**Extreme Catalysts from Low-Temperature Environments**

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Cold-loving or psychrophilic organisms are widely distributed in nature as a large part of the earth’s surface is at temperatures around 0°C. To maintain metabolic rates and to prosper in cold environments, these extremophilic organisms have developed a vast array of adaptations. One main adaptive strategy developed in order to cope with the reduction of chemical reaction rates induced by low temperatures is the synthesis of cold-adapted or psychrophilic enzymes. These enzymes are characterized by a high catalytic activity at low temperatures associated with a low thermal stability. A study of protein adaptation strategies suggests that the high activity of psychrophilic enzymes could be achieved by the destabilization of the active site, allowing the catalytic center to be more flexible at low temperatures, whereas other protein regions may be destabilized or as rigid as their mesophilic counterparts. Due to these particular properties, psychrophilic enzymes offer a high potential not only for fundamental research but also for biotechnological applications.

**Key words:** psychrophile, extremophiles, cold adaptation, enzyme kinetics, flexibility strategy

Life under low-temperature conditions was identified as early as 1840 by Hooker, who observed that algae were associated with sea ice. In 1887, Forster was the first who reported that microorganisms isolated from fish could grow well at 0°C (1). The term “psychrophilic” was first used in 1902 by Schmidt-Nielsen to describe such cold-adapted organisms (2). Psychrophile is defined as an organism, prokaryotic or eukaryotic, living permanently at temperatures close to the freezing point of water in thermal equilibrium with the medium. Thus psychrophiles are numerous, including a large range of species of gram-positive and gram-negative bacteria, yeast, algae, marine invertebrates, insects and polar fish, and are widely distributed (3). Psychrophiles have developed mechanisms of adaptation to temperature including a huge range of structural and physiological adjustments in order to cope with the deleterious effect of low temperatures. Indeed, they display metabolic fluxes at low temperatures that are more or less comparable to those exhibited by closely related mesophiles living at moderate temperatures (4–6). This is explained by the capability of these psychrophilic organisms to produce “cold-adapted” enzymes which are able to cope with the reduction of chemical reaction rates induced by low temperatures. However, most cellular adaptations to low temperatures and the underlying molecular mechanisms are not fully understood and are still being investigated. Moreover, a study of proteins and enzymes from cold-adapted organisms is not only useful in the understanding of some general processes related to the protein structure and function but also in protein folding investigations. In addition, cold-active and heat-labile psychrophilic enzymes possess an interesting biotechnological potential.

**I. PSYCHROPHILIC ORGANISMS**

Cold-adapted organisms prosper at temperatures close to the freezing point of water and have successfully colonized permanently cold habitats such as polar and alpine regions or deep-sea waters (7–9). These cold habitats constitute more than three-quarters of the earth’s surface and are exposed to temperatures that are more or less permanently below 5°C. Microbial growth and metabolic activities have been recorded beneath 3–6 m of ice in permanently frozen lakes, in subglacial ice and sediments, in surface snow at the South Pole where the highest summer temperatures remain well below zero (at least –45°C). Moreover, biological activities have been recorded in the brine veins of sea ice at temperatures as low as –20°C (10). More often, psychrophilic microorganisms are not only adapted to low temperatures but also to other environmental constraints. In deep-sea water, for example, they have to be adapted to extremely high pressure and therefore must be, at the same time, psy-
Psychrophiles and piezophiles (11). Temperature is one of the most important environmental factors for life, as it influences most biochemical reactions. In order to overcome the detrimental effect of low temperatures, psychrophiles have developed various adaptive strategies. One of the strategies used in order to withstand temperatures below freezing is the protection of the cells from ice formation by the synthesis of antifreeze molecules and cryoprotectors (12). Another problem is the decrease in membrane fluidity with temperature. Indeed, major cellular functions, including passive and active permeabilities, nutrient uptake, electron transport, environmental sensing, photosynthesis, and recognition processes, require the maintenance of membrane fluidity. The membrane of microorganisms contains a lipid bilayer that must have the proper fluidity to support most biological functions. As temperature is decreased, fatty acid side chains in membrane bilayers undergo a change of state from a disordered liquid crystalline state to a gel-like state (13). To maintain the membrane stability, psychrophilic microorganisms increase the ratio of unsaturated to saturated fatty acids and modulate the activities of the enzymes involved in fatty acid and lipid biosyntheses (3).

Sudden decreases in temperature will elicit a specific alteration in gene expression known as the cold-shock response, which is distinct from the general stress response (heat shock) in terms of both the nature of the proteins induced and regulatory aspects (3). The cold-shock response involves the induction and the synthesis of cold-shock proteins. The regulation of cold-shock protein synthesis is multifactorial, being controlled at the levels of both transcription and translation, as well as mRNA and protein stabilities. The main functions of cold-shock proteins (Csp) are in the regulation of cellular protein synthesis, at the level of transcription as well as the initiation of translation, and in mRNA folding, acting as a chaperone preventing the formation of an mRNA secondary structure (14). However, in psychrophilic microorganisms, the synthesis of housekeeping gene products is not inhibited by cold shock, and the number of cold-shock proteins is usually higher and increases with the severity of the cold shock. In contrast, in these microorganisms, one set of proteins, known as cold-acclimation proteins (CAPs), is permanently induced by continuous growth at low temperatures (15). One acclimation protein is known to be analogous to CspA, a major cold-shock protein in Escherichia coli (16) but the cellular functions of acclimation proteins remain not well understood. Low temperatures promote unfavourable RNA secondary structures, which are likely to interfere with translation, and strengthen the interactions between DNA strands in the double helix and in the supercoiled state, therefore impairing unwinding and access to RNA polymerase. Nucleic-acid-binding proteins relieve the adverse effects of low temperatures and could play a central role in the cold adaptation of psychrophiles (17, 18). Other adaptive strategies developed by psychrophilic organisms involve the regulation of ion channel permeability, seasonal dormancy and microtubule polymerization (19–21). In addition to these cellular adaptations, a key adaptive strategy of psychrophiles is the modification of enzyme kinetics, allowing the emergence of metabolic rates compatible to life at low temperatures. Indeed, low temperatures strongly inhibit the rate of chemical reactions. The temperature dependence of chemical reactions is described by the Arrhenius equation $k = Ae^{-E_a/RT}$, in which $A$ is the pre-exponential factor (related to steric factors and molecular collision frequency), $E_a$ is the activation energy, $R$ is the gas constant (8.314 J K$^{-1}$ mol$^{-1}$) and $T$ is the absolute temperature in Kelvin. Thus, any decrease in temperature will induce an exponential decrease in the reaction rate and for most biological systems, a decrease of $10^\circ$C depresses the rate of chemical reactions by a factor ranging from 2 to 3 corresponding to $Q_{10}$ (which expresses the ratio of reaction rates measured at an interval of temperature of $10^\circ$C). Nevertheless, psychrophiles succeed to maintain an appropriate rate for enzyme-catalyzed reactions that are involved in essential cellular processes by synthesizing cold-active, but heat-labile, enzymes with an activity that is up to 10 times higher at low temperatures than that of their mesophilic homologues. The synthesis of such cold-active or psychrophilic enzymes is the main physiological adaptation at the enzyme level.

