Temperature adaptation of DNA ligases from psychrophilic organisms

Kristel Berg, Ingar Leiros, Adele Williamson*

Current address: Department of Chemistry, The University of Tromsø - The Arctic University of Norway, N-9019 Tromsø, Norway
Correspondence: adele.k.williamson@uit.no

Abstract
DNA ligases operating at low temperatures have potential advantages for use in biotechnological applications. For this reason, we have characterised the temperature-optima and thermal stabilities of three minimal Lig E-type ATP-dependant DNA ligase originating from Gram-negative obligate psychrophilic bacteria. The three ligases, denoted Vib-Lig, Psy-Lig and Par-Lig show a remarkable range of thermal stabilities and optima, with the first bearing all the hallmarks of a genuinely cold-adapted enzyme, while the latter two have activity and stability profiles more typical of mesophilic proteins. A comparative approach based on sequence comparison and homology modelling indicates that the cold-adapted features of Vib-Lig may be ascribed to differences in surface charge rather than increased local or global flexibility: which is consistent with the contemporary emerging paradigm of the physical basis of cold adaptation of enzymes.

Keywords
ATP-dependent DNA ligase; psychrophile; enzyme activity; temperature optima

Introduction
DNA ligases are DNA-joining enzymes essential for survival of all organisms, due to their critical roles in DNA replication and repair. Using ATP or NAD⁺ as a cofactor, DNA ligases catalyze the formation of a phosphodiester bond between the 5’ phosphate of one DNA strand and the hydroxyl group at the 3’ end of the other DNA strand, producing an intact sugar-phosphate backbone. The enzymatic reaction mechanism can be divided into three nucleotidyltransfer steps (Ellenberger and Tomkinson 2008); the first involves the activation of the enzyme through a nucleophilic attack by a lysine residue to the adenosine cofactor ATP or NAD⁺, releasing nicotinamide mononucleotide for NAD-dependent ligases (NDLs) or di-phosphate in the case of ATP-dependent ligases (ADLs). Next, the nucleophilic 5’-phosphate of the DNA attacks the phosphoramidate bond to form an adenylated-DNA intermediate. The final step involves attack of the 3’-nucleophilic hydroxyl group on the new pyrophosphate bond, forming a phosphodiester bond between the 5’ and the 3’ position of the DNA and releasing the AMP. All three chemical steps depend on a divalent cation, which is usually Mg²⁺ or in some cases Mn²⁺.

DNA ligases are divided into two main classes based on the cofactor required in step 1 of the enzymatic reaction. The ADLs use ATP and are found in all phylogenetic kingdoms, with eukaryotes, archaea and many viruses possessing at least one ADL that is essential for DNA replication (by joining Okazaki fragments), and some encode multiple forms with dedicated roles in DNA repair (Ellenberger and Tomkinson 2008). NDLs meanwhile are found almost exclusively in bacteria where they function...
in both replication and repair (Dwivedi et al. 2008; Wilkinson et al. 2001). In the cases where accessory ADLs are identified in bacteria, it is always in addition to the essential NDLs (Pitcher et al. 2007b).

Since the first X-ray crystal structure of an ADL was solved two decades ago from bacteriophage T7 (Subramanya et al. 1996), numerous structural analyses of bacterial, archael and eukaryotic ADLs have followed (Nishida et al. 2006; Pascal et al. 2004) (Kim et al. 2009; Nishida et al. 2006; Pascal et al. 2004; Petrova et al. 2012) (Akey et al. 2006; Kaminski et al. 2018; Pascal et al. 2006; Shi et al. 2018; Williamson et al. 2018; Williamson et al. 2014), and the wide variety of domains and gene arrangements between the different classes of ligases has become evident. Crystallographic studies of bacteriophage T7 (Doherty and Wigley 1999; Subramanya et al. 1996) revealed a common core architecture of two essential catalytic core domains: the adenylation domain (AD) directly involved in catalysis and the site of step 1 enzyme-adenylation, and the smaller oligonucleotide/oligosaccharide binding domain (OB) that is also required for activity (Doherty and Suh 2000; Doherty and Wigley 1999). These core catalytic domains include six conserved motifs (I, III, IIIa, IV, V, and VI) which are involved in one or more steps of the ligation pathway (Shuman 2009). The AD- and OB domains are connected by a flexible linker that allows them to reorient during DNA binding. An additional N-terminal DNA binding domain has been described in the larger ADLs active in DNA replication in Eukarya and Archaea, and additional enzymatic domains with end-repair functions are appended to the large LigD enzymes involved in bacterial non-homologous end joining (Pitcher et al. 2007a). The Lig E group of ADLs, found predominantly in Gammaproteobacteria, have no additional DNA-binding of enzymatic domains, and may serve as a model for the minimal functional unit of the ATP-dependent ligases. The ADL from the marine psychrophile Psychromonas sp. strain SP041 (Psy-Lig) is the smallest DNA ligase that has been structurally studied, being 41 residues shorter than the minimal ChlV-Lig protein (Williamson et al. 2014). Recent structure-function analysis of Psy-Lig and the closely related Ame-Lig demonstrated a novel mode of ligase engagement with its DNA substrate that relies on well-ordered side-chain contacts on the surface of the conserved domains, rather than re-ordering of flexible loop regions to achieve encirclement of the DNA duplex as was previously observed for minimal viral ligases (Nair et al. 2007; Williamson et al. 2018). All Lig E-type ADLs have strong predictions for N-terminal leader sequences proposed to direct them to the periplasm. Proposed biological functions of such secreted ligases include competence and DNA uptake in the periplasm (Magnet and Blanchard 2004), and the demonstrated increase in activity and solubility when this predicted leader was not included in recombinantly-produced Aliivibrio salmonicida (hereafter referred to as Vib-Lig) supports such signal processing (Williamson and Pedersen 2014).

