these data verify that Mpa Pol, Lig and Ku are bona fide NHEJ repair proteins and establish that a functional NHEJ complex exists in an archaeal species.

**Phosphoesterase removal of 3’-phosphate assists DNA gap-filling and ligation**

As discussed, Mpa PE possesses both phosphodiesterase and monophosphatase activities. PE is capable of removing a 3’-phosphate group from any DNA primer strand configuration (Fig. S7). This confirms that, unlike its ribonuclease activity, monophosphoesterase activity does not require a 2’-hydroxyl (OH) ribose moiety on the penultimate nucleotide. PE’s ability to remove 3’-phosphates, in a context that is different to its ribonuclease activity, suggests that PE’s phosphatase activity may participate in distinct processing events during end-joining. The likely role of the phosphatase is to facilitate 3’ phosphate removal at damaged termini to expose a free 3’-OH, thus allowing the DNA termini to be receptive to subsequent modifications by the polymerase and ligase whose activities are dependent on the availability of a OH group.

To scrutinize this further, we incubated a single nucleotide gapped DNA substrate, containing both 3’ and 5’-phosphate groups, with Mpa Pol, Lig and PE (+/-) (Fig. 2D). Double-stranded DNA with gaps are frequently used as substrates for
NHEJ enzymes as they mimic annealed breaks formed during DSB repair (13, 24). As predicted, the presence of a 3'-phosphate prevented NTP incorporation by *Mpa* Pol (Fig. 2D), due to the lack of a 3'-OH required for nucleophilic attack during extension. *Mpa* Lig was also unable to ligate this DNA, as ligation mechanisms also requires a free 3'-OH. This impediment was overcome by the addition of PE, which catalysed removal of the 3'-phosphate producing a phosphate shift (annotated as n-OH; Fig. 2D) followed by subsequent gap-filling and ligation. These data demonstrate that *Mpa* PE performs additional DNA modification tasks in NHEJ, aside from its ribonuclease function, thus acting as a phosphatase that can operate at early stages of DSB repair to facilitate further processing and ligation of the break termini.

**Mpa** Pol, a non-processive polymerase with strand displacement activity

As discussed, *Mpa* Pol incorporated NTPs onto a primer strand in a template-dependent, but non-processive, manner (Fig. 1D) *Mpa* Pol inserted only 3-4 NTPs opposite a template strand (19 bases long) and efficiency was even further reduced with dNTPs, as noted above. This limited extension may result from the formation of DNA/RNA hybrids, switching the DNA conformation from B- to A-form, thus encouraging polymerase dissociation (25). *Mpa* Pol is also capable of catalyzing non-templated NTP incorporation on blunt-ended DNA, another activity shared with bacterial NHEJ polymerases (Fig. S8) (13).

To determine if *Mpa* Pol displays strand displacement activity following gap-filling (13, 26), we measured incorporation on a single nucleotide gap DNA substrate.
Although *Mpa* Pol efficiently filled in the gap, it also synthesised additional longer products, incorporating up to three nucleotides (Fig. 3A). This implies that, following gap-filling, *Mpa* Pol proceeds to incorporate further NTPs, presumably by displacing the downstream (D) strand. Notably, when *Mpa* Pol was assayed with a nicked DNA substrate (Fig. 3B), containing no gap, it also inserted up to 3 nucleotides. Even with nicked DNA, the NTP was incorporated in a template-dependent manner (Fig. S11B). Strand displacement synthesis is also intrinsic to the mycobacterial NHEJ polymerases (*Mtu* PolDom, Fig. S9) and has also been described in other DNA repair pathways (27). It is unclear what precise role displacement synthesis plays during NHEJ repair, at least in the absence of a 5′ exonuclease, which could resect the displaced DNA flap. However, in addition to gap-filling, these polymerases also play important roles in end-synapsis, by directly facilitating the annealing of break termini (24). Therefore, Pol may maintain intrinsic displacement activity to enable it to open up breaks with limited complementarity, e.g. blunt ends. Pol may ingress into the termini, to uncover regions of microhomology that can then be annealed together to facilitate end-synapsis (24). It may also enable these enzymes to locate an internal 5′ phosphate, if a terminal one is not present, to allow it to bind more securely to a DSB (24, 28). However, although it may be beneficial for some purposes, strand displacement can lead to some undesirable consequences, discussed below.

**Phosphoesterase removes ribonucleotides from annealed DNA breaks**

Although PE resects terminal ribonucleotides from recessed 3′ primer strands (Fig. 1D), the specific role of this nucleolytic activity in processing NHEJ intermediates has
remained elusive. Given that polymerase incorporation of NTPs is the likely source for the presence of small tracts of RNA at sites of DSBs, we asked if PE’s ribonuclease activity is required for processing these DNA-RNA intermediates? To address this question, we assayed PE’s ability to resect an annealed DNA break substrate containing RNA, a diribonucleotide at the 3’ end of the primer strand, and a D-strand. Notably, PE readily removed NMPs from the 3’ DNA-RNA primer strand on this “annealed” break, leaving a single 3’ ribonucleotide. Significantly, PE’s resection activity was far more efficient than on an equivalent 3’ primer lacking an adjacent D-strand (Fig. 3C, 3D).