## II. PSYCHROPHILIC ENZYMES

### Structural factors involved in the stability and the activity of psychrophilic enzymes

**Structural adaptations related to stability** The attenuation in strength and number of structural factors known to stabilize proteins could be involved in the low stability of psychrophilic enzymes (6, 22, 23). One mutational study has shown that the accumulation of several mutations, which stabilize weak interactions in the psychrophilic $\alpha$-amylase, resulted in a strongly stabilized mutant displaying kinetic parameters close to those of the mesophilic homologue (24, 25). In the $\alpha$-amylase, the lower stability of the psychrophilic enzyme originates from a general weakening of intramolecular forces which contribute to the stability of the native protein molecule. In this case, the overall decrease in the thermostability of the cold-adapted protein provides a sufficient plasticity around the catalytic residues to render the enzyme efficient at low temperatures. The decrease in global stability has been observed in particular for psychrophilic enzymes which operate on large substrates such as $\alpha$-amylases and proteases. The lower stability of cold-active enzymes could also originate from weakened interactions in one or a few regions of the structure. Localized flexibility has been demonstrated by microcalorimetric studies of multidomain proteins (26–28). In psychrophilic chitobiase and the $\alpha$-amylase precursor, differences in stability can be found between regions carrying the active site and the other moieties of the protein: The active site is embedded in a heat-labile domain, whereas the stability of other domains is unaffected or even enhanced. In psychrophilic phosphoglycerate kinase, the two domains are connected by a hinge region that bends during catalysis forming the active site at the interface of both the heat-labile and heat-stable domains. The heat-labile domain could provide the required flexibility around the active site and favour the reaction rate, whereas the heat-stable domain, having the same stability as that of the mesophilic enzyme, could improve substrate binding as a result of its rigidity and appropriate comple-
mentarities with the substrate (28).

Each protein uses one or a combination of possible structural adjustments to reduce the local or global stability of the molecular edifice. In addition to mutational studies, the involvement of these factors in protein adaptation is suggested from comparative studies between psychrophilic enzymes and their heat-stable homologues based on homology models and 3D structures. An increased number of crystallographic structures of psychrophilic enzymes from bacteria and cold-adapted fishes have been previously elucidated (29–41). Among these structures, seven have been compared with mesophilic or thermophilic homologues. These include the bacterial xylanase (31), α-amylase (42), citrate synthase (32), malate dehydrogenase (35), alkaline Ca$^{2+}$-Zn$^{2+}$ protease (30), triose-phosphate isomerase (34), insect lysozyme (41), fish uracil-DNA glycosylase (29) and fish trypsin (37). Other structures should become available soon, as crystals of three other cold-active enzymes have been obtained (43–45).

Compared to their mesophilic and thermophilic counterparts, structural modifications found in psychrophilic enzymes involve the clustering of glycine residues (providing local mobility), a decrease in the number of proline residues in loops (providing enhanced chain flexibility between secondary structures), a decrease in the number of arginine residues which are capable of forming multiple salt bridges, and hydrogen bonds, as well as a lower number of ion pairs, aromatic interactions or hydrogen bonds, and the weakening of charge-dipole interactions in α-helices. Nonpolar core clusters can have a weaker hydrophobicity, making the protein interior less compact, the binding of stabilizing cofactors such as metal ions can be weaker and loose protein extremities appear to be preferential sites for unzipping. However, a few common trends in the structural strategies used in the adaptation to temperatures are observed: a decrease in the number of ionic interactions in psychrophilic enzymes compared to their mesophilic and thermophilic homologues, a decrease in the fraction of accessible surface of side chains and a reduced apolar fraction of the buried surface with decreasing apparent optimal temperature (23).

Comparisons of some psychrophilic enzymes with their mesophilic and thermophilic homologues showed not only that each cold-adapted enzyme adopts its own adaptive strategy but also that there is a continuum in the strategy of protein adaptation to temperatures as illustrated below. In the dimeric citrate synthase, as one ascends the temperature ladder, the subunit interface of this enzyme and loop regions are reinforced by complex electrostatic interactions, and there is a reduced exposure of the hydrophobic surface. There is a progressive pattern of stabilization through multiple additional interactions at solvent-exposed, loop and interfacial regions (46). A comparison of the structure of *Pseu-doalteromonas haloplanktis* Cel A from *Clostridium thermocellum* suggested that cold adaptation is achieved by a decrease in the number of salt bridges and an increased exposure of hydrophobic residues (31). Compared to its mesophilic and thermophilic homologues AKsub and AKste, the psychrophilic adenylate kinase Asklo has the most apolar exposed surface area, although it has the fewest apolar atoms. A strong correlation is found between the thermal stability and apolar buried surface area of the adenylate kinases (47). The determination of the composition of the accessible surface of DNA ligases Phlig and Ts lig showed a decrease in charged surface in the order thermophile to psychrophile (48). Such a decrease could lead to a decrease in the strength of the ion pair network at the surface of the psychrophilic ligase compared to its thermophilic homologue and could improve the resilience of the external shell of the cold-active enzyme.