In the present study we have characterised the temperature-optima and thermal stability of Psy-Lig and Vib-Lig, both of which originate from obligate psychrophiles, along with a third homolog from Pseudoalteromonas artica (hereafter Par-Lig), isolated from sandy beach sediment on the Arctic island of Svalbard (Al Khudary et al. 2008). In order to understand structural determinants for low-temperature activity, possible psychrophilic properties of these enzymes were studied. This builds on previous work by Georlette et al. who conducted biophysical analyses and biochemical comparisons of larger, more complex NDLs from species spanning a range of growth temperature optima (Georlette et al. 2003; Georlette et al. 2000).

Living and thriving at low temperatures requires that both enzyme kinetics and protein stability are adapted accordingly. It is now widely accepted that structural differences between cold-active
enzymes and their mesophilic counterparts enable high specific activity at low temperatures, with a lower energy cost (D’Amico et al. 2002; Feller 2003; Struvay and Feller 2012). The physical origin of decreased temperature optima imparted by these structural changes are an active area of contemporary investigation (Åqvist et al. 2017; Arcus et al. 2016; Isaksen et al. 2016; Saavedra et al. 2018; van der Kamp et al. 2018), but it is generally observed that improved catalytic efficiency is accompanied by a reduced thermal stability and weaker substrate affinity, compared to thermophiles and mesophiles at the opposite end of the temperature scale (Struvay and Feller 2012). For this reason we have also carried out in silico comparisons of these Arctic-derived ADLs with mesophilic-derived counterparts from human pathogens.

DNA ligases adapted to low temperatures offer novel potential advantages for use of these enzymes in biotechnological applications. Recently, the thermolability of a cold adapted DNA ligase was used to develop a novel temperature-sensitive vaccine for tularemia (Duplantis et al. 2011), showing great potential in the biomedical science and other applications where bacterial growth control is crucial. Further, the enzymatic activity performed by DNA ligases in DNA replication and repair makes them useful tools in molecular biology and biotechnology applications, such as genetic engineering and next-generation DNA sequencing technologies (Chambers and Patrick 2015; Shuman 2009; Tanabe et al. 2015). Cold-adapted enzymes have a potential advantage over mesophilic homologs by increasing yields of product at low temperatures, while suppressing contaminating nuclease activity. Finally, should the cold-active ligases be highly active, protocols may be carried out with smaller amounts of enzyme, due to better activity rates. In particular, short base-pair overhangs, i.e. ‘sticky ends’ generated by many restriction enzymes will be stabilized due to the low melting temperature of short tracts of base-pairing involved. For these reasons, improving our understanding of temperature adaptation and identification of psychrophilic traits that could be used directly, or reverse-engineered into commercial ligase scaffolds has important biotechnological applications.

Methods

Protein expression and purification
ADLs from Psychromonas spp. strain SP041 (Psy-Lig) and Aliivibrio salmonicida (Vib-Lig) were expressed and purified as described previously (Williamson and Pedersen 2014; Williamson et al. 2014). The gene encoding the Lig E-type ADL from Pseudoalteromonas artica (WP_010555135; Par-Lig), without the leader peptide, was synthesized by Life Technologies as the mature His-tagged, TEV-cleavable form with codon optimization for E. coli and supplied in the donor vector pDONR221. Transfer to the PHMGWA vector was done using Gateway® cloning (Thermo Fisher), and all steps including expression of the MBP-fusion, purification and tag removal were carried out as described for Psy-Lig and Vib-Lig.

Enzyme assays
Gel-based endpoint assays were carried out as described previously using 20 nt + 20 nt oligomers to form 40 nt product (Williamson et al. 2018; Williamson et al. 2014). Details of substrate preparation are given in Table S1. Reactions contained 80 nM substrate, 1 mM ATP, 10 mM MgCl₂, 10 mM DTT, 50 mM NaCl, 50 mM Tris-HCl pH 8.0. Enzymatic activity was detected by conversion of the FAM-labeled 20 nt substrate oligonucleotide into a 40 nt product, resolved by denaturing electrophoresis, detected by fluorescence on a Pharos FX Plus imager (Biorad) and quantified by band intensity using
the extent of ligation activity was calculated from the ratio of these bands and expressed as a percentage. The temperature dependence of ligase activity was investigated by assaying for 15 min at temperatures between 5 °C to 55 °C for nicked substrates and 5 °C to 35 °C for cohesive substrates. Reactions were allowed to equilibrate for 1 min to the assay temperature, and then the assay was started by addition of the enzyme.

**Differential scanning calorimetry**

Differential scanning calorimetry (DSC) experiments were carried out using an N-DSC III differential scanning calorimeter (Calorimetry Sciences Corporation). Purified ligases with concentrations of 1-2 mg ml⁻¹ were extensively dialyzed against 50 mM HEPES pH 8.0, 100 mM NaCl to ensure complete equilibration. The enzymes were filtered through a 0.2 µm syringe filter (Millipore, Billerica, USA) and degassed for approximately 15 min before being loaded into the sample cell. The dialysis buffer was used as reference for baseline subtraction. Data analysis was performed using the program NanoAnalyse 2.4 (TA instruments). For each protein sample scanned the corresponding buffer baseline was subtracted, and the data were normalized to the molar protein concentration calculated from the absorbance at 280 nm after dialysis and filtration. The calorimetric enthalpy was determined directly from the experimental data, and a theoretical two-state model was fitted using the routines provided in the program for determination of the van ’t Hoff enthalpy.