Given the striking stimulation of PE’s resection activity in the presence of a D-strand, we assayed a variety of different D-strands (ranging from a 2 base 5’ flap (+2) to a 2 base (-2) gap) to identify PE’s preferred substrate (Fig. 3E). A nicked substrate was the most efficiently processed intermediate, with almost 100% resection observed. A one base 5’ flap (+1) and gapped substrate were processed less efficiently but considerably better than the remaining substrates (Fig. 3E). Notably, the activity of PE decreased drastically if the gap was larger than +1, whereas a 2 base 5’ flap did not greatly suppress activity. These data demonstrate that PE is most efficient at resecting RNA incorporation at annealed breaks with little or no gap, even preferring to remove RNA that is displacing DNA rather than from a gap of 2, or more, bases.

Together, these data establish that PE preferentially resects RNA from the 3’ primer strand when a D-strand is present, and not from a recessed 3’ strand of a free
end, as previously reported (14, 29, 30). This observed preference for 3’ resecting RNA from ‘nicked’ DNA supports a model in which PE specifically operates, not on break termini, but after DSB synapsis and gap filling has already occurred.

**PE regulates incorporation of RNA at annealed breaks**

The roles of NHEJ Pol and PE are antagonistic, as one inserts ribonucleotides at DSBs whilst the other one removes them. It is therefore critical to understand the context and co-ordination of these opposing end-processing activities to delineate how they co-operate to repair DSBs. As shown, PE prefers to resect RNA from the 3’ end of primer strands at annealed breaks, indicating that it may be required to modulate the length of extension products incorporated during gap-filling. To test this model, a fixed concentration of PE was incubated with an increasing concentration of Pol and two different DNA substrates (+/- D-strand) containing a 1-nucleotide gap to assess the effects on NTP incorporation, if any (Fig. 4A, 4B). Notably, PE limited the amount of NTP incorporation, at approximately equimolar concentrations, on both substrates. PE significantly limited NTP incorporation in the presence of a D-strand, with the major product at +1, even at saturating Pol concentrations (Fig. 4B). These data establish that PE directly modulates the length of extension of primer strands at annealed DSBs by resecting the 3’ RNA products incorporated by NHEJ polymerases.

**PE resection of strand displacement intermediates during NHEJ repair**

Although NHEJ polymerases are proficient at filling in gaps formed at annealed breaks (24), they also catalyse, less desirably, strand displacement synthesis on
gapped substrates. This is a particularly dangerous activity to possess given that it can lead to the formation of unligatable intermediates, containing a 5' flap, that are refractory to repair. A mechanism is therefore required to minimize the production of such abortive genotoxic intermediates, allowing end-ligation to proceed in a favourable way.

Given that PE regulates the amount of nucleotides incorporated by Pol into gapped NHEJ substrates, we assessed if PE provides a mechanism for modulating strand displacement synthesis thus preventing the formation of unligatable DNA intermediates (Fig. 4C, left panel). A single nucleotide gapped substrate was pre-incubated with Mpa Pol (0, 15, 30 and 60 minutes), with manganese and NTPs, before addition of Mpa PE and Lig. Whilst NTP incorporation by Pol remained at +3 or +4 across the time course, the amount of ligated product reduced dramatically over time. For example, after 15 minutes Mpa Pol had catalyzed almost total strand displacement synthesis, rendering most of the DNA inert to ligation (Fig. 4C, left panel). However, addition of Mpa PE (after 15, 30 or 60 minutes) rescued a considerable proportion of the non-repairable DNA (Fig. 4C, left panel), presenting Mpa Lig with a nicked substrate optimised for ligation. Indeed, addition of PE directly following addition of Pol (0 min; Fig. 4C) yielded a much greater quantity of ligated substrates than present in reactions lacking it. Similar results were also observed for Mtu PolDom (Fig. S10). End-joining substrates with 3’ overhangs were also assayed as described above (S12A, S12B) and the results are consistent with the gap-filling and ligation assays in figure 4C, with PE counteracting Pol’s downstream displacement activity. Together, these data robustly support a model in which PE is
required to optimally resect RNA-containing NHEJ intermediates, particularly those produced by strand displacement synthesis, to ensure the remodelled annealed break is amenable to ligation (Fig. 4C, right panel). The availability of this novel editing mechanism may be particularly important for ensuring the repair of all DSBs as even a single unrepaired DNA break can be lethal.

**Concluding remarks**

The potentially lethal nature of DNA double-strand breaks has ensured that all organisms have evolved repair pathways to effectively mend these potent lesions. Although recombination-based repair mechanisms are conserved across all living organisms, NHEJ break repair pathways were considered to operate exclusively in eukaryotes. However, in recent years it has been recognized that most bacterial species also utilize a closely related NHEJ repair apparatus to mend breaks. In contrast, there was limited evidence to support the existence of NHEJ repair pathways in the third major domain of life, archaea. This is particularly surprising given that many archaeal species thrive in some of the harshest environmental niches, e.g. deep sea thermal vents, significantly putting their genomes at increased risk of breakage. This current study resolves this apparent discrepancy with the discovery of a fully functional NHEJ repair apparatus in archaea thus establishing the conservation of this pathway across all three domains of life.

