

The thermal limits to life on Earth

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Abstract: Living organisms on Earth are characterized by three necessary features: a set of internal instructions encoded in DNA (software), a suite of proteins and associated macromolecules providing a boundary and internal structure (hardware), and a flux of energy. In addition, they replicate themselves through reproduction, a process that renders evolutionary change inevitable in a resource-limited world. Temperature has a profound effect on all of these features, and yet life is sufficiently adaptable to be found almost everywhere water is liquid. The thermal limits to survival are well documented for many types of organisms, but the thermal limits to completion of the life cycle are much more difficult to establish, especially for organisms that inhabit thermally variable environments. Current data suggest that the thermal limits to completion of the life cycle differ between the three major domains of life, bacteria, archaea and eukaryotes. At the very highest temperatures only archaea are found with the current high-temperature limit for growth being 122 °C. Bacteria can grow up to 100 °C, but no eukaryote appears to be able to complete its life cycle above \( \sim 60 \) °C and most not above 40 °C. The lower thermal limit for growth in bacteria, archaea, unicellular eukaryotes where ice is present appears to be set by vitrification of the cell interior, and lies at \( \sim -20 \) °C. Lichens appear to be able to grow down to \( \sim -10 \) °C. Higher plants and invertebrates living at high latitudes can survive down to \( \sim -70 \) °C, but the lower limit for completion of the life cycle in multicellular organisms appears to be \( \sim -2 \) °C.

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Introduction

If we wish to assess the possibility of life elsewhere in the universe, we need first to understand the physical factors that constrain the existence of life in the only example available for study, namely here on Earth. Limits to life could theoretically be set by one of any number of environmental factors, or combination of these, such as pressure, pH, ionizing radiation, or the presence of particular elements or compounds. In this paper, I examine the limits to life on Earth set by one such factor: temperature.

Temperature attracts particular attention from astrobiologists because it sets the boundary conditions for the existence of liquid water, which is known to be essential for life on Earth. Astrobiologists have considered the possibility of life based on solvents other than water (Clarke 2003; Bains 2004; Benner et al. 2004; Ward & Baross 2007), but at present these remain purely theoretical possibilities. The search for exoplanets that might support life is thus essentially a search for rocky planets with surface temperatures allowing the presence of liquid water.

Temperature is a thermodynamic state variable and appears in all of the equations of state that describe the universe. It affects every aspect of existence, as it influences all of the physical features of the environment as well as the inside of the cell (Clarke 2003). Examples of physiologically important physical features governed by temperature include the viscosity, density, pH, and dielectric permittivity of water, as well as its capacity to dissolve solutes and gases.

This pervasive influence of temperature sets a profound methodological challenge to any investigation of the effects of temperature on life. Because every aspect of the physical environment is affected by temperature, it is impossible to design a fully controlled experiment to explore the effects of temperature, where the only difference between the treatment and the control is a difference in temperature. For example, two aquatic animals exposed to different temperatures will also experience differences in viscosity, gas solubility and pH. Similarly two terrestrial plants exposed to different temperatures will also be subject to differences in relative humidity, gaseous diffusion rates and air density. This makes the interpretation of many experimental investigations of temperature challenging, and sometimes equivocal.

Definitions and principles

Life, like time, is an elusive concept that we feel we understand intuitively and yet find difficult to explain to others. But if we are to discuss life as a general concept, then we need to know exactly what we are talking about; the problem here is that philosophers and scientists have yet to produce a universally agreed definition of life (Benner 2010). What we can do, however, is arrive at a working description of life on Earth.


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Table 1. A working description of life on Earth

<table>
<thead>
<tr>
<th>Feature</th>
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<tbody>
<tr>
<td>General features of a living entity</td>
</tr>
<tr>
<td>Isolated from the rest of the universe by a partition</td>
</tr>
<tr>
<td>An internal composition that is non-equilibrium thermodynamically, maintained by a flow of materials and the dissipation of energy</td>
</tr>
<tr>
<td>Reacts to changes in the external world</td>
</tr>
<tr>
<td>Reproduces itself, based on a set of internal instructions</td>
</tr>
<tr>
<td>Specific features of life on Earth</td>
</tr>
<tr>
<td>Based on a few small, common atoms (C, H, N, O, P, S), covalently bonded</td>
</tr>
<tr>
<td>Energy supplied and transferred as electrons</td>
</tr>
<tr>
<td>Transition metals important in internal transfer of energy</td>
</tr>
<tr>
<td>Energy stored in proton gradients (short-term) or covalent bonds (long-term)</td>
</tr>
<tr>
<td>Metabolism utilizes a limited number of basic molecules</td>
</tr>
<tr>
<td>Metabolism driven by small changes in free energy</td>
</tr>
<tr>
<td>Covalent bonds thermally stable, non-covalent bonds break and reform rapidly</td>
</tr>
<tr>
<td>Water provides the solvent for many molecules, is involved as a reactant in all four major chemical transitions involved in metabolism (oxidation, reduction, condensation, hydrolysis), and typically proves the electrons and protons used for energy transport and storage within the cell</td>
</tr>
</tbody>
</table>

Table 1 provides one such description, emphasizing those aspects which are important in terms of understanding what sets the thermal limits to its continued existence.