**Thermofluor**

Thermal denaturation of the purified ADLs with different buffers were examined by the thermofluor assay as described previously (Ericsson et al. 2006). Briefly, 5 µl of protein (1.0-1.5 mg ml⁻¹) was mixed with 1 µl of 300 x Sypro-Orange, 12.5 µl of 50 mM HEPES pH 8.0, 200 mM NaCl, added to the wells of a 96-well PCR plate (Bio-Rad) and sealed with Microseal® 'B' Adhesive Seals from Bio-Rad. Melting curves were recorded from 20 °C to 90 °C in increments of 0.3°C per sec using a MiniOpticon Real-Time PCR System with both FAM and HEX dye channels selected. Tₘ was determined using the supplied instrument software and monitoring the fluorescence of the HEX channel.

**Sequence comparison**

The amino acid sequences of Par-Lig, Psy-Lig and Vib-Lig, were aligned with the Lig E sequence from *Vibrio cholera* (Vch-Lig; gi|147674166). N-terminal leader sequences were predicted using SignalP 4.1 and omitted from further analyses (http://www.cbs.dtu.dk/services/SignalP/) (Petersen et al. 2011). The ClustalW alignment tool in BioEdit was used to determine sequence identities and similarities. Conserved domains were analysed by Pfam protein families database at EMBL-EBI (http://pfam.xfam.org).

**Homology modelling and analysis**

Homology models of Vib-Lig, Par-Lig and Vch-Lig were built based on the deposited crystal structure of Psy-Lig (4D05; (Williamson et al. 2014)). The sequences were uploaded to the Swiss-Model homology modeling server (Biasini et al. 2014). The A-chain of the deposited structure of Psy-Lig was selected as a modeling template for all modeled structure as it has overall superior quality than the B-chain (with lower overall B-factor and amino acid residues generally better defined in electron density).

HBPLUS Hydrogen Bond Calculator v 3.2 (McDonald and Thornton 1994) was used to calculate hydrogen bonds in all PDB-files. The hydrogen bonds included were those fulfilling the criteria for
parameters donor (D), acceptor (A), acceptor antecedents (AA) and calculated hydrogen (H): 
maximum distance for D-A, 3.5 Å and H-A, 2.5 Å; minimum angle for D-H-A, D-A-AA and H-A-AA of 90°. Ion-pair-interactions were investigated using the WHAT IF Web Interface (http://swift.cmbi.ru.nl/servers/html/index.html) (Vriend 1990), where interatomic distances between the side-chains of the negatively charged Asp and Glu, and the positively charges Arg, Lys and His were tabulated with respect to being <4 Å and <6 Å. The APBS plugin in Pymol was used to estimate electrostatic surface potentials (Dolinsky et al. 2007).

Results

Temperature optimum and thermal stability

The aim of this study was to understand the determinants of low-temperature adaptation among DNA ligases. We chose to investigate the temperature optimum and thermal stability of Lig E ADLs from Psychromonas spp. strain SP041, Aliivibrio salmonicida, and Pseudoalteromonas artica, delineated Psy-Lig, Vib-Lig and Par-Lig respectively as these represent psychrophilic species of bacteria isolated from a consistently low-temperature environments (Al Khudary et al. 2008; Egidius et al. 1986) (Error! Reference source not found.).

To analyze the temperature optima for ligase activity, gel-based endpoint assays were performed, both with single nicked and overhanging substrates. Nick sealing activity was measured by temperature intervals of 5°C, ranging from 5 to 60 °C, or until the activity was abolished. For ligation of single-nicked substrates (Figure 1a), there is a sharp peak of more than 50% ligation activity at around 20°C for Vib-Lig, quickly declining to 10% activity at 30°C, whereas the activity of Psy-Lig and Par-Lig increases with temperature from 15 °C up to an optimum of 35-40°C, above which a sharp decline is observed. Although all ligases were cloned from psychrophilic organisms with similar growth temperatures, the T_{opt} of their ligases for nicked substrates are different (Table 1).

The characterized ligases Psy-Lig, Par-Lig and Vib-lig show a similar and relatively broad temperature optimum on the overhang substrate tested, with approximately 60-80 % ligation activity from 5°C to 25-30°C, followed by a sharp decline at higher temperatures. As they all show better activity on overhang breaks at lower temperatures, we suggest that substrate stability rather than enzyme activity is the driving feature here. However, the enzymatic reaction will work very slowly at the low temperature, requiring a longer incubation time.

DSC experiments were performed to obtain a complete thermodynamic profile of the protein unfolding process of Psy-Lig, Par-Lig and Vib-Lig. The melting temperature (T_m) was estimated to be significantly lower for Vib-Lig, 30.7 °C, compared to Psy-Lig and Par-Lig with a T_m of 46.0 °C and 53.7 °C, respectively (Figure 2a). All three ligases measured show a ratio > 1 between the van’t Hoff enthalpy derived from fitting a two-state model, and the calorimetric enthalpy, derived by integration of the area under the excess heat capacity. Such temperature profile indicates that unfolding proceeds as a higher order oligomer; however, the irreversibility of the unfolding transition precluded detailed thermodynamic analysis.