Although many archaeal DNA metabolism pathways, e.g. DNA replication, share many more common genes with eukaryotes rather than with bacteria, archaeal NHEJ is much more homologous to the bacterial DSB repair complexes, particularly
in terms of its conservation of the key repair proteins and their common intrinsic biochemical activities. Therefore, these closely related break repair pathways can be grouped together as archaeo-prokaryotic (AP) NHEJ to differentiate it from the more divergent eukaryotic pathway. Notably, a complete LigD AP-NHEJ apparatus is present in some plants, e.g. castor bean, suggesting that this pathway diverged quite late in evolution. The discovery of a shared end-joining repair apparatus in both archaeal and prokaryotic organisms further supports the hypothesis that a primordial NHEJ repair pathway arose early in evolution, was maintained and further evolved in lower eukaryotes, before eventually becoming the predominant DSB repair pathway in mammalian cells.

Although bacterial LigD has proven invaluable in providing insights into AP-NHEJ repair mechanisms, it also has some major experimental limitations. However, unlike LigD, the Mpa archaeon end-joining machinery is not fused into a single protein. This physical disconnectivity of the NHEJ activities has provided a tractable in trans repair apparatus that now allows dissection of the interplay between the end-processing enzymes to elucidate how they co-operate together to co-ordinate break repair. Although the AP-NHEJ phosphoesterase family was identified over a decade ago, its specific role in break repair has remained elusive. The modularity of the archaeal end-joining complex has enabled us to delineate a major function for PE in break repair. We demonstrate that the propensity of NHEJ polymerases to insert excessive ribonucleotides into gaps by strand displacement synthesis, following end-synapsis, is strongly counteracted by the resection activity of PE. NHEJ phosphoesterases limit excessive ribonucleotide incorporation by using a
ribonucleolytic resection mechanism, thus ensuring that nicked DNA substrates are available for ligation, preventing the accumulation of potentially genotoxic NHEJ intermediates (Fig. 5). The principle of strand displacement in DNA repair is not novel and, in fact, is well established in the eukaryotic base-excision repair (BER) pathway (27). Pol β actively displaces downstream damaged DNA as an interim repair step of BER, before FEN-1 endonucleolytically resects the 5’ damaged flap. The PE activity associated with AP-NHEJ offers an alternative resolution of strand displaced intermediates. Resection of the newly synthesised ingressing DNA strand by PE facilitates reannealing of the displaced strand to produce a nicked template that is optimal for end-ligation.

Replicative DNA polymerases possess a 3’ proofreading exonuclease (Exo) function to maintain fidelity. AP-NHEJ polymerases also maintain an associated 3’-Exo activity but to limit displacement synthesis rather than for maintaining fidelity. This functional association of these atypical nucleases with Pol and Lig is supported by the genomic distribution of PE genes, which are almost exclusively found in an operonic arrangement with Pol and Lig genes, indicating that a selective genetic requirement maintains PE in association with these NHEJ processing enzymes. PE-like genes also exist in some yeast species although their role is unknown.

An intriguing feature of AP-NHEJ repair pathways is the preferential insertion, removal and ligation of ribonucleotides at DSBs by the break repair enzymes. Utilisation of ribonucleotides during break repair is a concept of growing recognition with reports of preferentially incorporation of ribonucleotides during eukaryotic NHEJ
(31, 32), suggesting that RNA-DNA intermediates are not limited to AP-NHEJ (33). There are several reasons why it may be favourable to incorporate RNA during DSB end-joining. NHEJ is the DSB repair pathway of choice for non-dividing cells (10), where NTPs are much more abundant than dNTPs in the available nucleotide pools (33). Another distinct possibility, driven by the availability of ribonucleotides, is that NHEJ repair pathways evolved to selectively incorporate RNA, over DNA, into repaired DSBs to demarcate the boundaries of any alterations made to the repaired sites. This “labelling” of the repaired break with ribonucleotides may be useful in a number of ways. Firstly, It prevents the loss of genetic material at sites of DSBs by preventing resection of the DNA termini by the ribo-centric NHEJ repair machinery. Significantly in this regard, PE cannot resect DNA ends and prefers to resect short RNA tracts incorporated by the polymerase. Secondly, efficient ligation of breaks is also predicated on incorporation of a terminal 3’ ribonucleotide that, potentially, acts as a molecular signal to indicate when a nick is ready for ligation thus preventing aberrant end-joining. A major consequence of AP-NHEJ repair is the incorporation of, at least, one ribonucleotide into ligated DSB junctions and it is likely that this RNA is subsequently excised from the repaired breaks (33, 34). Studies in murine systems have revealed that RNase HII performs an essential role in ribonucleotide removal from genomic DNA (35). Whilst this role is thought largely to protect from RNA misincorporated during replication, RNase HII (or a similar enzyme) may also resolve the final complication of AP-NHEJ repair but this remains to be established.
Methods