The first four items comprise the essential general features of life as we know it on Earth, and may be characterized as a containment, metabolism and programme (CMP) description of life (Bedau 2010). It implies constraints on the physical size of a living entity, which cannot be too small (to avoid fatal stochastic imbalances in the internal environment) or too large (such that internal integration becomes compromised).

Almost all discussions of the nature of life refer back to Erwin Schrödinger’s classic short book *What is life?* (Schrödinger 1944) in which he emphasized the role of free energy and entropy in maintaining the viability of living organisms and explored the nature of the chemistry underpinning the coding of genetic information. At that time the precise chemical nature of the genetic material was unknown, and was widely believed to be protein. It was only later that it was shown to be nucleic acid (Avery et al. 1944), and the nature both of the code and its manner of replication recognized with the elucidation of the structure of DNA (Watson & Crick 1953).

Although prescient in his analysis of the thermodynamic aspects of life, Schrödinger confounded the two important features of heredity, namely the code itself and the translation and execution of that code. Von Neumann corrected this error in his consideration of the theory of automata (von Neumann 1951, 1966), emphasizing that the genetic material must both be copied, so that its integrity is preserved, and translated (that is, interpreted). Von Neumann’s analysis thus recognized the critical distinction between what we would now call software (the coded instructions embodied in nucleic acids) and hardware (the protein machinery produced by translating and interpreting that software).

Taking these insights together suggests a tripartite description of life (Fig. 1). A living entity on Earth comprises three essential features: the genetic information encoded in nucleic acids (software), the structural elements of the cell in the protein and lipid architecture, and the protein functional elements (hardware), and energy flux. This last dynamic element is required by thermodynamics: the interior of the cell is maintained in a thermodynamically non-equilibrium state and for this locally low entropy environment to be maintained there must be a continuous dissipation of energy. Without this, the internal composition of the cell cannot be maintained, the organism dies and free energy is then dissipated as its internal composition decays to the equilibrium state.

Some authors have used such thermodynamic considerations as the basis of a definition of life (e.g. Lovelock 1965). Such definitions, however, also apply to systems that generate order through the dissipation of energy but which we would not regard intuitively as alive.

An organism is alive only if it exhibits all three of the features shown in Fig. 1. A virus has only software: it contains the genetic instructions for the construction of new viruses, but lacks the machinery to do this and exhibits no energy flow. Viruses are obligate parasites of cells, which they utilize for both the machinery of construction and the energy flux to drive that construction. Based on the description of life in Table 1 and Fig. 1, viruses are not alive.

A resting spore, such as those of bacteria, fungi, eukaryotic algae or the brine shrimp *Artemia salina* contains both the software (DNA) and the hardware (proteins, membranes), but exhibits no energy flux. Like many plant seeds, tardigrade tuns or dehydrated nematodes, *Artemia* spores have the potential (or capacity) to be alive when circumstances change, but they are not alive in themselves. This context dependence of the living state sets an interesting challenge for those wrestling with the definition of life.

A living entity thus has software, hardware and exhibits energy flux (metabolism). While necessary, however, these are not sufficient. Thermodynamic considerations dictate that no living entity can perpetuate itself forever (Bains in press), and the fourth essential component of life is thus reproduction.
This reflects a second key distinction identified by von Neumann (1951, 1966), namely that between metabolism and replication. All living entities on Earth reproduce themselves; for a bacterium or archaean this means cell division to produce daughter cells, for a sexually reproducing eukaryote it means completion of the cycle from zygote to zygote.

Given that the copying of the internal instructions in DNA will always be imperfect, and in a resource-limited world not all offspring will survive, evolutionary change is inevitable. Evolutionary change starts with variation in the genetic material, but also requires that this variation results in changes in organism performance (fitness), and that the changes in the genetic material are heritable. A subtle but important feature here is that the basis of the description of life on Earth in Table 1 is the individual entity, and an individual organism does not evolve. Neither does a change in genetic makeup between parent and offspring constitute evolution. Evolution is the change in the frequency of genes in a defined group of individuals over time: it is a feature of populations, not of individuals. Darwinian evolution is thus a population consequence of the general features of life, rather than a defining characteristic of an individual entity (as proposed in the NASA definition of life: Luisi 1998).

**Temperature thresholds**

In relation to life on Earth, we can therefore draw an important distinction between two different threshold temperatures: a threshold for completion of the life cycle, \( T_L \), and a threshold for metabolism, \( T_M \) (Fig. 2). Between \( T_L \) and \( T_M \), the organism is viable and metabolizing, but unable to complete its life cycle. This might be because of insufficient energy to produce gametes, or initiate cell division, or it might be caused by a temperature-related failure of a key physiological process involved in reproduction but which is not in itself lethal.

A third important threshold is that for survival, \( T_S \). This does not exhibit a consistent relationship with \( T_L \) or \( T_M \); for example the lower \( T_S \) may be at \( T_L \) or \( T_M \). Thus in a chilling-sensitive organism, where a critical physiological system fails at a relatively high temperature, and \( T_S \), \( T_L \) and \( T_M \) may even coincide. By contrast, in a cold-hardy organism, \( T_S \) may be below \( T_M \). Between \( T_M \) and \( T_S \) the organism is in a state of suspended animation, but can resume metabolism once the temperature increases and recrosses the \( T_M \) threshold. An important difference between high- and low-temperature limits for life is that at high temperatures \( T_S \) is always reached; in contrast at low temperatures an organism may never reach its \( T_S \), even when taken close to absolute zero (Morowitz 1968).