The thermal stability in various buffer systems was measured by a thermofluor assay to confirm the DSC results and exclude the possibility that low thermal stability of observed for Vib-Lig is caused by non-ideal buffer conditions as it has a significantly lower pI (predicted to be 5.5) relative to Psy-Lig and Par-Lig (both greater than 9.0). Thermofluor data (Figure 2b) suggest that stability of the various
ligases does not vary between pHs 6.5 and 9, with the exception of Psy-Lig which is extremely
unstable in phosphate buffer at pH 7.0. Otherwise, Psy-Lig shows stability up to 46 °C and Par-Lig up
to 53 °C, which is in line with DSC unfolding temperature. Also consistent with DSC data, Vib-Lig
shows a lower thermal stability relative to Psy-Lig and Par-Lig with a maximum at 23 °C in all buffers
down to pH 6.5. Below this, no transition could be observed, indicating that Vib-Lig was already
unfolded.

**Sequence comparison**

Cold-active enzymes may combine rigidity and stability with a high level of flexibility. To gain further
insight into the activity/stability/flexibility relationship and cold adaptation, interesting sequence and
structural differences were identified by sequence alignments and homology modelling.

The enzymes studied are of similar size and share all properties common to minimal ADLs but exhibit
different temperature optima and stabilities. A structure-based sequence alignment was generated
(Figure 3). Lig E from *V. cholera* (Vch-Lig) was included as this human pathogen has growth
temperature between 20 and 45 °C and is unable to survive at 4 °C for extended periods of time
(Martinez et al. 2010). Pairwise comparison of the three experimentally-examined Lig Es together
with Vch-Lig show that all sequence pairs have identities in the 40-49% range. Consistent with both
enzymes deriving from members of the genus *Vibrio*, Vib-Lig and Vch-Lig share the highest homology
in terms of sequence identity (48.4 %), although they are adapted to different habitats and
temperatures; thus Vch-Lig represents a phylogenetically-related mesophilic homolog of Vib-Lig. All
four Lig Es contain the conserved nucleotidyltransferase family motifs I-VI and align with very few
insertions or deletions, giving high confidence in placement of secondary structural elements by
homology modeling (described below). Further, the sequence alignment revealed high conservation
of amino acids involved in substrate binding, metal binding and enzymatic activity.

Several studies have indicated increased occurrence of some residues in cold-adapted proteins and
decreased frequency of others, which has been rationalized by physical properties of their sidechains
influencing flexibility and stability of the protein. This includes fewer salt bridges, fewer hydrogen
bonds, a lower content of proline residues, a reduced Arg/(Arg + Lys) ratio, lower (Leu + Ile)/(Leu + Ile
+ Val) ratio and increased glycine content (Aghajari et al. 1998; Collins et al. 2005; Huston et al. 2004;
Metpally and Reddy 2009; Russell et al. 1998; Saavedra et al. 2018). For this reason, we compared
the amino acid content of the four proteins; however most classic sequence ‘traits’ of cold-
adaptation, including increased glycine, decreased proline and less-packed hydrophobic core, were
not apparent in Vib-Lig. Instead, higher sequence conservation appeared to be with the more
phylogenetically-related Vch-Lig than the other psychophile-derived ADLs. For example, a lower
number of Gly residues is often pinpointed as a typical cold adapted trait, however this did not
correlate with thermal stability of these ADLs, and most Gly residues are conserved, especially
between the psychrophilic Vib-Lig and the mesophilic Vch-Lig (Table 2). Likewise, decreased Pro
content has also been related to cold-adaptation (Wallon et al. 1997; Zhao et al. 2010), but as Vch-
Lig has fewer Pro than Vib-Lig (11 versus 13) Pro content is not an evident factor.

Another ‘typical’ feature of cold-adapted enzymes is a decreased number of Arg residues, which may
increase stability through its capability to form hydrogen bonds and salt bridges (Ait Talebeh et al. 1997).
In line with this, we observed the highest Arg count in the presumably mesophilic Vch-Lig (Table 2).
The number of Arg is significantly lower for Vib-Lig (11), Psy-Lig (12) and Par-Lig (11) compared to
Vch-Lig (18). This is also reflected by the ratio Arg/(Lys+Arg) per residue, which is 0.53 in Vch-Lig compared to 0.40, 0.39 and 0.39 in Psy-Lig, Vib-Lig and Par-Lig, respectively, also supporting an overall better stability of the mesophilic molecule. Arg can contribute in more interactions with surrounding amino acids than lysine. However, Arg may also interact with water on the surface. Interestingly, the multiple sequence alignment (Figure 3) shows that Arg in Vch-Lig are frequently substituted with hydrophobic residues in Vib-Lig.

Homology modeling and comparison to the crystal structure of Psy-Lig

To identify positions in the three-dimensional ligase structure where relevant amino acid substitutions occurred, homology models of Par-Lig, Vib-Lig and Vch-Lig were built based on the deposited structure of Psy-Lig 4D05; (Williamson et al. 2014). Increased local and/or global flexibility can be achieved by destabilization of the structure through a reduction in intramolecular forces such as salt-bridges, ion-pair networks, hydrogen bonds and aromatic interaction, and increased length of loop regions (Davail et al. 1994; Feller 2003; Russell 2000). Hydrogen bond analysis shows that Vib-Lig is possibly destabilized by fewer hydrogen bonds per residue in total, compared to Par-Lig and Psy-Lig (0.715/0.778/0.759). In comparison, the mesophilic Vch-Lig has the highest ratio of hydrogen bonds per residue (0.816). It is interesting to note that the ratios correlate well with the measured melting temperatures Vib-Lig, Psy-Lig and Par-Lig with low ratios giving low melting temperatures. In particular, the number of side-chain to main-chain hydrogen bonds is lower for the cold-adapted Vib-Lig.