**Structural adaptations related to activity** Psychrophilic enzymes are characterized by destabilizing structural adjustments but also by adaptations in their active site. Frequently, their catalytic site reveals an increased size and a better accessibility to ligands when compared to their mesophilic or thermophilic homologues (22, 30, 32). This is achieved in various ways such as by the deletion of loops bordering the active site, by the distinct conformation of these loops and by the replacement of bulky side chains by smaller groups at the entrance of the active site. This adaptation could allow the accommodation of the substrate at low energy cost and facilitate the release and exit of the reaction products. Nevertheless, the opposite situation can be observed: In psychrophilic xylanase pXyl, the accessibility of the substrate-binding region is substantially lower than that in the thermophilic homologue, which is mainly the result of a loop folding over the groove in pXyl (31). Although all reactive side chains as well as most side chains pointing towards the catalytic activity are strictly conserved in psychrophilic enzymes compared with their mesophilic and thermophilic homologues, changes occurring elsewhere in the molecule can be responsible for the optimization of the catalytic parameters of these enzymes as shown for the psychrophilic α-amylase (24, 49). Electrostatic potential in and around the catalytic site can be improved so that it may facilitate the interaction of the oppositely charged ligands with the surface of the enzyme (29, 31, 32, 35, 38). However, in psychrophilic DNA ligase Phlig, the adenylation domain contains a lower number of charged residues than that of the thermophilic enzyme Ts lig. This is in agreement with the fact that $k_{cat}$ is believed to be the most important adaptive parameter for Phlig, while for the DNA ligase Ts lig, the adaptive parameter is expected to be $K_m$ for nicked DNA, which is prone to melting at high temperatures. The adenylation domain of the psychrophilic enzyme is destabilized by an excess of hydrophobic surface residues (48). The flexibility of specific residues in the substrate-binding site also seems to be a cold-adapted feature (30, 31, 35). In the psychrophilic xylanase pXyl, two aromatic residues lining the subsites +1 and +2 were shown to adopt a double conformation. The increased flexibility of these residues may assist the accommodation of the substrate (31). In the psychrophilic cellulase Cel5G, structural adaptation to cellulose hydrolysis at low temperatures is observed in the unusually long linker and in the extended shape of the entire molecule compared with that of the mesophilic Cel45 from *Humicola insolens*. The unusually long linker of Cel5G can provide a significant steric optimization of the cellulolytic activity by increasing substrate accessibility (Sonan, G., personal communication).
Oligomeric enzymes Intersubunit interaction seems to be a major stabilization mechanism of hyperthermophilic oligomeric proteins. The hydrophobic core and ion-pair network at the subunit interface are the major stabilizing factors which stabilize the intersubunit interface (50). Moreover, oligomerization can be a significant stabilization mechanism for hyperthermophilic enzymes (50–53). On the basis of stability studies of dimeric globular proteins, it was calculated that quaternary interactions could provide 25% to 100% of the conformational stability in protein dimers (54).

A comparison of the E. coli 3-isopropylmalate dehydrogenase with the Thermus thermophilus enzyme showed a more hydrophobic subunit interface for the thermophilic protein (55). These hydrophobic interactions seemed to make the dimer more resistant to dissociation (56). Similarly, a comparison of the psychrophilic triosephosphate isomerase from Moraxella sp. TA137 and of the thermophilic TIM from Bacillus stearothermophilus with the enzyme from yeast shows three more hydrophobic residues in the B. stearothermophilus TIM which enhances the stability of its dimers (57). The alignment of the amino acid sequences of Moraxella sp. TA137 and B. stearothermophilus with respect to that of chicken, the 3D structure of which was already known, showed two small insertions in the loops taking part in the monomer–monomer interface of the dimeric psychrophilic enzyme. These insertions may increase the flexibility of the loops and thereby could lead to the destabilization of the dimer due to a greater loss of entropy upon dimer formation (57).

A comparison of the dimeric cold-active citrate synthase from the antarctic bacterium Arthrobacter DS2-3R with the homologous enzyme from the hyperthermophilic Pyrococcus furiosus shows a decrease in the number of subunit interactions, without any intersubunit ion-pair network in the cold enzyme. This difference and the presence of isoleucine and tyrosine clusters at the interface of the enzyme from P. furiosus may account for the difference in thermal stability of each enzyme (32). In the hyperthermophilic citrate synthase, salt bridges organized in networks are largely clustered in the active-site region and at the dimer interface. In the case of the enzyme from P. furiosus, electrostatic interactions may contribute to the integrity of both the protein dimer and the active site by possibly reducing conformational disorder at high temperatures. In contrast, in the psychrophilic enzyme, electrostatic interactions are more dispersed throughout the structure and the psychrophilic enzyme contains a greater number of charged residues with large destabilizing electrostatic free-energy contributions. The active-site charged residues of the psychrophilic enzyme show a greater extent of destabilization than those of the thermophilic enzyme and its charged residues on its surface may form favorable electrostatic interactions with the surrounding water molecules along with those among the charged residues themselves. Therefore, in the psychrophilic citrate synthase at low temperatures, electrostatics may contribute to enhance protein solvation and to ensure active-site flexibility (58).