Purification of *Mpa* Lig, Pol, PE and Ku proteins

All *Mpa* NHEJ genes were cloned into pET 28A, the histidine-tagged proteins expressed in B834S (DE3) and purified using the same chromatography protocols, except where noted. Pelleted cells containing the appropriate plasmid were grown in Terrific Broth and pelleted cultures resuspended in lysis buffer (50mM Tris-HCl pH7.5, 500mM NaCl, 30mM Imidazole, 10% glycerol, 17mg/mL PMSF, 34mg/mL benzamidine). Cells were disrupted by sonication and soluble proteins were isolated by centrifugation. The supernatant was applied to a 25mL NTA (Qiagen) column equilibrated in IMAC buffer A (as lysis buffer) and eluted in IMAC buffer B (as A, with 300mM imidazole). *Mpa* Ku supernatant was subjected to an extended wash period in IMAC buffer C (as A, with 1M NaCl) to remove excess DNA bound to Ku proteins. Protein containing fractions were loaded onto 5mL fast flow Q and S (Lig, Pol), or Q (PE and Ku) columns (GE Healthcare) pre-equilibrated in IEx buffer A (50mM Tris-HCl pH 7.5, 10% glycerol) for separation by ionic charge. Lig, Pol and PE were eluted in the flow through without binding the column, Ku was eluted in 10% IEx buffer B (50mM Tris-HCl pH7.5, 2M NaCl, 10% glycerol). Removal of any remaining contaminating proteins was achieved by gel filtration on a Superdex S200 column (GE Healthcare) equilibrated in GF buffer (25mM Tris pH 7.5, 500mM NaCl, 10% glycerol). Eluted proteins were analysed for purity using SDS-PAGE and concentrations ascertained by spectrophotometry of samples at 280nm.

EMSA

Fluorescein labelled DNA primer (5'-CATATCCGTGTCGCCCTTATTCCGATAGTGACTACA) was annealed to DNA template (5'-TGTAGTCACTATCGGAATAAGGGGCACACGGATATG) creating Oligo 20 for DNA binding assays. 60nM DNA was incubated with 0.2-1.6µM *Mpa* Ku in 50mM Tris pH 7.5 and 5% glycerol for 30 minutes at room temperature in a volume of 20µL. Samples were separated by electrophoresis on a native 5%
polyacrylamide gel in 0.5x TBE buffer for 2hr. Fluorescently labelled DNA oligomers were detected by scanning using a Fujifilm FLA-5100.

**DNA ligation assay**

Fluorescein labelled DNA primer (5'-CTATGAGCGAATCGCrC) was annealed to DNA template (5'-AGTCGCATAGTGAGTCGGGGCGATTCGCTCATAG) with a downstream sequence (5'-P-CGACTACACTATGCAACT) creating Oligo 60. 30nM DNA was incubated with 300nM Mpa Lig in 50mM Tris HCl pH 7.5, 5mM MnCl₂ for 30, 60 and 90 minutes at 37ºC in a volume of 20µL. The reactions were stopped by addition of Stop buffer (95% formamide, 0.09% xylene cyanol) and boiled at 95ºC for 10 minutes. Samples were separated by electrophoresis on an 8M Urea, 15% polyacrylamide gel in 1xTBE buffer for 2hr. Fluorescently labelled oligos were detected by scanning using a Fujifilm FLA-5100.

**DNA extension assay**

Fluorescein labelled DNA primer (5'-CTATGAGCGAATCGCC) was annealed to DNA template (5'-AGTCGCATAGTGAGTCGGGGCGATTCGCTCATAG) creating Oligo 9 for extension. To this DNA oligo, further downstream sequences were annealed (5'-CCGACTACACTATGCAACT, 5'-P-CGACTACACTATGCAACT) creating Oligos 11 and 37 for gap filling assays. 30nM DNA was incubated with 300nM Mpa Pol in 50mM Tris HCl pH 7.5, 5mM MnCl₂, 250µM NTP mix for 1hr at 37ºC in a volume of 20µL. The reactions were stopped by addition of Stop buffer (95% formamide, 0.09% xylene cyanol) and boiled at 95ºC for 10 minutes. Samples were separated by electrophoresis on an 8M Urea, 15% polyacrylamide gel in 1xTBE buffer for 2hr. Fluorescently labelled oligos were detected by scanning using a Fujifilm FLA-5100.
Ribonuclease and phosphatase assay

Fluorescein labelled DNA-RNA primer (5’-CTATGAGCGAATCGrCrC) was annealed to DNA template (5’-AGTCGCATAGTGTAGTCGGGGCGATTCGTCATAG) creating Oligo 3 for phosphoesterase assays. To this oligo further downstream sequences were annealed (5’-CTACCGACTACACTATGC GACT 5’-CCGACTACACTATGC GAC, 5’-P-CGACTACACTATGC GACT 5’-GACTACACTATGC GACT) creating oligos 25, 62, 2, 38 and 43 respectively. 30nM DNA was incubated with 300nM Mpa PE in 50mM Tris pH 7.5 and 5mM MnCl₂ at 37°C for varying time points across 90 minutes. The reactions were quenched by addition of Stop buffer buffer (95% formamide, 0.09% xylene cyanol) and boiled at 95°C for 10 minutes. Samples were separated by electrophoresis on an 8M Urea, 15% polyacrylamide gel in 1xTBE buffer for 2hr. Fluorescently labelled oligos were detected by scanning using a Fujifilm FLA-5100.