Examination of the structural models also revealed that the substituted arginines described in the preceding section are generally located on the surface, thus introducing hydrophobic surface patches in Vib-Lig (Figure 4). Calculations by POPS (Parameter OPtimsed Surfaces (Fraternali and Cavallo 2002)) showed that the overall total area of exposed hydrophobic residues were similar among all ligases, thus, unique exposed hydrophobic patches in Vib-Lig appear to be local. Interesting Arg substitutions in Vib-Lig compared to Vch-Lig include Arg95 to Ala90, Arg167 to Ile162, Arg193 to Thr188, Arg209 to Gln204 and Arg257 to Ala252 (Figure 4). For Par-Lig and Psy-Lig, three of these Arg are substituted with Leu/Lys. The percentage of hydrophobic residues is slightly higher for Vib-Lig (42.80%) and Psy-Lig (41.25%) compared to Par-Lig (39.25%) and Vch-Lig (39.69%), possibly reflecting the substitutions of polar residues with hydrophobic residues on the surface compared to Vch-Lig. In combination, the elevated number of hydrophobic residues described above, the unique local hydrophobic surface patches and the lower number of Arg, may impart local flexibility to the Vib-Lig structure compared to its mesophilic counterpart Vch-Lig.

Electrostatic surface potential

Some cold adapted enzymes feature an overall excess of negative charges at the surface of the protein, with a pl frequently more acidic than that of their mesophilic homologues(Feller 2003; Leiros et al. 1999; Russell 2000). Higher frequency or patches of acidic residues on the surface may increase solvent interactions and thereby lead to an overall destabilization of the enzyme by charge-charge repulsion, observed in cold-adapted trypsin and β-lactamase (Feller 2003; Leiros et al. 1999). The calculated pl of 5.3 for Vib-Lig is significantly more acidic compared to its counterparts, and also correlates with the substitution of basic arginine residues at the surface with hydrophobic amino acids. Further examination of the charge distribution on the surface of the Vib-Lig model (Figure 5) indicates that the DNA-binding faces of Vib-Lig remain positively charged as seen for structures of other Lig Es, while surfaces not involved in DNA binding are more positively charged compared with
the more thermostable Psy-Lig. This suggests that charges in the binding surfaces of Vib-Lig are conserved and the majority of variation located in distant areas of the protein.

**Conservation of Active site and DNA-binding surface**

It is often suggested that low-temperature adaptation of enzymes is driven by increased local flexibility at the active site (D’Amico et al. 2002; D’Amico et al. 2006; Struvay and Feller 2012), therefore we examined three key areas of the Vib-Lig enzyme that are essential for activity: the region surrounding the AMP-binding pocket where the enzyme is covalently adenylated in the first step of the ligase reaction, the inter-domain linker region which undergoes significant structural changes during the catalytic cycle and the surfaces of the adenylation (AD-) and oligonucleotide-binding (OB-) domains that are in contact with double-strand DNA during nick-sealing. Our comparisons reveal that the active site is strictly conserved, except for Lys 41 in Psy-Lig which is replaced by the chemically-similar Arg in the other three ADLs (Fig 3, supplementary figure S1a). The sequence alignment shows that the flexible linker regions connecting the two core domains are similar, preserving the hydrogen bonding pattern observed in Psy-Lig, with the exception of Par-Lig where the equivalent of Lys 176 (Psy-Lig) is replaced by Pro (Fig 3, supplementary figure S1b). Lig E-type ligases efficiently ligate DNA breaks without any additional DNA-binding domains or large flexible loop regions, instead using interactions with shorter highly structured motifs and specific charged residues found on the DNA-binding surface of the core catalytic domains (Williamson et al. 2018; Williamson et al. 2014). In general these motifs are well conserved between the three variants, consistent with both the equivalent positively-charged DNA binding surfaces of Vib-Lig and Psy-Lig and previous observations of consensus between Lig Es in this region (Fig 5) (Williamson et al. 2018).

**Discussion**

In this study, biochemical and biophysical characteristics of ATP-Dependent Ligases (ADLs) from psychrophilic organisms were analyzed in an attempt to identify typical cold-adaptation features. Vib-Lig, originating from the psychrophilic fish pathogen Aliivibrio salmonicida which has a growth range of 1-22 °C and an optimum of 15 °C (Egidius et al. 1986) exhibits classical features of cold adaptation including a low temperature optimum of activity compared to homologous enzymes and decreased thermostability. In contrast, despite being derived from psychrophilic organisms, the Psy-Lig and Par-Lig enzymes are not themselves cold-adapted as both have temperature optima in the range of 35-40 °C and unfolding temperatures greater than 45 °C. This has been observed for many psychophile-derived enzymes such as L-haloacid dehalogenase from Psychromonas ingrahamii, alcohol dehydrogenase of Flavobacterium frigidimarvis and KUC-12-keto acid decarboxylases derived from Psychrobacter, where the individual enzymes remain active and stable at temperatures well above the survival limit of the host organism (Kazuoka et al. 2007; Novak et al. 2013; Wei et al. 2013). The simplest rationale in the case of the ADLs is that although the temperature optimum is relatively high, the 30–40% activity recorded below 15 °C is sufficient for the biological purposes of the bacterium in its native environment, although the effects of different conditions on in vivo activity are also possible.

During ligation of double-strand breaks with cohesive ends, low temperature is an advantage to stabilize base-pairing between short stretches of complementary nucleotides at the break site. This must be balanced against decreased enzyme activity at lower temperatures. Lower temperatures allow DNA overhangs to base-pair and remain annealed long enough for the ligase to join them, at the
expense of reduced ligase activity. This is directly observed in the present study during ligation of substrates with 4nt overhangs as optimal activities are shifted to lower temperatures for all three enzymes measured, despite of their individual $T_{\text{opt}}$ varying when measured with a nicked substrate.