The upper and lower \( T_L \) thresholds thus mark the limits to life on Earth, as existence outside these boundaries does not allow for completion of the life cycle. The \( T_L \) thresholds are those which define a habitat capable of sustaining life, and are thus the temperature thresholds of most significance to astrobiology. While studies of extremophile microbes have tended to concentrate on establishing \( T_L \) by determining the thermal limits to cell division, studies of plants and animals have tended to concentrate on limits to survival (\( T_S \)). Coupled with the highly variable thermal environment on land, this makes determination of \( T_L \) in these organisms very difficult, but in terms of defining the thermal thresholds for life, it is \( T_L \) we need to know.

**Domains of life**

Living organisms on Earth range in size and complexity from bacteria and small mycoplasma to whales and redwoods. These organisms show a wide range of internal structure and, in the case of multicellular organisms, tissue architecture. We should therefore not expect them all to respond to temperature in the same way, and indeed they do not. However, life on Earth can be grouped into a relatively small number of functional groups, based on their response to temperature (Table 2).

**Temperature extremes on earth**

An important context for understanding the thermal limits to life on Earth is set by the range of temperatures available. If we find habitats with liquid water but no life, then we can conclude that these habitats may exceed the temperature limits for life. If on the other hand, we find life everywhere we cannot say whether other, more extreme, environments might harbour life; we may not have sampled the full range of temperatures to which life could adapt.

Life evolved in the sea, and the emergence of first plants and then animals onto the land was one of the most important steps in the evolution of life on Earth. The thermal challenges of land and sea are, however, very different, and the two realms therefore need to be considered separately.

The mean surface temperature of the Earth over the period 1980–2013 was 14.4 °C (data from the Reanalysis Project of the European Centre for Medium-Range Weather Forecasts), but there is much variation with latitude, altitude and season. In addition, the data show an overall increase in mean global temperature from 14.1 °C in the early 1980s to 14.5 °C in 2012. The lowest air temperature yet measured at the surface of the Earth is −89.2 °C, recorded at Vostok Station, Antarctica on 21 July 1983 (Bodetsky 1984). The lowest mean monthly
temperature for Vostok is $-65 \, ^\circ C$ (August), indicating that this was a truly unusual event. Turner et al. (2009) discuss the atmospheric conditions leading to this record low, and speculate that at nearby Dome A similar circumstances could lead to surface air temperatures as low as $-100 \, ^\circ C$.

Vostok and Dome A are deep in the interior of Antarctica. There is no native biota at either site, though this is likely the result of the isolation and the lack of habitat rather than temperature per se. It is possible that the biota of isolated nunataks (mountain peaks exposed above the ice plateau) in continental Antarctica experience temperatures similar to those of continental locations such as Vostok, but as yet the data are lacking. The lowest temperature for a location with native flora and fauna is $-67.7 \, ^\circ C$, recorded for two locations in the Sakha (Yakutia) Republic of Russia: Verkhoyansk on 5 and 7 Feb 1892, and at Oymyakon on 6 Feb 1933. Earlier records of $-69.8 \, ^\circ C$ and $-68.8 \, ^\circ C$ at Verkhoyansk in 1892 are now regarded as unreliable. The daily mean temperatures for January are $-45.5 \, ^\circ C$ (Verkhoyansk) and $-46.4 \, ^\circ C$ (Oymyakon) (all data from the World Meteorological Association).

The interior of Alaska also gets very cold in winter and here the record low is $-62.2 \, ^\circ C$ recorded at Prospect Creek on 23 Jan 1971 (data from NOAA). This is only just above the record low for North America, which is $-63.4 \, ^\circ C$ recorded at Snag airport, Yukon, on 3 Feb 1947. The interior of continental North America also gets very cold in winter, and there are record low temperatures below $-50 \, ^\circ C$ recorded for many states such as Montana, Utah, Wyoming and Colorado.

The highest surface air temperature ever recorded is 56.7°C, measured at Furnace Creek Ranch (formerly Greenland Ranch), Death Valley, California on 10 July 1913, a previous record of 58°C from El Azizia, Libya, now being regarded as unreliable (El Fadli et al. 2013). These are, of course, meteorological air temperatures taken at 1.5 m above the ground surface. While these temperatures are relevant for familiar animals and plants, many organisms are small or very small, and live on or very close to the ground surface. Here temperatures can be very different, and satellite observations indicate that surface temperatures frequently exceed 50°C. Surface temperatures reflect heating of the ground by solar radiation and the highest temperatures will occur where skies are clear, the soils are dry and winds are light. The hottest surface temperatures on Earth determined by remote sensing are regularly recorded from the Lut Desert in Iran, where in 2005 a record of 70.7 °C was achieved (data from NASA MODIS satellite).

The oceans are the largest habitat on the face of the Earth. The overall range of temperatures is much narrower than for terrestrial habitats and the high thermal capacity of water means that rates of change, both across space and in time, are much slower than in air. Surface oceanic waters vary in temperature with depth, latitude and season. The enormous size of the oceans, coupled with their great depth, make it difficult to calculate an overall temperature for seawater. Below the seasonally warmed and cooled surface layers, seawater remains permanently between $-1 \, ^\circ C$ and $+4 \, ^\circ C$, except for the deep Mediterranean and Red Sea and the immediate environs of hydrothermal vents (Gage & Tyler 1991). Overall, the mean temperature of oceanic water is $<4 \, ^\circ C$; the largest habitat on the face of the Earth is cold.