DNA end joining assay

Fluorescein labelled DNA and DNA-RNA primers (5’-CATATCCGTGTCGCCCTTTATTCGATAGTGACTArC 5’-CATATCCGTGTCGCCCTTTATTCGATAGTGACTCrA 5’-CATATCCGTGTCGCCCTTTATTCGATAGTGACTACGCG) were annealed to DNA templates (5’-P-CATGGTAGTCACACTATC GAGATAAGGGCGACACCGGATAT 5’-P- TGTAGTCACACTATCGGATAAGGGCGACACCGGATAT) to form Oligos 45, 68 and 76 respectively, for DNA end joining assays. 300nM Mpa Lig was incubated with 200-800nM Mpa Ku, and 300nM Mpa Pol where indicated, in 50mM Tris pH 7.5, 5mM MnCl₂, 1mM DTT, 5% glycerol, 0.1mgml⁻¹ BSA, 250μM NTPs (Pol assay only) at 37°C in a final volume of 20μL. The reactions were quenched by addition of Stop buffer (95% formamide, 0.09% xylene cyanol) and boiled at 95°C for 10 minutes. Samples were separated by electrophoresis on an 8M Urea, 15% polyacrylamide gel in 1xTBE buffer for 2hr. Fluorescently labelled oligos were detected by scanning using a Fujifilm FLA-5100.
Phosphatase, gap filling and ligation assay

Fluorescein labelled DNA primer (5'-CTATGAGCGAATCAGC-P) was annealed to DNA template (5'-AGTCGAGTCAGTGTCGGGGCGATTCGCTCATAG) and downstream strand (5'-P-CGACTACACTATCGACT) creating Oligo 63 for phosphatase, gap filling and ligation. 300nM of Mpa Lig, Pol and PE were incubated where indicated with 50mM Tris HCl pH 7.5, 5mM MnCl$_2$, 250µM NTP mix for 1hr at 37°C in a volume of 20µL. The reactions were quenched by addition of Stop buffer buffer (95% formamide, 0.09% xylene cyanol) and boiled at 95°C for 10 minutes. Samples were separated by electrophoresis on an 8M Urea, 15% polyacrylamide gel in 1xTBE buffer for 2hr. Fluorescently labelled oligos were detected by scanning using a Fujifilm FLA-5100.

Polymerase DNA extension, gap-filling and phosphoesterase assay

Oligos 9, 11 and 37, as described above, were used for DNA extention and phosphoesterase assays. 30nM DNA was incubated with 300nM Mpa Pol and PE where indicated in 50mM Tris pH 7.5 and 5mM MnCl$_2$ at 37°C for varying time points across 90 minutes. The reactions were quenched by addition of Stop buffer buffer (95% formamide, 0.09% xylene cyanol) and boiled at 95°C for 10 minutes. Samples were separated by electrophoresis on an 8M Urea, 15% polyacrylamide gel in 1xTBE buffer for 2hr. Fluorescently labelled oligos were detected by scanning using a Fujifilm FLA-5100.

DNA gap-filling, phosphoesterase and ligation assay

Oligo 37, as described above, was used for gap-filling, phosphoesterase and ligation assays. 30nM DNA was incubated with 300nM Mpa Pol in 50mM Tris pH 7.5 and 5mM MnCl$_2$ at 37°C for 0, 15, 30 and 60 minutes. 300nM Mpa Lig and PE were then added and incubation continued for a further 60 minutes. The reactions were
quenched by addition of Stop buffer buffer (95% formamide, 0.09% xylene cyanol) and boiled at 95°C for 10 minutes. Samples were separated by electrophoresis on an 8M Urea, 15% polyacrylamide gel in 1xTBE buffer for 2hr. Fluorescently labelled oligos were detected by scanning using a Fujifilm FLA-5100.
Acknowledgements

We declare that none of the authors have a financial interest related to this work.

Work in the AJD laboratory is supported by grants from Biotechnology and Biological Sciences Research Council (BBSRC) and indirectly by a centre grant from the MRC. EB was supported by a BBSRC PhD studentship. We would like to thank Dr H. Imachi for providing Methanocella paludicola genomic DNA.

Conflict of interest

All authors declare no conflicts of interest.
References


Figure legends

Figure 1. Biochemical activities associated with *Mpa* NHEJ repair proteins. (A) Comparison of gene arrangement in a NHEJ operons from *Methanocella paludicola* (*Mpa*) and *Mycobacterium tuberculosis* (*Mtu*). *Mpa* NHEJ factors are expressed as discrete proteins, whilst *Mtu* LigD is a multi-domain single polypeptide. Ku is present in both genomes, and is operonic with LigD in *Mtu*. (B) Ku DNA binding assays, 100nM 5′-fluorescein labelled 33mer dsDNA was incubated with 0, 200, 400, 800 and 1600nM *Mpa* Ku protein. (C) Ligase reactions contained 30nM 5′-fluorescein labelled 16mer DNA-RNA (D15R1) annealed to template DNA and 5′-phosphorylated D-strand to create a nicked substrate, and 300nM *Mpa* Lig protein. Ligated product is 35 bases in length. (D) Polymerase primer extension reactions contained 30nM 5′-fluorescein labelled 16mer DNA annealed to template DNA, and 300nM *Mpa* Pol protein. Incubations included 250µM NTPs/dNTPs and 5mM MnCl$_2$ or MgCl$_2$ where indicated. (E) Phosphoesterase reactions contained 30nM 5′-fluorescein labelled 16mer DNA/RNA (D14R2) annealed to template DNA, and 300nM *Mpa* PE protein. n-R-R is unmodified substrate, n-R-P represents removals of the terminal RMP to yield a terminal phosphate group, and n-R-OH is the product following phosphate removal leaving a terminal hydroxyl group. All reactions performed in the presence of manganese (5mM).