The Lig E enzymes compared in our study have moderate sequence identities (40-50%) and likely highly similar structures. The analyses performed indicated some sequence differences that potentially lower $T_{\text{opt}}$ of Vib-Lig relative to homologs. One difficulty in such comparative analyses is distinguishing between substitutions imparting psychrophilicity and those that have occurred through genetic drift. To exclude possible false-positive findings based on phylogenetic resemblance, we included Lig E from *V. cholera* in our sequence comparison as previous phylogenomic studies have placed this close to Vib-Lig in evolutionary terms (Williamson et al. 2016), thus representing a genus-related but mesophilic organism. Coming from a mesophilic human pathogen, Vch-Lig is not anticipated to exhibit cold-adapted characteristics. The major differences in Vib-Lig appear to be in non-DNA-binding surface exposed residues. Arginines are generally located on the surface of Psy-Lig, Par-Lig and Vch-Lig, and substitution of these positions introduces hydrophobic or uncharged surface patches in Vib-Lig. This is consistent with the investigation of three structurally homologous NAD$^+$-dependent DNA ligases (NDLs) adapted to different temperatures, where specific surface areas revealed a significant increase of exposed hydrophobic residues to solvent, in contrast to a more hydrophilic and charged surface area in thermophiles (Georlette et al. 2003), indicating an entropy-driven destabilization of the protein structure. Likewise, replacements of lysine with arginine in the psychrophilic $\alpha$-amylase from *Pseudoalteromonas haloplanktis* resulted in a more stabilized enzyme with mesophilic properties, demonstrating the relevance of arginine content in cold adaptation (Siddiqui et al. 2006). It is interesting to note that these substitutions in Vib-Lig are unevenly distributed between the two domains with only two occurring in the larger catalytic adenylation domain (approximately 170 residues), and four on the smaller oligonucleotide domain (approximately 80 residues). Recent work demonstrated that substitutions increasing flexibility in different domains of adenylylate kinase gave rise to different temperature effects on substrate binding and catalysis (Saavedra et al. 2018). As with DNA ligases, adenylylate kinase activity involves coordinated reorientations between discrete protein domains, and it is interesting to consider whether this distribution reflects tuning of the oligonucleotide binding domain for DNA binding/product release rather than catalysis, as the former is are the rate limiting processes (Bauer et al. 2017; Lohman et al. 2011).

Calculations of the electrostatic surface potential revealed that the cold-active Vib-Lig displays a positively charged surface near the active site and on the binding face of the OB-domain, which is important for binding of the negatively charged DNA substrate (Fig 5), despite its overall more acidic pl. Similar results were observed for the cold-adapted uracil-DNA N-glycosylase (cUNG) from Atlantic cod (Leiros et al. 2003), indicating increased affinity for the negatively charged DNA compared with mesophilic homologues. The number and nature of residues around the active site are conserved among the homologous ADLs adapted to different temperatures, suggesting that local cold adapted residues are not directly involved in catalysis, but influence flexibility indirectly at some distance apart. The psychrophilic Vib-Lig is further characterized by a decreased number of hydrogen bonds, which correlates with an increase in overall flexibility of the enzyme and affects protein water surface interactions.
Although the decreased temperature optima of psychrophile-derived enzymes is commonly attributed to an increase in flexibility, either global or local, which causes a concomitant lowering of thermal stability, (Smalas et al. 2000) (D'Amico et al. 2002; Feller 2003; Struvay and Feller 2012) many enzymes are inactivated by temperatures below those inducing denaturation. A comparison of the NDL from the psychrophile *Pseudoalteromonas haloplanktis* with that of mesophilic NDL of *E. coli* and the thermophilic NDL of *Thermus scotoductus* found that structural differences imparted a temperature optimum of 18 °C, compared to 30 °C for *E. coli* and more than 60 °C for *T. scotoductus* (Georlette et al. 2000). This is accompanied by a decrease in $T_m$ in the *P. haloplanktis* enzyme (33 °C) compared to the ones from *E. coli* (54 °C) and *T. scotoductus* (95-101 °C) (Georlette et al. 2003). The temperature optimum for activity of the *E. coli* NDL corresponds to the beginning of the thermal unfolding. *P. haloplanktis* NDL, however, shows a different link between activity and thermal adaptation; optimal activity is reached 10 °C before unfolding and the enzyme is inactivated at the beginning of the unfolding transition. A similar behaviour is observed for the activity and stability of Vib-Lig, Psy-Lig and Par-Lig, where a decrease in activity above $T_{opt}$ is observed in the absence of denaturation/unfolding. Recently, new paradigms have been suggested to explain this behaviour, as the classical (two-state) model is limited to enzymes where increased catalytic activity is directly followed by thermal inactivation. These include macromolecular rate theory (MMRT), which provides a rationale for the curved temperature-rate plots observed for enzymes, independent of denaturation, and describes the temperature dependence of enzyme-catalyzed rates in the absence of denaturation by the difference in heat capacity between the enzyme substrate complex and the enzyme transition state species (Arcus et al. 2016). The three-state equilibration model (EM) (Daniel and Danson 2013) has also been suggested to explain the temperature dependence of enzyme-catalyzed rates in the absence of denaturation. EM introduces a reversible inactivated (not denatured) form of the enzyme ($E_{inact}$) as an intermediate in rapid equilibrium with the active form ($E_{act}$), which adds a thermal buffer effect that protects the enzyme from thermal inactivation.