Surface oceanic waters can reach $\sim 30 \, ^\circ C$ in tropical regions and $-1.96 \, ^\circ C$ (the freezing point of seawater at atmospheric pressure) in high latitudes. In waters close to the Antarctic continent, bottom temperatures can be $\sim -2 \, ^\circ C$ year-round (Clarke et al. 2009).

### High-temperature limits to life

The earliest record of life in hot springs appears to be that of Pliny the Elder, who noted in his Natural History (probably written around 77–79 AD) that green plants could be found growing in the hot springs at Padua. These springs are still active, and undeveloped sections contain cyanobacterial mats (which are presumably the ‘green plants’ recorded by Pliny) (Brock 1978).

There are many places on Earth where geothermally heated water reaches the surface, including Iceland, New Zealand, Kamchatka and the hydrothermal vent fields associated with the spreading ridge systems of the deep ocean. However, the basis of our understanding of microbial life at high temperatures stems from the pioneering work of Thomas Brock and colleagues in Yellowstone Park (Brock 1978). The value of Yellowstone as a location for this work is the existence of a

<table>
<thead>
<tr>
<th>Domain</th>
<th>Key features</th>
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<tbody>
<tr>
<td>Archea</td>
<td>No nucleus, free ribosomes, ether-linked membrane lipids</td>
</tr>
<tr>
<td>Bacteria</td>
<td>No nucleus, free ribosomes, acyl-linked membrane lipids</td>
</tr>
<tr>
<td>Eukarya</td>
<td>Nucleus, mitochondria, ribosomes on internal membranes, acyl-linked membrane lipids</td>
</tr>
<tr>
<td>Unicellular eukaryotes</td>
<td>Cell membrane in direct contact with external environment</td>
</tr>
<tr>
<td>Multicellular eukaryotes</td>
<td>Cell membrane exposed to internal body environment</td>
</tr>
<tr>
<td>Lichens</td>
<td>Symbiosis between fungal hyphae and algae; often highly resistant to dehydration</td>
</tr>
<tr>
<td>Mosses and liverworts</td>
<td>Non-vascular (no xylem), absorbing water and nutrients through leaf surfaces</td>
</tr>
<tr>
<td>Higher plants</td>
<td>Vascular tissues for distributing resources through plant</td>
</tr>
<tr>
<td>Invertebrates</td>
<td>No backbone; cardiovascular system for distributing nutrients</td>
</tr>
<tr>
<td>Ectothermic vertebrates (fish, amphibians, reptiles)</td>
<td>Cardiovascular system for distributing nutrients</td>
</tr>
<tr>
<td>Endothermic vertebrates (mammals, birds)</td>
<td>Maintain internal body temperature above $\sim 30 , ^\circ C$</td>
</tr>
</tbody>
</table>

Table 2. The three major domains of life on Earth, with Eukarya subdivided into categories with differing features of potential importance to their thermal ecology
large number geothermal springs with a wide range of temperatures, pH and mineral contents. Although Brock himself ceased working in there in 1975, Yellowstone continues to provide important insights into thermal physiology and ecology. Brock first reported microbes growing at high temperatures in the 1960s (Brock & Brock 1966; Brock 1967). Shortly afterwards together with a colleague Hudson Freeze he isolated a bacterium growing at 70 °C which they described as Thermus aquaticus (Brock & Freeze 1969). This organism has proved to be of enormous significance as the source of the DNA polymerase which underpins the genomic revolution. Based on his experiences of life in hot springs in Yellowstone and elsewhere, Brock (1978) predicted that life would be found wherever water was liquid, a prediction that was vindicated spectacularly by the discovery of microbial life in the range 65 °C (Sokolova et al. 2006). This difference between archaea and bacteria in sensitivity to high temperatures is evident in the distribution of the two groups within active vents, where there can be a transition from a mixed assemblage of archaea and bacteria near the cooler exterior of the chimney, to primarily archaea in the hotter interior (Schrenk et al. 2003).

The very hot water, often at temperatures >300 °C, that emerges from hydrothermal vents mixes with the local seawater and this leads to very strong thermal gradients. The very hottest waters appear to be abiotic, but areas where the water has cooled are characterized by extensive microbial mats. These have yielded a wide range of hyperthermophiles, both archaeans and bacteria. Hyperthermophiles are defined as organisms with an optimal growth temperature at or above 80 °C. They require liquid water so growth above 100 °C is possible only where pressure keeps water liquid. They are found in a wide variety of terrestrial and marine habitats, all associated with geothermal sources of heat. The low solubility of oxygen at these high temperatures and the frequent presence of large amounts of reducing gases mean that most habitats for hyperthermophiles are anaerobic. Most hyperthermophiles utilize inorganic redox reactions as the sources of energy, and CO₂ as the sole carbon source.

The current record for a high-temperature growth is Methanopyrus kandleri, originally isolated from a vent in the Gulf of California and found to grow between 84 and 110 °C (Huber et al. 1989; Kurr et al. 1991). However a strain of M. kandleri isolated subsequently from the Kairei vent field on the Central Indian Ridge was found to grow at 122 °C under 40 MPa pressure (Takai et al. 2008), just surpassing the previous record for 121 °C for Geogemma barossii strain 121 (Kashefi & Lovley 2003).