Figure 2. *Mpa* NHEJ complexes repair a diverse range of DNA breaks *in vitro*. (A) End-joining reaction mixtures contained 30nM 5′-fluorescein labelled 36mer DNA-RNA (D35R1) annealed to 5′-phosphorylated 40mer template DNA to create self-complementary 4mer overhang, with 300nM *Mpa* Lig and 200-800nM *Mpa* Ku proteins. Ligated products were 76 bases in length. (B) End-joining reaction mixtures contained 30nM 5′-fluorescein labelled 36mer DNA-RNA (D35R1) annealed to 5′-phosphorylated 36mer template DNA to create blunt ended 5′-phosphorylated DNA, with 300nM *Mpa* Lig and 200-800nM *Mpa* Ku proteins. Ligated products were 72 bases in length. (C) End-joining reaction mixtures contained 30nM 5′-fluorescein labelled 42mer DNA annealed with 5′-phosphorylated 37mer template DNA to create 5mer 3′-overhang DNA, four bases of which are self-complementary, with 300nM *Mpa* Lig and Pol and 200-800nM *Mpa* Ku proteins. Ligated products were 79 bases
in length. (D) 3'-phosphatase, gap filling and ligation reaction mixtures contained 30nM 5'-fluorescein labelled 16mer with a 3'-phosphate annealed to template DNA and 5'-phosphorylated D-strand, with 300nM Mpa Lig, Pol and PE where indicated. n-P indicates unmodified substrate, n-OH indicates removal of 3'-phosphate to reveal 3'-OH which can then receive incoming NTPs for n+1. All reactions performed in the presence of manganese (5mM).

**Figure 3. Characterization of polymerase strand displacement and phosphoesterase activities.** (A, B) Gap filling reactions mixtures contained 30nM 5'-fluorescein labelled 16mer DNA annealed to template DNA and D-strands to create 1 nucleotide gap DNA (A) or nicked DNA respectively (B), and 300nM Mpa Pol protein. (C, D, E) Phosphoesterase reaction mixtures contained 5'-flourscein labelled DNA-RNA (D14R2) annealed to a template with no D-strand (C), with D-strand (D), with D-strands producing 2 and 1 base nucleotide 5' flaps and 1 and 2 nucleotide gaps in respect to the DNA-RNA primer (E) and 300nM Mpa PE. Products as described in Fig. 1E. All reactions performed in the presence of manganese (5mM).

**Figure 4. PE resection of strand displacement intermediates is required for NHEJ repair.** (A, B) Polymerase and phosphoesterase reaction mixtures contained 5'-fluorescein labelled 16mer DNA primers annealed to a template with no D-strand (A) and with a 1 nucleotide gap (B), and 300nM Mpa PE and 0.1-1.4µM Mpa Pol. (C, left panel) Gap filling, phosphoesterase and ligation reaction mixtures contained 5'-fluorescein labelled 16mer DNA annealed to template DNA and a D-strand producing a single nucleotide gap, 300nM Mpa Lig, Pol and PE where indicated. Pre-incubated with Mpa Pol at 37°C for 0, 15, 30 and 60 minutes before addition of Mpa Lig and PE and incubation at 37°C for 1hr. All reactions performed in the presence of manganese (5mM). (C, right panel) Illustration of rescue of abortive NHEJ repair intermediates which cannot be ligated. Downstream DNA displaced by incorporation of NTPs prevents ligation, however Mpa PE resects the RNA tract and leaves a monoribonucleotide. Downstream DNA can then reanneal and the substrate is ready for ligation with the 5'-phosphate opposite the 3'-hydroxyl group.
**Figure 5. Model of AP-NHEJ using RNA repair intermediates.** The initial step in AP-NHEJ repair is the binding of Ku homodimers to broken ends and bringing them into close proximity. Ku and Pol promote microhomology-mediated synapsis of overhanging termini, if present, to anneal the break. Pol fills in any resulting gaps, at either end of the annealed break, by a template-dependent RNA synthesis. Pol may displace several bases of downstream DNA using its distinct ability to mobilise and direct DNA strands, offering a greater flexibility to repair a variety of configurations of damaged DNA. Strand displacement RNA synthesis by Pol is regulated by PE, which can remove unnecessary NMPs and allow the displaced DNA to realign with the template. PE always leaves a single terminal 3’ ribonucleotide in place in resected breaks, presenting Lig with an optimised nicked substrate for ligation. Although the broken DNA is reconnected, repair may not be fully completed as several RNA bases still resided in the duplex. RNA removal may be performed by additional enzymes, such as RNase HII, before other repair polymerases fill in the gaps with DNA.
Figure 1
Figure 2
Figure 3
Figure 5

AP-NHEJ

Double Strand Break

Break annealed

dsDNA

RNA removal?

Pol

Ku

RNA insertion

RNA

Ligation

RNA optimization

PE

Alternative repair?
Supporting information:
Ribonucleolytic resection is required for repair of strand displaced NHEJ intermediates, Edward Bartlett, Nigel C. Brissett and Aidan J. Doherty

S1. A, B, C & D Protein sequence alignments of Mpa Lig, Pol, PE and Ku with Mtu, Msm, Pae LigD and Ku.  
(A) Methanocella paludicola (Mpa) Ligase protein sequence alignment with Ligase D (LigD) ligase domains (LigDom) of Mycobacterium tuberculosis (Mtu), Mycobacterium Smegmatis (Msm) and Pseudomonas Aeruginosa (Pae).  
B. Mpa polymerase (Pol) protein sequence alignment with LigD polymerase domains (PolDom) of Mtu, Msm and Pae.  
C. Mpa phosphoesterase (PE) protein sequence alignment with LigD phosphoesterase domains (PEDom) of Mtu, Msm and Pae.  
D. Mpa Ku protein sequence alignment with Ku of Mtu, Msm and Pae. All alignments were made with TCoffee and edited in JalView.