A comparison of the modeled structure of the tetrameric psychrophilic β-galactosidase from P. haloplanktis (59) with the structure of its homologous mesophilic enzyme from E. coli (60) showed the disappearance of two ion pairs, as well as one hydrophobic cluster of 14 residues, at the level of the long interface of the cold-active enzyme, which is responsible for the formation of dimers. At the level of the activating interface, which induces the formation of tetramers and is responsible for the activity, the number of hydrogen bonds is decreased by a factor of five and one hydrophobic cluster is also deleted. This reduced number of stabilizing subunit interactions in the psychrophilic enzyme could be responsible for the decrease in its stability.

Kinetic optimization and heat-labile activity of cold-active enzymes The main physiological adaptation to cold of psychrophilic enzymes is a high activity ($k_{cat}$) at low temperatures which can be 10 times higher than that of their mesophilic homologues. Psychrophilic enzymes succeed to reduce the deleterious effect of low temperatures on enzyme reaction rates by reducing the temperature dependence of their specific activity ($k_{cat}$), which correlates with a decrease in the activation energy of chemical reactions. The catalytic constant $k_{cat}$ or turnover number corresponds to the maximum number of substrate molecules converted to product per active site per unit of time. In the Michaelis–Menten equation, the catalytic constant $k_{cat}$ is the first-order rate constant for the chemical conversion of enzyme-substrate complex ES to enzyme and product. The following equation is equivalent to the Arrhenius law and describes the effect of temperature on enzyme activity:

$$k_{cat} = \frac{k_0 T}{h} e^{-\Delta G^* / RT}$$

where $k_{cat}$ is the turnover number, $k_0$ is the transmission coefficient, $k_{0}$ is the Bolzmann constant (1.3805 $\times$ 10$^{-23}$ J K$^{-1}$), $h$ is the Planck constant (6.6256 $\times$ 10$^{-34}$ J s), $R$ is the gas constant, $\Delta G^*$ is the free energy of activation and $T$ is the absolute temperature. The free energy of activation $\Delta G^*$ is the variation of the Gibbs energy between the activated enzyme–substrate complex ES$^*$ and the ground-state ES according to the transition state theory which assumes the existence of an activated complex ES$^*$ in equilibrium with the ground-state ES:

$$E + S \leftrightarrow \text{ES} \leftrightarrow \text{ES}^* \rightarrow E + P$$

The thermodynamic parameters of the reaction ($\Delta G^*$, $\Delta H^*$, $\Delta S^*$) can be calculated as described by Lonhienne et al. (61). However, the activated state ES$^*$ does not really exist as the reagents only flow around this state and an ensemble of a multitude of activated conformations should be possibly considered. Moreover, the transition state theory contains several well-known simplifications and the real and absolute significance of the thermodynamic parameters has to be taken with caution (61). Particularly, the transmission coefficient $k$ is generally considered to be close to one and is, in general, ignored. However, as the viscosity of solvents markedly changes with temperatures, the frictional forces cannot always be neglected especially at low temperatures and should be taken into account according to the Kramers theory (62). Nevertheless, the errors inherent to the transition state theory have been attenuated by the analysis of the variation of the thermodynamic parameters obtained for psychrophilic enzymes with the variation of those obtained...
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for mesophilic enzymes: \(\Delta \Delta G^\text{p–m}\), \(\Delta \Delta H^\text{p–m}\), and \(\Delta \Delta S^\text{p–m}\) (61). The comparative studies show that psychrophilic enzymes are characterized by lower \(\Delta G^\text{p–m}\) values, compared to their mesophilic homologues. As expected, \(\Delta \Delta G^\text{p–m}\) is negative and its small value originates from large differences in the enthalpic \(\Delta \Delta H^\text{p–m}\) and entropic contributions \(\Delta \Delta S^\text{p–m}\) between psychrophilic and mesophilic enzymes (61). The decrease of the activation enthalpy \(\Delta H^\text{a}\) corresponds to the lower activation energy of the reaction and indicates a reduced temperature dependence of \(k_{\text{cat}}\). The low activation enthalpy \(\Delta H^\text{a}\) is the main adaptive parameter to low temperatures in a reaction catalyzed by psychrophilic enzymes and is achieved structurally by a reduced number of enthalpy-driven interactions that have to be broken to reach the transition state (63). Thus the enzyme regions that are involved in the conformational changes occurring during the catalytic reaction can provide sufficient flexibility to increase \(k_{\text{cat}}\) but should be less stable than that of mesophilic enzymes. Consequently, the activity of psychrophilic enzymes is heat-labile and this characteristic is generally well illustrated by the curves representing the thermal dependence of the specific activities of the cold-active enzymes and their heat-stable homologues (Fig. 1). Indeed, these curves show that the apparent maximal activity of cold-active enzymes is shifted towards low temperatures, suggesting a low stability of these enzymes. Interestingly, recent experiments have shown that psychrophilic enzymes are inactivated at temperatures well below that at which the protein unfolds (Fig. 2). This is not the case for their mesophilic and thermostable homologues which show a temperature of maximal activity corresponding, in general, to the unfolding transition suggesting that the observed loss of activity is due to protein unfolding (48, 64, 65). These results confirm the concept of “localized flexibility”, suggesting that the high activity at low temperatures is determined by a low stability at or near the active site of psychrophilic enzymes associated or not with a decrease in stability in the regions not involved in the catalysis. The entropic contribution \(\Delta \Delta S^\text{p–m}\) is always negative and could be a consequence of the active-site flexibility. Indeed, the ground-state enzyme–substrate complex ES occupies a broader distribution of conformational states than the mesophilic homologues. The entropic contribution of activation has an unavoidable counter effect on the acquisition of a specific activity of psychrophilic enzymes but retaining stability in regions of protein not involved in the catalysis could contribute to a reduction in \(T \Delta S^\text{p–m}\) (61).