Microbes growing at the very highest growth temperatures all appear to be archaeans, but there are some bacteria are able to grow to ~100 °C, with the current record being Geothermobacterium ferrireducens, which was isolated from Obsidian Pool in Yellowstone National Park (Kashefi et al. 2002). Two other taxa, Aquifex aeolicus and Thermotoga maritima can grow at 90 °C or above (Table 3), and there are a range of Fe(III)-reducing thermophilic bacteria with $T_L$ values in the range 65–75 °C (Sokolova et al. 2006). This difference between archaea and bacteria in sensitivity to high temperatures is evident in the distribution of the two groups within active vents, where there can be a transition from a mixed assemblage of archaea and bacteria near the cooler exterior of the chimney, to primarily archaea in the hotter interior (Schrenk et al. 2003).

The water issuing from hydrothermal vents can exceed 300 °C, and with the surrounding seawater at less than 4 °C this leads to very steep and highly variable thermal gradients that make it extremely difficult to quantify the precise thermal environment of the microbes growing there. Schrenk et al. (2003) reported intact microbes growing in water ‘that must have been well in excess of 100 °C’ but the only reliable technique to establish unequivocal growth temperatures is culture in vitro. Baross et al. (1982) sampled the microbial flora from hydrothermal vents along the East Pacific Rise, where
The current accepted upper limit for microbial growth is 122 °C. Below that limit for microbial growth may be as high as 150 °C; the thermal limit for microorganisms can be as high as 150 °C, the upper limit for the rhodophyte *Cyanidium caldarium*, although optimal (maximum) growth was at 45 °C (Doemel & Brock 1970, 1971). Over a century ago, however, Dallinger (1887) reported an experiment in which he raised the temperature of a culture of *‘monads*** (unicellular flagellates, including *Tetramitus rostratus*, *Monas dallingeri* and *Dallingeria drysdali*), inspecting the cultures for morphology (especially the presence of vacuoles), activity, fission and sexual fusion after each rise in temperature. Over the course of 7 years (the experiment ran from 1880 to 1886, allowing the cultures to acclimate after each small increment in temperature) Dallinger reported that the flagellates were still active and reproducing at 70 °C. This intriguing experiment needs repeating with replication, modern means of temperature control and documentation of growth from cell counts, as it may well establish a new thermal maximum for growth in unicellular eukaryotes.

**Eukaryotes at high temperature**

Eukaryotes appear not to be able to live at the very highest temperatures that characterize geothermal water. The highest *T_L* for a unicellular eukaryote appears to be 55–56 °C, which is the upper limit for the rhodophyte *Cyanidium caldarium*, although optimal (maximum) growth was at 45 °C (Doemel & Brock 1970, 1971). Over a century ago, however, Dallinger (1887) reported an experiment in which he raised the temperature of a culture of ‘monads’ (unicellular flagellates, including *Tetramitus rostratus*, *Monas dallingeri* and *Dallingeria drysdali*), inspecting the cultures for morphology (especially the presence of vacuoles), activity, fission and sexual fusion after each rise in temperature. Over the course of 7 years (the experiment ran from 1880 to 1886, allowing the cultures to acclimate after each small increment in temperature) Dallinger reported that the flagellates were still active and reproducing at 70 °C. This intriguing experiment needs repeating with replication, modern means of temperature control and documentation of growth from cell counts, as it may well establish a new thermal maximum for growth in unicellular eukaryotes.

Slightly higher temperatures appear to be tolerated by filamentous fungi, and a survey of a range of high-temperature habitats revealed species able to grow at 55–60 °C (Tansey & Brock 1972). Both unicellular algae and fungi can produce resting stages (spores) when environmental conditions become unfavourable. The upper thermal limit for survival of these spores is unclear, though resistance to 115 °C has been reported in spores of saprotrophic fungi in the Western Ghats of Southern India (Suryanarayanan *et al.* 2011). This extreme thermostolerance may be an adaptation to surviving the dry season wildfires that characterize the Western Ghats; the growing fungi themselves are mesophilic.

Higher plants can be found growing close to hot springs in Yellowstone, and in the perennial grass *Dichanthelium lanuginosum* (intriguingly named ‘hot springs panic grass’). In the hot springs panic grass the thermal tolerance is mediated through a mutualistic endophytic fungus *Curvularia protuberata* (Redman *et al.* 2002) and a mycovirus (Márquez *et al.* 2007). With both the fungus and mycovirus present, plants can grow in soils up to 65 °C; with either missing the plants are unable to grow above 38 °C (Márquez *et al.* 2007). Hot springs also provide the hottest habitats inhabited by invertebrates and vertebrates. Temperature in these springs may reach over 50 °C, and the fauna includes crustaceans, chironomid larvae, nematodes and molluscs, as well as fish. It is difficult to establish *T_L* values for these; although many secondary and anecdotal sources quote a range of temperatures for hot springs, there are very few primary sources with data for both temperature and fauna. Two nematodes, *Rhodtus terrestris* and *Udonchus tenacuatus*, appear to be ubiquitous in thermal springs and have been recorded as living up to 42.8 °C in Granada, Spain (Ocaña 1991), Wharton...
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(2002) records *Aphelenchoides parientus* living in hot springs up to 51 °C, and *Darylaimus thermus* is found in Yellowstone in waters up to 53 °C (Hoeppli 1926; Hoeppli & Chu 1932). Hot springs contain a range of other aquatic invertebrates, including crustaceans such as the isopod *Thermosphaeroma subequatum*, insect larvae (especially chironomids), and molluscs such as the springsnail *Tryonia julimensis*; all of these will have similar *T_l* values. The highest temperature for completion of the life cycle in an invertebrate may be for nematodes of the genus *Aphelenchoides* and *Panagrolaimus*, which tolerate temperatures of 60 °C in compost heaps (Steel et al. 2013).