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SDS-PAGE of gel filtration chromatography fractions of respective purifications of Mpa Lig, Pol, PE and Ku. Migration sizes of the proteins were: Mpa Lig ~40kD, Mpa Pol ~34kD, Mpa PE ~24kD and Mpa Ku ~30kD (monomer) ~60kD (dimer). The gels indicate the high purity of the proteins.

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Three key conserved motifs (I, II and III) confer membership to the AEP superfamily. Here Mpa Pol is aligned with Mycobacterium tuberculosis and Pseudomonas Aeruginosa NHEJ Pols, and Homo sapiens, Mus musculus, Drosophila melanogaster, Saccharomyces cerevisiae and Schizosaccharomyces pombe primase small sub-units.

S4. Mpa PE ion preference.  
Phosphoesterase reaction mixtures contained 50mM Tris pH 7.5, 5mM MnCl$_2$, MgCl$_2$, CoCl$_2$, CuCl$_2$, CaCl$_2$, NiCl$_2$ or Zn(O$_2$CCH$_3$)$_2$ where indicated, 30nM 5'-fluorescein labelled 16mer DNA/RNA (D14R2), and 300nM Mpa PE protein. Incubated at 37°C for 1 hour 30 minutes and electrophoresed on a 15% denaturing polyacrylamide gel.

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Six key conserved motifs (I, Ia, III, IIIa, IV and V) confer membership to the polynucleotide ligase and mRNA capping superfamily. Motif VI is also included with highly conserved residues. Here Mpa Lig is aligned
Mycobacterium tuberculosis and Pseudomonas aeruginosa Ligase D; Enterobacterio phage T7, Homo sapiens, Mus musculus DNA ligase; and Homo sapiens, Mus musculus, Drosophila melanogaster, Saccharomyces cerevisiae and Schizosaccharomyces pombe mRNA capping enzymes.

S6. Mpa Lig preferable ligation with Mn, not Mg
Ligase reaction mixtures contained 50mM Tris pH 7.5, 5mM MnCl$_2$ or MgCl$_2$, 30nM 5'-fluorescein labelled 16mer oligomer (D16, D15R1,D14R$_2$, D12R4 or D8R8 where indicated) and 19mer 5'-phosphorylated DNA, and 300nM Mpa Lig protein. Incubated at 37°C for 2 hours and electrophoresed on a 15% denaturing polyacrylamide gel.

S7. Mpa PE phosphomonoesterase activity does not require 2'-OH
Phosphatase reaction mixtures contained 50mM Tris pH 7.5, 5mM MnCl$_2$, 30nM 5'-fluorescein labelled 16mer DNA with 3'-phosphate, and 250, 500 and 1000nM Mpa PE protein. Incubated at 37°C for 1 hour 30 minutes and electrophoresed on a 15% denaturing polyacrylamide gel.

S8. Mpa Pol non-templated NTP incorporation
Polymerase non-templated extension reaction mixtures contained 50mM Tris pH7.5, 5mM MnCl$_2$, 62.5µM ATP, CTP, GTP, UTP or 250µM mixed NTPs where indicated, 30nM 5'-fluorescein labelled 36mer DNA, and 300nM Mpa Pol protein. Incubated at 37°C for 1 hour and electrophoresed on a 15% denaturing polyacrylamide gel.

S9. Mtu PolDom strand displacement
Polymerase strand displacement reaction mixtures contained 50mM Tris pH 7.5, 5mM MnCl$_2$, 250µM NTPs, 30nM 5'-fluorescein labelled 16mer DNA, and 300nM Mpa Pol or Mtu PolDom protein where indicated. Incubated at 37°C for 1 hour and electrophoresed on a 15% denaturing polyacrylamide gel.

S10. Mpa PE can rescue NHEJ intermediates formed by Mtu PolDom and Mpa Pol.
Gap-filling, phosphoesterase and ligation reaction mixtures contained 50mM Tris pH 7.5, 5mM MnCl$_2$, 250µM NTPs, 30nM 5'-fluorescein labelled 16mer DNA, and 300nM Mpa Pol or Mtu PolDom, 300nM Mpa Lig and PE protein where indicated. Incubated at 37°C for 1 hour and electrophoresed on a 15% denaturing polyacrylamide gel.

S11. Mpa Pol gap filling and strand displacement at nicks is template dependent.
Polymerase gap filling and strand displacement reaction mixtures contained 50mM Tris pH 7.5, 5mM MnCl₂, 62.5µM of the indicated NTP, 30nM 5’-fluorescein labelled 16mer DNA (A, B) and 47mer DNA (C), and 300nM Mpa Pol protein. Incubated at 37°C for 1 hour 30 minutes and electrophoresed on a 15% denaturing polyacrylamide gel.

S11A and S11B demonstrate template dependent incorporation of NTPs onto a substrate with a downstream strand. Although Mpa Pol is capable of incorporating NTPs to a DNA primer in a template-independent manner (S8), these assays indicate that base selection is performed primarily by template dependence, even in the presence of an annealed sequence of downstream DNA.