These considerations lead to the idea that the thermode-
Dependence of the reaction rate is only dependent on the catalytic constant $k_{cat}$. However, an enzymatic reaction is characterized by two fundamental kinetic parameters, $k_{cat}$ and $K_m$ (which usually represents a measure of the affinity of the enzyme for the substrate), and the specific constant $k_{cat}/K_m$ is a better indicator of catalytic efficiency than $k_{cat}$ alone, especially for enzymes working at low substrate concentrations. Among psychrophilic enzymes (for compilations, see Refs. 22 and 66), two main trends are observed: improving $k_{cat}$ at the expense of $K_m$ (both $k_{cat}$ and $K_m$ increase) and improving the $k_{cat}/K_m$ ratio (increase in $k_{cat}$ and decrease in $K_m$). However, most psychrophilic enzymes have higher $K_m$ values than their mesophilic counterparts (67) and increase $k_{cat}$ at the expense of $K_m$. This weaker substrate binding is predicted to accompany the broader distribution of the ground-state ES if increases in $K_m$ are achieved through active-site resilience. If the substrate is weakly bound, the ground-state ES falls in a shallower energy pit, reducing the energy barrier $\Delta G^\circ$ of the reaction and increasing $k_{cat}$. This is generally observed in psychrophilic enzymes, such as $\alpha$-amylase, which display identical substrate-binding sites and active-site architecture when compared with their mesophilic homologues (42). A correlation between $k_{cat}$ and $K_m$ was demonstrated by reintroducing in the psychrophilic $\alpha$-amylase from P. haloplanktis, some stabilizing weak interactions that were missing when compared to the mesophilic enzyme from the pig pancreas (24). The stabilized mutants displayed decreased $k_{cat}$ and $K_m$ and support the concept of adaptive drift in the case of enzymes improving $k_{cat}$ at the expense of $K_m$. However, some enzymes react against this adaptive drift of $K_m$ in order to maintain or improve the substrate-binding affinity. The optimization of the substrate-binding affinity can be obtained by amino acid substitutions within the active site of the psychrophilic enzymes. This is the case for the cold-active extracellular chitinase from the psychrophilic Arthrobacter sp. TAD20 which improves the $k_{cat}/K_m$ ratio (increase in $k_{cat}$ and decrease in $K_m$) (26). In the mesophilic chitinase, two tryptophan residues are the main substrate-binding ligands, which perform hydrophobic interactions with the substrate. These residues are substituted by polar residues in the psychrophilic enzyme and this substitution could lead to the decrease in $K_m$ at low temperatures. Compared to hydrophobic interactions, the polar residues are able to perform, at least within a certain temperature range, stronger interactions as the temperature decreases. Indeed, two types of interaction are involved in the binding of a substrate by enzymes. Hydrophobic interactions occur endothermically around room temperatures and are weakened by a decrease in temperature. Interactions of an electrostatic nature (such as hydrogen bonds and ion pairs) occur exothermically and are stabilized by moderately low temperatures. Consequently, temperature will not only affect $k_{cat}$ but also $K_m$ and the effect of temperature on $K_m$ could depend on the type of interaction involved in substrate binding. Evolution through an increase in $k_{cat}$ and a decrease in $K_m$ should be particularly crucial for enzymes typically operating at subsaturating substrate concentrations, such as most intracellular enzymes. Some cold-active intracellular enzymes improve the $k_{cat}/K_m$ ratio (28, 59, 68) but some metabolic enzymes such as the psychrophilic dihydrofolate reductase from Moritella profunda sp. nov. (52) and the ornithine carbamoyltransferase from the psychrophilic Moritella abyssi (52) do not optimize the catalytic efficiency $k_{cat}/K_m$. Both heat-labile enzymes appear to be suboptimal in their physiological temperature range as far as the $k_{cat}/K_m$ ratio is concerned, supporting the concept that complete metabolic adaptation to cold may remain elusive (52). However, it should be noted that the improvement of the catalytic efficiency $k_{cat}/K_m$ could be related to the global or localized decrease in the stability of psychrophilic enzymes. Indeed, the psychrophilic $\alpha$-amylase and xylanase (global low stability) improve the $K_m$ value at low temperatures at the expense of the $K_m$ value, probably as a result of a less efficient substrate binding into the flexible active site. In contrast, the cold-active phosphoglycerate kinase (localized flexibility) displays an improved $K_m$ value with respect to its mesophilic homologue and it has been proposed that keeping a part of the protein rigid may improve the binding of the substrate (65).

**Stability of psychrophilic enzymes** Psychrophilic enzymes are characterized by their low stability in comparison with their mesophilic and thermophilic homologues. The lower stability of cold-active enzymes arises from the weakening of intramolecular forces contributing to the cohesion of the native protein molecule. The decreased stability of psychrophilic enzymes is well demonstrated by differential scanning calorimetry (DSC) which allows the direct analysis of thermodynamic parameters. The thermal unfolding of proteins adapted to extreme temperatures is particularly well illustrated by thermograms from the psychrophilic, mesophilic and thermophilic $\alpha$-amylases (64), xylanases (65) and DNA ligases (48) recorded by microcalorimetry (Fig. 3). The analysis of the parameters used to characterize the stability of proteins shows common trends. First, the unfolding of cold-active enzymes occurs at lower temperatures, as indicated by the melting point, $T_m$, at the top of the transition. Second, the area below the transition (calorimetric enthalpy of unfolding $\Delta H_{unf}$), representing the total heat absorbed during the unfolding, is much lower for psychrophilic enzymes and there is a clear trend for increasing $\Delta H_{unf}$ values from psychrophile to thermophile. Third, the unfolding of psychrophilic enzymes is often a cooperative, all-or-nothing process, devoid of stability domains, whereas, for example, mesophilic and thermophilic $\alpha$-amylases and DNA ligases show two and three transitions reflecting the presence of structural units of distinct stabilities. The analysis of the thermodynamic parameters of activation for the irreversible unfolding of $\alpha$-amylases (mutant N12R of the psychrophilic $\alpha$-amylase), xylanases and DNA ligases shows a higher denaturation rate for the psychrophilic enzymes and correspondingly a lower energy barrier ($\Delta G^\circ$) (Table 1). However, $AH$ and $TAS$ are highest for the psychrophilic enzymes, indicating that their lower thermostability is due to an unfavorable entropic contribution. These parameters show an increasing stability of mesophilic and thermophilic enzymes with temperature and support the concept of a continuum in protein adaptation to temperature. According to these results, the cold adaptation of the psychrophilic $\alpha$-amylase, xylanase and DNA ligase is clearly
correlated to the general weakening of intramolecular interactions, leading to an overall decrease in the thermostability of the protein. A decrease in global stability may in turn provide the appropriate plasticity around the catalytic residues necessary to adapt the catalytic efficiency to low temperatures. However, as previously mentioned, the reduced stability related to the low stability of the psychrophilic enzyme may not necessarily arise from the general reduction in the strength of intramolecular forces, but from the weakened interactions in one or a few important regions of the structure (localized flexibility) (26, 28, 32, 63). Lines of direct evidence of differences in stability between some domains and the entire structure have been provided by DSC studies (28, 26). In these cases, cold-adapted enzymes contain some elements controlling protein stability and affinity for the substrate and other elements conferring the required flexibility for an efficient catalysis at the environmental temperature.