Hot springs also support populations of fish, and the classic high-temperature fish are the desert pupfish of the genus *Cyprinodon*. These fish live in shallow geothermal springs, where the temperatures are high but vary both spatially and throughout the day and with season. *Cyprinodon pachycephalus* from the hot springs of San Diego de Alcalá, Chihuahua, México lives in waters of 39.2–43.8 °C (Minkley & Minkley 1986; Miller et al. 2005), and *Cyprinodon julimae* recently described from the hot springs of Julimes, Chihuahua, México lives at temperatures of between 38 and 46 °C (Montejano & Absalon 2009). In contrast to terrestrial vertebrates which can use shade to avoid the heat of the sun, and which cool off by night, desert pupfish spend their entire life at these high temperatures. While the water temperatures do vary a little diurnally and the fish often select the cooler water (Montejano & Absalon 2009), these two species of *Cyprinodon* are believed to be the fish with the highest *T_l* on Earth. They also limited to a few small springs and pools and are consequently highly endangered.

The hottest marine environments are hydrothermal vents, and these have a spectacularly rich and abundant fauna that includes a range of crustaceans, molluscs and worms (Van Dover 2000). As with the microbial flora within the vent chimneys, the very steep thermal gradients make it difficult to assess precisely what temperatures any given animal is experiencing. Behavioural observations and associated temperature measurements suggest that many motile vent animals select warm but not hot locations, and that they are very sensitive to changes in temperatures.

The most studied vent animal in this regard is the Pompeii worm, *Alvinella pompeiana*. This polychaete lives in a tube through which vent fluids pass, and from which it emerges to forage. Recordings with a temperature probe indicated that at the mouth of the tube temperatures averaged 22 °C (Cary et al. 1998). These data indicate that *Alvinella* is subject to a quite remarkable thermal gradient along its body (roughly 60 K). However, it is difficult to assess its *T_l*, both because of this gradient but also because the worm leaves its tube to forage in much cooler water (2 °C: Lutz 2012). Indeed studies of individual proteins indicate quite low thermal denaturation temperatures: ~ 50 °C for haemoglobin and ATP generation by isolated mitochondria, and ~ 45 °C for collagen. These data would indicate that in the long term, *Alvinella* is limited to temperatures below ~ 50 °C (Desbruyères & Laubier 1991). A recent study (Ravaux et al. 2013) has shown that long-term survival, as assessed by a 2 h ramped thermal exposure, is above 42 °C but below 50 °C. Similarly, another vent polychaete *Paralvinella sulfunctica*, can be found in waters up to 88 °C, but has an upper incipient lethal temperature (at which 50% of the population cannot survive indefinitely) of only 45 °C (Dilly et al. 2012). Although the actual *T_l* values for *Alvinella* or *Paralvinella* are unknown, current data suggest that they may hold the record *T_l* for an aquatic animal, and are also probably some of the most eurythermal metazoans on the planet (Lutz 2012).

Away from geothermal areas, the hottest terrestrial environments are deserts and directly measured surface temperatures in places can reach 75–80 °C (Ward & Seeley 1966). Deserts are, however, often only hot during the day; by night and under a clear sky temperatures can drop below freezing (Ward 2009).

This combination of high daytime temperatures and low night-time temperatures poses severe physiological problems for organisms living in deserts. Some motile forms are active by day and can tolerate brief periods of very high temperatures. For example, the Saharan silver ant *Cataglyphis bombycina* forages for very short periods in air temperatures up to 55 °C (Wehner et al. 1992). Similarly, *Oeomyrmex barbiger*, an ant from the Namib Desert, forages in air temperatures up to 67 °C (Marsh 1985), and the pseudoscorpion *Eremogarypus perfectus* only goes into heat coma at 65 °C (Heurtault & Vannier 1990). Being small, these ants have a very low thermal mass, and in consequence they heat up and cool down quickly. These, and other small arthropods active in the desert heat thus minimize the period of time for which they are exposed to the highest temperatures, and they climb frequently up stems of vegetation where the air is cooler.

Not all desert animals are so active and able to use behavioural means to minimize their exposure to the heat of the sun. Some such as land snails simply have to sit it out. The desert snail *Sphicterochila boissieri* minimizes its body temperatures by having a highly reflective shell, which allows it to maintain a tissue temperature of 50 °C in direct sunlight despite a local air temperature of 43 °C and a surface temperature of 65 °C (Schmidt-Nielsen et al. 1971). A significant physiological problem for organisms that cannot move underground to avoid temperature extremes is that gas exchange inevitably also results in water loss. Many arthropods minimize this water loss by restricting the periods of tracheal respiration (an adaptation referred to as discontinuous gas exchange). While these examples indicate that many desert invertebrates can tolerate very high temperatures, at least for short periods, they cool down by night and it is not clear what is the highest temperature at which any of these organisms can complete their life-cycles.