S11C demonstrates the template dependent incorporation of NTPs onto a substrate with a self-annealing microhomology of 4 base pairs, which creates a single nucleotide gap on either side. Proficiently incorporation of UTP required prior formation of the microhomology, since Mpa Pol can only extend from a 3’-hydroxyl and template-independent incorporation function is limited to a single addition onto a double-stranded blunt DNA termini. Furthermore, preferential incorporation of UTP indicates strong template dependence.

Gap-filling, phosphoesterase and ligation reaction mixtures contained 50mM Tris pH 7.5, 5mM MnCl₂, 250µM NTPs, 30nM 5’-fluorescein labelled 42mer DNA (A) and 48mer DNA (B), and 300nM Mpa Pol, Lig and PE and 400nM Ku protein where indicated. Mpa Pol pre-incubated with the substrate for 30 minutes where indicated before addition of Mpa Lig, PE and Ku. Electrophoresed on a 15% denaturing polyacrylamide gel.
Supporting information:

Ribonucleolytic resection is required for repair of strand displaced NHEJ intermediates, Edward Bartlett, Nigel C. Brissett and Aidan J. Doherty

S1A  
Mpa Ligase alignment

(A) Methanocella paludicola (Mpa) Ligase protein sequence alignment with Ligase D (LigD) ligase domains (LigDom) of Mycobacterium tuberculosis.
(Mtu), *Mycobacterium Smegmatis* (Msm) and *Pseudomonas Aeruginosa* (Pae). B. *Mpa* polymerase (Pol) protein sequence alignment with LigD polymerase domains (PolDom) of Mtu, Msm and Pae. C. *Mpa* phosphoesterase (PE) protein sequence alignment with LigD phosphoesterase domains (PEDom) of Mtu, Msm and Pae. D. *Mpa* Ku protein sequence alignment with Ku of Mtu, Msm and Pae. All alignments were made with TCoffee and edited in JalView.
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Phosphatase reaction mixtures contained 50mM Tris pH7.5, 5mM MnCl₂, 30nM 5'-fluorescein labelled 16mer DNA with 3'-phosphate, and 250, 500 and 1000nM Mpa PE protein. Incubated at 37°C for 1 hour 30 minutes and electrophoresed on a 15% denaturing polyacrylamide gel.
S8. *Mpa* Pol non-templated NTP incorporation
Polymerase non-templated extension reaction mixtures contained 50mM Tris pH7.5, 5mM MnCl$_2$, 62.5µM ATP, CTP, GTP, UTP or 250µM mixed NTPs where indicated, 30nM 5′-fluorescein labelled 36mer DNA, and 300nM *Mpa* Pol protein. Incubated at 37°C for 1 hour and electrophoresed on a 15% denaturing polyacrylamide gel.

S9. *Mtu* PolDom strand displacement
Polymerase strand displacement reaction mixtures contained 50mM Tris pH7.5, 5mM MnCl$_2$, 250µM NTPs, 30nM 5′-fluorescein labelled 16mer DNA, and 300nM *Mpa* Pol or *Mtu* PolDom protein where indicated. Incubated at 37°C for 1 hour and electrophoresed on a 15% denaturing polyacrylamide gel.
**S10.** *Mpa* PE can rescue NHEJ intermediates formed by *Mtu* PolDom and *Mpa* Pol.

Gap-filling, phosphoesterase and ligation reaction mixtures contained 50mM Tris pH7.5, 5mM MnCl₂, 250µM NTPs, 30nM 5'-fluorescein labelled 16mer DNA, and 300nM *Mpa* Pol or *Mtu* PolDom, 300nM *Mpa* Lig and PE protein where indicated. Incubated at 37°C for 1 hour and electrophoresed on a 15% denaturing polyacrylamide gel.
Polymerase gap filling and strand displacement at nicks is template dependent. Polymerase gap filling and strand displacement reaction mixtures contained 50mM Tris pH7.5, 5mM MnCl₂, 62.5µM of the indicated NTP, 30nM 5' fluorescein labelled 16mer DNA (A, B) and 47mer DNA (C), and 300nM Mpa Pol protein. Incubated at 37°C for 1 hour 30 minutes and electrophoresed on a 15% denaturing polyacrylamide gel.

S11A and S11B demonstrate template dependent incorporation of NTPs onto a substrate with a downstream strand. Although Mpa Pol is capable of incorporating NTPs to a DNA primer in a template-independent manner (S8), these assays indicate that base selection is performed primarily by template dependence, even in the presence of an annealed sequence of downstream DNA.

S11C demonstrates the template dependent incorporation of NTPs onto a substrate with a self-annealing microhomology of 4 base pairs, which creates a single nucleotide gap on either side. Proficient incorporation of UTP required prior formation of the microhomology, since Mpa Pol can only extend from a 3'-hydroxyl and template-independent incorporation function is limited to a single addition onto a double-stranded blunt DNA termini. Furthermore, preferential incorporation of UTP indicates strong template dependence.

Gap-filling, phosphoesterase and ligation reaction mixtures contained 50mM Tris pH7.5, 5mM MnCl₂, 250µM NTPs, 30nM 5’-fluorescein labelled 42mer DNA (A) and 48mer DNA (B), and 300nM Mpa Pol, Lig and PE and 400nM Ku protein where indicated. Mpa Pol pre-incubated with the substrate for 30 minutes where indicated before addition of Mpa Lig, PE and Ku. Electrophoresed on a 15% denaturing polyacrylamide gel.