Another explanation invokes a tuning of surface mobility through alteration of regions spatially removed from the active site which affect the overall enzyme dynamics (Åqvist et al. 2017; Isaksen et al. 2016). Computer simulations and Arrhenius plots suggest that surface rigidity/flexibility outside the catalytic region affects the enthalpy/entropy balance. Key single distant mutations may disrupt surface hydrogen binding networks and alter the protein water surface interactions (Isaksen et al. 2016) which may be the case with arginine substitutions in our study.

### Conclusions

We have described the temperature optima and thermal denaturation profiles of three psychrophile-derived ADLs of the minimal Lig E-type. In the course of this work we determined that two of the three, Par-Lig and the structurally-characterized Psy-Lig did not exhibit marked psychrophilic properties, while the third had typical low-temperature characteristics such as low $T_{opt}$ and low thermal stability. Sequence comparison and homology modeling identified surface-exposed patches with greater hydrophobicity in Vib-Lig, relative to homologs, which we suggest are relevant for the experimentally-observed psychrophilic properties.

Consistent with our observations for Vib-Lig and Vch-Lig, catalytic sites may be strictly conserved between homologs with different activity optima, meaning explanations for the markedly lower $T_{opt}$ of Vib-Lig relative to Psy-Lig and Par-Lig cannot invoke specific increases in active-site flexibility. We hope
that future application of more sophisticated computational methods, coupled with specific mutational studies may elucidate general principles imparting low-temperature activities which can be transferred to commercially-relevant DNA ligases allowing us to tailor their activity optima in biotechnological applications.

Figures

Figure 1. Temperature optimum of Psy-Lig, Par-Lig and Vib-lig by ligase activity assay. (a) Percentage of ligated single-nicked substrate (b) Percentage of ligated cohesive substrate. Ligase activity was quenched after 15 minutes at various temperatures and quantified as percentage ligation by the intensity of the upper band relative to the sum of the two bands on the TBE-UREA gel. Ligase concentration was 2.5 µM for the nicked substrate, and 100 µM for the cohesive substrate.

Figure 2. Biophysical data. (a) Thermal unfolding monitored by DSC (b) Thermal stability measured by thermofluor.

Figure 3. Amino acid sequence alignment comparing mature ATP-dependent ligases from Psychromonas spp. strain SP041 (Psy-Lig), Aliivibrio salmonicida (Vib-Lig), Pseudoalteromonas artica (Par-Lig) and Vibrio cholera (Vch-Lig). Identical residues are shaded with red and similar residues are shown in red text. Spirals indicate α-helices and arrows indicate β-strands. Boxed amino acids represent conserved motifs of the nucleotidyltransferase enzymes. The DNA-binding elements of Lig Es are boxed with dashed lines. Surface-exposed substitutions of basic to uncharged residues in Vib-Lig are indicated by blue circles.

Figure 4. Sequence variability mapped onto molecular surface representations of Vib-Lig, Psy-Lig, Par-Lig and Vch-Lig. The top and bottom panels are rotated 180° views, while the middle panel shows melting temperature and substituted amino acids in selected positions for the four enzymes. Color codes: Blue: positively charged residues; Green: polar residues; Orange: hydrophobic residues. Vib-Lig Residue numbers are included for reference between the panels. There is an apparent correlation between reduced thermostability and substitution from charged residues into more hydrophobic ones.

Figure 5. Structure of Psy-Lig (top) and model of Vib-Lig (lower) colored surface charge. The surface potential was generated using APBS (Dolinsky et al. 2007), with positively charged areas shown in blue and negatively charged areas in red.

Tables

Table 1. Literature and experimental data showing host optimal growth temperature, ligase temperature optimum for nick sealing and melting temperature.

<table>
<thead>
<tr>
<th>Ligase</th>
<th>Species of origin</th>
<th>Optimal growth (°C)</th>
<th>T_max (°C)</th>
<th>T_opt (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Psy-Lig</td>
<td>Psychromonas spp. strain SP041</td>
<td>15&lt;sup&gt;b&lt;/sup&gt;</td>
<td>35&lt;sup&gt;a&lt;/sup&gt;</td>
<td>46&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Par-Lig</td>
<td>Pseudoalteromonas artica</td>
<td>10-15&lt;sup&gt;c&lt;/sup&gt;</td>
<td>35-40&lt;sup&gt;a&lt;/sup&gt;</td>
<td>53&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Vib-Lig</td>
<td>Aliivibrio salmonicida</td>
<td>15&lt;sup&gt;d&lt;/sup&gt;</td>
<td>20&lt;sup&gt;a&lt;/sup&gt;</td>
<td>30&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup>This study
<sup>d</sup>Egidius et al, (1986) (Egidius et al. 1986)
Table 2. Brief summary of extracted sequence features and characterization data for Psy-Lig, Vib-Lig, Par-Lig and Vch-Lig, respectively.