Desert plants must be able to withstand the same temperatures as the animals, but without the ability to move about to alleviate the direct effects of heat. One must assume that some plants living in the hottest deserts have thermal tolerances at least comparable with those living around
geothermal springs, but data on the maximum temperatures at which plants can complete their life cycle are very difficult to find.

Plants exposed to direct solar heating can reach temperatures well above that of the surrounding air. Fleshy leaves can reach internal temperatures of 40–50 °C when the surrounding air is only 20–30 °C, and the cambium of the sunny side of some trees may reach 55 °C (Öpik & Rolfe 2005). The record appears to be held by the cactus Opuntia, several species of which can reach internal temperatures up to 65 °C (Smith et al. 1984).

Current knowledge of the upper thermal limits to life is summarized in Table 4. Although there is a diverse literature on thermal limits to survival ($T_S$), data on the thermal limits to the completion of the life cycle ($T_L$) are far more difficult to obtain, and for some groups the data in Table 4 rely principally on the data collated over 40 years ago by Precht et al. (1973).

### What sets upper thermal limits?

The main physiological challenge from increased temperature comes in the form of greater thermal motion. In particular, the enhanced internal molecular motions increase the likelihood of thermal denaturation, in that the molecule may move to a conformation that is neither functional nor allowing return to a functional state.

The higher level structure of cellular macromolecules and the association between molecules critical to physiological processes are all dependent on weak bonds, such as Van der Walls interactions, hydrogen bonds, electrostatic (ionic) bonds and hydrophobic interactions. These all have low enthalpies of formation, typically in the range $<4$ kJ mol$^{-1}$ (Van der Waals) to 10–30 kJ mol$^{-1}$ (hydrogen bonds) and $<40$ kJ mol$^{-1}$ (hydrophobic interactions). These values are of the same order of magnitude as the mean thermal energy of molecules at 25 °C, and hence are likely to break more frequently at higher temperatures.

The most common adaptations to allow macromolecules to function at high temperatures include changes in the number of residues influencing overall hydrobicity, and an increase in the number of weak interactions to increase stability. These adaptations result in homologous enzymes having similar flexibility, and hence similar metabolic activity, at the optimal growth temperature (Tehei & Zaccai 2007). Whilst some proteins that are stable to ~140 °C can be isolated from hyperthermophiles, not all proteins appear to be sufficiently stable to retain their structure, and hence function, at very high temperatures (Lang 1986; Committee on the Origins and Evolution of Life 2007). In these, structural integrity is associated with the presence of large molecular chaperones and a range of small compatible solutes (Sterner & Liebl 2001).

The thermostability of nucleic acids has been linked to the content of guanine and cytosine, which, having three hydrogen bonds, are thermally more stable than the pairing of adenine and thymine, which have only two. However, there is no correlation between genome GC content and growth temperatures (Hurst & Merchant 2001; Hickey & Singer 2004). There is a correlation in structural RNAs although this is limited to the double-stranded stem regions of the molecules (Galtier & Lobry 1997).

Much attention has been directed at the unique membrane architecture of archaea in respect of the ability of some taxa to live at very high temperatures. Archaeal membranes consist of isoprenoid hydrocarbon chains, which are attached to the glycerol moiety by an ether link. Furthermore, the glycerol moiety has a different stereochemistry from that in bacteria and eukaryotes, and in some archaeans the isoprenoid chains are fused to form a single layer (as distinct from the bilayer that is found in all other organisms). While this unique membrane architecture is found in hyperthermophilic archaeans, it is also

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### Table 4. High-temperature limits for life on Earth

<table>
<thead>
<tr>
<th>Taxon</th>
<th>$T_L$ (°C)</th>
<th>$T_S$ (°C)</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Archaea</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>M. kandleri</td>
<td>122</td>
<td>&lt;130</td>
<td>Strain isolated from Kairei hydrothermal field, Central Indian Ridge (Takai et al. 2008)</td>
</tr>
<tr>
<td><strong>Bacteria</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>G. ferrireducens</td>
<td>100</td>
<td>nd</td>
<td>Isolated from Obsidian Pool, Yellowstone Park (Kashefi et al. 2002)</td>
</tr>
<tr>
<td><strong>Eukarya</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unicellular algae</td>
<td>60</td>
<td>nd</td>
<td>C. caldarium, isolated from acid hot springs in Yellowstone (Doemel &amp; Brock 1970, 1971)</td>
</tr>
<tr>
<td>Yeasts</td>
<td>60–62</td>
<td>nd</td>
<td>Filamentous fungi isolated from a range of geothermal sites in Yellowstone (Tansey &amp; Brock 1972)</td>
</tr>
<tr>
<td>Lichens</td>
<td>~ 45°</td>
<td>nd</td>
<td>Thermal tolerance depends on state of hydration (Precht et al. 1973); no recent data</td>
</tr>
<tr>
<td>Macroalgae</td>
<td>~ 45</td>
<td>nd</td>
<td>(Precht et al. 1973); no recent data</td>
</tr>
<tr>
<td>Mosses</td>
<td>~ 50°</td>
<td>nd</td>
<td>(Precht et al. 1973); no recent data</td>
</tr>
<tr>
<td>Angiosperms</td>
<td>65</td>
<td>nd</td>
<td>D. lanuginosum in Yellowstone National Park (Redman et al. 2002; Márquez et al. 2007)</td>
</tr>
<tr>
<td>Terrestrial invertebrates</td>
<td>~ 60</td>
<td>~ 70</td>
<td>Nematodes in compost heaps (Steel et al. 2013)</td>
</tr>
<tr>
<td>Freshwater invertebrates</td>
<td>~ 46</td>
<td>nd</td>
<td>Crustaceans and molluscs living alongside pupfish in hot springs</td>
</tr>
<tr>
<td>Marine invertebrates</td>
<td>&gt;42</td>
<td>~ 90</td>
<td>Polychaete Alvinella pompejana from hydrothermal vents (Cary et al. 1998; Ravaux et al. 2013)</td>
</tr>
<tr>
<td>Ectothermic vertebrates</td>
<td>~ 46</td>
<td>nd</td>
<td>Desert pupfish, Cyprinodon species (Miller et al. 2005; Montejano &amp; Absalón 2009)</td>
</tr>
<tr>
<td>Endothermic vertebrates</td>
<td>nd</td>
<td>nd</td>
<td>Cell temperatures in range ~30-45 °C</td>
</tr>
</tbody>
</table>