Among psychrophilic enzymes showing a global reduction in stability, some proteins unfold according to a reversible pathway without any stable intermediates between the native and unfolded states. This is the case of the psychrophilic α-amylase from *P. haloplanktis* that unfolds reversibly according to a two-state mechanism, while its mesophilic and thermophilic homologues unfold irreversibly with a number of transitions. This reversibility associated with a two-state unfolding allows the study of the temperature-dependent thermodynamic stability function according to the Gibbs free-energy change between the denatured and native states (\(\Delta G_{\text{U-N}}\)). This reversible character can be related to the weak hydrophobicity of the core clusters in the cold-adapted enzyme and to the low melting temperature, at which hydrophobic interactions are restrained because, unlike mesophilic α-amylases, aggregation does not occur. The stability curves in Fig. 4 represent the Gibbs free energy of unfolding as a function of temperature and correspond to the energy required to disrupt the protein structure at any given temperature. Compared to mesophilic and thermophilic proteins, the cold-active α-amylase shows a global collapse of its stability curve and its conformational energy (\(\Delta G_{\text{U-N}}=92\text{ J/mol/residue}\)) is the lowest value reported for two-state proteins (70). The psychrophilic α-amylase has reached a state close to the lowest possible stability of the native state, thereby preventing any further decrease in stability. If an increase in flexibility is gained at the expense of stability, this implies that the actual native state precludes any further adaptation toward a more flexible structure.

### Table 1. Thermodynamic parameters for irreversible thermal unfolding

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>(\Delta G^\circ) (kJ mol(^{-1}))</th>
<th>(\Delta H^\circ) (kJ mol(^{-1}))</th>
<th>(T\Delta S^\circ) (kJ mol(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>pXyl</td>
<td>82.0</td>
<td>414.0</td>
<td>332.0</td>
</tr>
<tr>
<td>CellA</td>
<td>113.0</td>
<td>345.0</td>
<td>232.0</td>
</tr>
<tr>
<td>Phlig</td>
<td>93.5</td>
<td>422.2</td>
<td>328.7</td>
</tr>
<tr>
<td>Eclig</td>
<td>98.1</td>
<td>162.1</td>
<td>64.0</td>
</tr>
<tr>
<td>AHA</td>
<td>85.7</td>
<td>458.5</td>
<td>372.8</td>
</tr>
<tr>
<td>PPA</td>
<td>89.8</td>
<td>354.0</td>
<td>264.2</td>
</tr>
<tr>
<td>BAA</td>
<td>95.8</td>
<td>310.1</td>
<td>214.3</td>
</tr>
<tr>
<td>α-Amylases</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ec lig</td>
<td>98.1</td>
<td>162.1</td>
<td>64.0</td>
</tr>
</tbody>
</table>

\(<\text{kJ mol}^{-1}\text{res}^{-1}\>) is the lowest value reported for two-state proteins (70). The psychrophilic α-amylase has reached a state close to the lowest possible stability of the native state, thereby preventing any further decrease in stability. If an increase in flexibility is gained at the expense of stability, this implies that the actual native state precludes any further adaptation toward a more flexible structure.

FIG. 3. Thermal unfolding of the psychrophilic, mesophilic and thermophilic α-amylases (A), DNA ligases (B) and xylanases (C) recorded by differential scanning calorimetry (DSC). AHA, Psychrophilic α-amylase; PPA, mesophilic α-amylase; BAA, thermostable α-amylase; Phlig, psychrophilic DNA ligase; Ec lig, mesophilic DNA ligase; Ts lig, thermostable DNA ligase; pXyl, psychrophilic xylanase; Xyl1, mesophilic xylanase; CellA, thermophilic xylanase. Adapted from Refs. 48, 64, 65.
Consequently, the specific activity of the cold-adapted enzyme at the environmental temperature is still much lower than that of the mesophilic counterpart at 37°C, reflecting the imperfect adaptation of psychrophilic enzymes. The stability curve is a parabolic function limited by the temperature of cold denaturation $T_m$ (below 0°C) and by the high temperature melting point $T_m$ at which by definition $\Delta G_{m, f} = 0$ (71). The curve shows that psychrophilic proteins are both heat labile and cold labile. Indeed, a decrease in stability, indicated on the left side of the stability curve of the psychrophilic enzyme, leads to cold denaturation. The extrapolation of the curve of the cold-active $\alpha$-amylase below 0°C predicts cold unfolding at $-10^\circ$C in vitro. It is thought that cold denaturation arises from the hydration of protein groups which destabilizes the protein at decreasing temperature (72). The hydration of polar and aromatic groups, in particular, is associated with a negative free energy and destabilizes the protein structure. The group hydration responsible for the decrease in $\Delta G_{m, f}$ at low temperatures could contribute to the conformational flexibility of psychrophilic enzymes in their natural environment. This is different from mesophilic and thermophilic enzymes, the latter taking advantage of the conformational entropy rise with temperature to gain plasticity. Group hydration seems to play an essential role in the acquisition of flexibility as improved interactions with the solvent have been noted for some cold-active enzymes. However, in certain cases, the higher fraction of the apolar molecular surface and the lower accessible area may prevent the effect of cold denaturation (23, 72). The psychrophilic adenylate kinase AKglo has been observed to have an approximately 200 square Å more apolar surface area than the mesophilic AKsub and the thermophilic AKste, suggesting that protection against cold denaturation might include a design selection that has a more “strategically” pre-exposed hydrophobic surface so as not to risk losing native configurations due to this effect (47).