<table>
<thead>
<tr>
<th></th>
<th>Psy-Lig</th>
<th>Vib-Lig</th>
<th>Par-Lig</th>
<th>Vch-Lig</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sequence length</td>
<td>257</td>
<td>257</td>
<td>260</td>
<td>262</td>
</tr>
<tr>
<td>$T_{\text{opt}}$ (°C)</td>
<td>35</td>
<td>20</td>
<td>35-40</td>
<td>-</td>
</tr>
<tr>
<td>$T_{\text{melt}}$ (°C)</td>
<td>46.0</td>
<td>30.7</td>
<td>53.7</td>
<td>-</td>
</tr>
<tr>
<td>Calculated pI</td>
<td>9.1</td>
<td>5.3</td>
<td>9.5</td>
<td>9.0</td>
</tr>
<tr>
<td>Net charge</td>
<td>+4</td>
<td>-9</td>
<td>+9</td>
<td>+4</td>
</tr>
<tr>
<td>Polar residues (%)</td>
<td>35.8</td>
<td>29.2</td>
<td>37.3</td>
<td>34.0</td>
</tr>
<tr>
<td>Hydrophobic residues (%)</td>
<td>41.3</td>
<td>42.8</td>
<td>39.2</td>
<td>39.7</td>
</tr>
<tr>
<td>Aromatic residues (%)</td>
<td>10.9</td>
<td>9.7</td>
<td>11.2</td>
<td>11.5</td>
</tr>
<tr>
<td>Gly (number and %)</td>
<td>17/6.6</td>
<td>24/9.3</td>
<td>20/7.7</td>
<td>23/8.8</td>
</tr>
<tr>
<td>Met (number and %)</td>
<td>7/2.7</td>
<td>8/3.1</td>
<td>3/1.2</td>
<td>6/2.3</td>
</tr>
<tr>
<td>Pro (number and %)</td>
<td>13/5.1</td>
<td>13/5.1</td>
<td>12/4.6</td>
<td>11/4.2</td>
</tr>
<tr>
<td>Arg (number and %)</td>
<td>12/4.7</td>
<td>11/4.3</td>
<td>12/4.6</td>
<td>18/6.9</td>
</tr>
<tr>
<td>Arg/(Lys+Arg)</td>
<td>0.40</td>
<td>0.39</td>
<td>0.39</td>
<td>0.53</td>
</tr>
<tr>
<td>$(\text{Leu+Ile})/(\text{Leu+Ile+Val})$</td>
<td>0.76</td>
<td>0.70</td>
<td>0.65</td>
<td>0.78</td>
</tr>
</tbody>
</table>

- a Residues R, K, D and E
- b Residues G, S, T, Y, N, Q and C
- c Residues A, V, L, I, W, F, P and M
- d F, W and Y

Table 3. Summary of calculated intramolecular interactions for Psy-Lig, Vib-Lig, Par-Lig and Vch-Lig, respectively.

<table>
<thead>
<tr>
<th></th>
<th>Psy-Lig</th>
<th>Vib-Lig</th>
<th>Par-Lig</th>
<th>Vch-Lig</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDB ID</td>
<td>4d05</td>
<td>Model</td>
<td>Model</td>
<td>Model</td>
</tr>
<tr>
<td>Resolution</td>
<td>1.65 Å</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>No. of residues in PDB file</td>
<td>257</td>
<td>256</td>
<td>257</td>
<td>250</td>
</tr>
<tr>
<td>No. of hydrogen bonds per residue</td>
<td>0.759</td>
<td>0.715</td>
<td>0.778</td>
<td>0.816</td>
</tr>
<tr>
<td>No. SS$^*$ hydrogen bonds per residue</td>
<td>0.086</td>
<td>0.066</td>
<td>0.066</td>
<td>0.104</td>
</tr>
<tr>
<td>No. SM$^*$ hydrogen bonds per residue</td>
<td>0.202</td>
<td>0.133</td>
<td>0.175</td>
<td>0.180</td>
</tr>
<tr>
<td>No. MM$^*$ hydrogen bonds per residue</td>
<td>0.471</td>
<td>0.516</td>
<td>0.537</td>
<td>0.532</td>
</tr>
<tr>
<td>No. ion pairs $&lt;$4/$&lt;$6 Å</td>
<td>8/19</td>
<td>11/18</td>
<td>10/16</td>
<td>12/23</td>
</tr>
<tr>
<td>No. 2 membered networks $&lt;$4.0 Å</td>
<td>6</td>
<td>5</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>No. 3 membered networks $&lt;$4.0 Å</td>
<td>1</td>
<td>3</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

- $^*$ SS, side-chain to side-chain hydrogen bonds.
- $^*$ SM, side-chain to main-chain hydrogen bonds.
- $^*$ MM, main-chain to main-chain hydrogen bonds.

Acknowledgements and Funding

This research was supported by Research Council Norway [244247, 2015]; Funding for open access charge was granted by the publication fund at the University of Tromsø.


References
14


Siddiqui KS et al. (2006) Role of lysine versus arginine in enzyme cold-adaptation: Modifying lysine to homo-arginine stabilizes the cold-adapted alpha-amylose from Pseudoalteremonas haloplanktis Proteins 64:486-501

Smalas AO, Leiros HK, Os V, Willassen NP (2000) Cold adapted enzymes Biotechnology annual review 6:1-57


Tanabe M, Ishino Y, Nishida H (2015) From Structure-Function Analyses to Protein Engineering for Practical Applications of DNA Ligase Archaea (Vancouver, BC) 2015:267570
doi:10.1155/2015/267570

doi:10.1038/s41467-018-03597-y

Vriend G (1990) WHAT IF: a molecular modeling and drug design program Journal of molecular graphics 8:52-56, 29

Wallon G et al. (1997) Sequence and homology model of 3-isopropylmalate dehydrogenase from the psychrotrophic bacterium Vibrio sp. IS suggest reasons for thermal instability Protein Eng 10:665-672

doi:10.1111/1574-6968.12208


doi:10.1093/nar/gky622

doi:10.1111/mmi.13229

doi:http://dx.doi.org/10.1016/j.pep.2014.02.008

doi:10.1107/S1399004714021099

doi:10.1371/journal.pone.0009109