Relationships between activity and stability  Relationships between activity and stability are well illustrated by the folding funnel model (64). In this model, the conformational stability of psychrophilic and thermophilic enzymes is represented as a function of conformational diversity in schematic energy landscapes (Fig. 5). The height of the funnel, which represents the conformational stability (free energy of folding), is higher for the thermophilic enzymes and the upper edge of the funnel, occupied by the unfolded state in a random coil conformation, is larger for the cold-adapted enzymes. During folding, the free-energy level decreases as well as the conformational ensemble. The funnel slopes of psychrophilic enzymes are steep and smooth and reflect the cooperative unfolding without intermediates characterizing the cold-adapted enzymes as a result of fewer stabilising interactions and stability domains. In contrast, the thermophilic proteins pass through intermediate states corresponding to the local minima of energy. These minima are responsible for the ruggedness of the funnel slopes and for the reduced cooperativity of the folding–unfolding reaction. The bottom of the funnel, which represents the stability of the native state ensemble, can be depicted as a single global minimum with a high energy barrier for a very stable and rigid thermophilic protein, whereas the bottom for an unstable and flexible psychrophilic protein is rugged and depicts a large population of conformers with a low energy barrier between them. The rigidity of the native state is therefore a direct function of energy barrier height (73, 74). In this context, the activity–stability relationships in these extremozymes depend on the bottom properties. Indeed, it has been argued that upon substrate binding to the association-competent subpopulation, the equilibrium between all conformers is shifted toward this subpopulation, leading to the active conformational ensemble (73–75). In the case of the rugged bottom of psychrophilic enzymes and in contrast with thermophilic proteins, this equilibrium shift only requires a modest free-energy change (low energy barriers) and a small enthalpy change for the interconversion of the conformations, but is accompanied by a large entropy change for fluctuations between the wide conformer ensemble.

The site-directed mutagenesis of the psychrophilic $\alpha$-amylase has provided strong experimental support for the adaptive relationships between stability and activity (24, 25). Mutants have revealed the individual contribution of a weak interaction and of an additional disulfide bond to protein stability as shown by the increases in $T_m$ (melting point) and
ΔH_{d(2)} (calorimetric enthalpy), by the modifications of unfolding cooperativity and reversibility, by the appearance of stability domains, and by the modification of the thermal inactivation rate constant k_{i} and of the kinetic parameters k_{cat} and K_{m}. The stabilization of the psychrophilic enzyme seems to decrease the activity by altering the dynamic properties of residues in the active site, but increases the affinity for the substrate, probably by improving its binding into the more rigid catalytic center of the mutant.

In contrast, laboratory evolution has shown that, in few cases, the stability and activity of proteins are not strictly linked, giving rise to the proposal that the low stability of cold-active enzymes results from a genetic drift originating from the lack of selective pressure for stable proteins (76). Directed evolution experiments can indeed yield synthetic catalysts that are thermostable and active either at high (77) or low (78, 79) temperatures. However, when random mutants of mesophilic and thermophilic enzymes were only screened for high activity at low temperatures during directed evolution (80–85), the evolved enzymes generally display a concomitant reduced thermostability. Moreover, in multidomain psychrophilic enzymes that contain both catalytic and noncatalytic domains as described for the psychrophilic chitobiase (26), the catalytic domain is always heat-labile, whereas the noncatalytic domain can be stable, even more than that of mesophilic proteins. It is therefore unlikely that genetic drift affects only the catalytic domain without modifying other regions of the protein. If it appears possible, at least under laboratory conditions, to uncouple activity and stability, enzymes displaying a high stability together with a high flexibility and activity do not exist in nature, except for thermophilic enzymes that catalyze the conversion of labile metabolic intermediates (86).

Thus the low stability of psychrophilic enzymes could be the simplest adaptive strategy that provides a gain in activity in the absence of selection for stable proteins in cold environments but in the presence of a strong selective pressure for highly active enzymes.

**Flexibility strategy** Maintaining the appropriate balance between molecular stability and structural flexibility is necessary for protein functions. Stability is needed to ensure the appropriate geometry for ligand binding, as well as to avoid denaturation, while flexibility is necessary to allow a catalysis at a metabolically appropriate rate (87). Psychrophilic enzymes are characterized by an increase in the plasticity or flexibility of all or some parts of the molecular structure in order to compensate for the lower thermal energy provided by the low-temperature habitat (88). This flexibility could be related to the low stability characterizing cold-adapted enzymes. Indeed, recent experiments involving the dynamic quenching of fluorescence, which shows an increased permeability to a small quencher, strongly support the concept of high flexibility of psychrophilic enzymes (48, 64, 65). In all cases, fluorescence quenching performed on psychrophilic, mesophilic and/or thermophilic enzymes revealed that the variation of fluorescence quenching between low and moderate temperatures decreases in the order psychrophile → mesophile → thermophile (Fig. 6). These results indicate a greater permeability of cold-adapted enzymes to the quencher, reflecting a strong correlation between flexibility and conformational stability.

In this technique, the decrease in fluorescence arising from diffusive collisions between the quencher and the fluorophore reflects the ability of the quencher to penetrate accessible regions of the protein structure and can consequently be viewed as an index of protein permeability. This is a dynamic process that allows the measurement of con-
formational motions over a large time scale (picoseconds to seconds). Nevertheless, it would be interesting to define the flexibility of cold enzymes in terms of type, amplitude and time scale of molecular motions. These aspects are now being studied by neutron scattering, which provides a unique tool to study atomic thermal motions in macromolecules, because neutron wavelengths and energies, respectively, match motion amplitudes and frequencies (89). Measured variables are the mean square amplitudes in a given time scale, as a function of temperature, from which an effective mean force constant, determining macromolecular resilience, can be calculated (90). Nevertheless, at this stage, the relationships between activity, stability and flexibility remain poorly understood in psychrophilic enzymes. These relationships have found some experimental support from the comparison of thermophilic and mesophilic enzymes (91–93), but the results obtained for probing the protein flexibility strategy in psychrophilic enzymes are controversial (87, 94, 95). This could be related to the technique used in these studies, global hydrogen–deuterium exchange measured over inappropriate time scales, while it is likely that the increase in the flexibility of cold-adapted enzymes is local and on the microsecond to millisecond time scale, i.e., directly related to the active site and catalytic activity. However, some studies report that thermostability is not incompatible with high activity at moderate temperatures (86, 96, 97) because protein stability can be achieved through an increase in the entropy of the native state as in the case of the β-glucosidase from Sulfolobus solfariticus (98). Moreover, other results show that relationships between flexibility and stability or activity are more complex than expected intuitively (99–101).

Potential for biotechnological applications One of the main interest of psychrophilic enzymes in biotechnological applications is the lack of requirement for expensive heating steps and consecutive economic benefits through energy saving. Other economic benefits originate from their abilities to function in a cold environment, to increase the reaction yield, to accommodate a high level of stereospecificity, and to minimize undesirable chemical reactions that can occur at higher temperatures (66). Moreover, they prevent the requirements of an increase in enzyme concentration to compensate for the lower efficacy when using mesophilic enzymes at low temperatures. These enzymes are also characterized by their thermal lability which allows an easy and rapid inactivation when required. This rapid inactivation of cold-active enzymes by mild heat treatment preserves product quality, permits selective enzyme inactivation in a complex medium, and does not require expensive heating/cooling systems (102). The ability to heat-inactivate cold-active enzymes is particularly useful in food industry where it is important to prevent any modifications of the original heat-sensitive substrates and products. This is also useful in sequential processes such as molecular biology where the actions of an enzyme must be terminated before the next process is undertaken (66). The biotechnological potential of psychrophilic enzymes is enhanced by the demonstration that recombinant cold-active enzymes are undistinguishable from their wild-type parent molecules with regards to kinetic parameters, folding properties (103) and three-dimensional structures (42), when the enzymes are expressed at sufficiently low temperatures (15–18°C). Moreover, expression at low temperatures is now possible as shown by the construction of a host-vector system that allows the overexpression of genes in psychrophilic bacteria (104). This study revealed a new biotechnological potential of psychrophilic strains as the expression at low temperatures prevents the formation of inclusion bodies and protects heat-sensitive gene products. Other host-vector systems for a temperature-regulated gene expression, allowing the overproduction of thermolabile enzymes originating from psychrophiles, have also been patented (Schwedel, T., patent DE10101266, 2002; Kann, T. and Schweder, T., patent EP1224307, 2002). In other respects, the improvement of the competencies of psychrophilic enzymes, according to the requirements of a biotechnological process, could be obtained by directed mutagenesis or perhaps more quickly by the directed evolution of the recombinant enzyme. Possible applications of cold-adapted enzymes can be found in numerous biotechnological and industrial fields as tools in molecular biology, food, drinks, textile and detergent industries, and bioremediation (for review, see Refs. 66, 102, 105). However, despite a large number of possible biotechnological applications, only a few cold-adapted enzymes are in commercial use. The costs of production and processing at low temperatures remain probably higher than those for the commercial enzymes that are presently in use. Nevertheless, patent protection for enzymes from psychrophilic organisms is slowly increasing (Hasan, A.K.M.Q. and Tamiya, E., patent WO9730172, 1997; Hasan, A.K.M.Q. and Tamiya, E., patent US6200793, 1998; Kubota, H. et al., patent WO9743406, 1996; François, J.M. et al., patent WO0104276, 2001; Genot, B. et al., patent WO2004023879, 2004).

III. CONCLUSIONS
The actual availability of crystal structures of psychrophilic, mesophilic and thermophilic homologous enzymes enables the investigation of the different aspects of protein adaptation to cold and the continuum in the strategy of protein adaptation to temperature. Moreover, the availability of the first genome-sequence data enables the investigation of more psychrophilic aspects of such as temperature sensors (18). The high activity characterizing psychrophilic enzymes is the main adaptive parameter to low temperatures and seems to be achieved by the destabilization of the active site or of the entire protein structure, allowing the catalytic center to be more flexible at low temperatures. However, the relationships between activity, flexibility and stability are still the subject of controversy. A definition of flexibility in terms of the type, amplitude and time scale of molecular motions related to the catalytic efficiency would be helpful in understanding these relationships. It was demonstrated that each psychrophilic enzyme adopts its own adaptive strategy. In terms of protein evolution, one could say that the adaptive strategy observed in natural cold environments could be, at least in some cases, a result of the lack of selective pressure for stable proteins, in conjunction with a strong selection for highly active enzymes. In addition to a powerful potential in fundamental research and particularly in protein dynamic and folding investigations, psychrophilic enzymes have a
high potential for biotechnological purposes.

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