nd: no data.

* Unclear whether the entire life cycle is completed at this temperature.
found in archaea that live at low temperatures, which suggests that it may not necessarily be an adaptation to high temperature but simply a feature of archaeans in general. Its greater stability at high temperatures does, however, allow archaea to live at higher temperatures than bacteria or eukaryotes.

While an upper limit to the stability of biomolecules may be set by processes such as hydrolysis, it is not yet clear whether the upper limit for survival of an organism is set by the fate of individual molecules or the intracellular structure. Hansen et al. (2009) used calorimetry to investigate denaturation of key physiological macromolecules at high temperature and showed that in both *Escherichia coli* and *Lactobacillus plantarum* ribosomes denatured at lower temperatures than either DNA or cell walls.

We cannot necessarily generalize from these results and assume that upper thermal limits will be set by the same processes in all organisms. Indeed the difference in the upper thermal limits for Archaea, Bacteria and Eukarya (Table 4) suggests that different processes may set the limits for different domains. It is tempting to ascribe the markedly lower upper thermal limit for eukaryotes to the presence of a nucleus, but there are many other differences in internal cellular complexity that might be responsible for the greater sensitivity to high temperatures, and in truth we do not know the cause. In multicellular organisms, the requirement for integration between cells in tissues, and between tissues within the organism, introduces further possibilities for limiting thermal sensitivity. It also introduces the possibility of sublethal damage that limits physiological function, and hence maybe preventing completion of the life cycle, but without causing organism death.

**The physiological challenge of low temperature**

Cells at low temperatures must cope with the reduced molecular kinetic energy of the environment and the consequent lower rate of many physical processes. In addition, cells must adjust to reduced membrane fluidity, changes in intracellular pH and loss of macromolecular integrity (Hochachka & Somero 2002). The molecular adaptations to maintain physiological function at low temperatures are in many cases simply the reverse of those involved in adaptation to high temperatures.

Additional challenges arise, however, when ice is present in the environment immediately external to the cell. Following ice nucleation, solutes are rejected from the growing ice crystal and are concentrated in the remaining liquid, which thus increases in osmotic strength and tends to pull water from inside the cell.

A critical distinction in considering how organisms react to the physiological challenge of ice nucleation is that between unicells and multicellular organisms (Table 2). In archaea, bacteria and unicellular eukaryotes, the cell membrane is exposed directly to the environment. Although a cell wall may be present, this has to be permeable to allow the cell to take up nutrients, and dump waste products. This means that when ice is present in the environment, the cell membrane will experience directly the associated changes in osmotic strength. In contrast, multicellular organisms have their internal cells and tissues bathed in a fluid over whose composition the organism can exert a degree of control. When ice is present in the external environment, the cells and tissues of a multicellular organism such as a plant, insect or frog are not necessarily exposed to that ice.

When ice is present in the immediate environment of the cell, that cell may be damaged through hydraulic stresses, osmotic changes or solute toxicity mechanisms. In addition, intracellular ice can form when the cooling rate is sufficiently high that the cell cannot maintain osmotic equilibrium with the environment (Mazur 2004). Intracellular ice is observed only rarely in the natural environment (Wharton & Ferns 1995). This is partly because the formation of intracellular ice generally requires faster cooling than is typical of the natural environment. For example in the yeast *Saccharomyces cerevisiae* intracellular ice formation requires cooling rates faster than 20 K min$^{-1}$ (Seki et al. 2009). Atmospheric cooling rates in the environment rarely exceed 1 K min$^{-1}$ (Clarke et al. 2013), but some specialized habitats such as rock or leaf surfaces can change temperature more rapidly (Strimbeck et al. 1993). Freezing of extracellular fluids does, however, occur in some multicellular plants and animals living in seasonally cold climates (Schmid 1982; Leather et al. 1993; Pearce 2004).

At the slow cooling rates typical of the natural environment, cells can often maintain osmotic equilibrium with the surrounding fluid. Under these circumstances, free-living cells such as bacteria, archaeans or unicellular eukaryotes may vitrify. Vitrification (also known as the glass transition) occurs when a liquid begins to behave as a solid during cooling, but without any substantial change in molecular arrangement or thermodynamic state variables (pressure, volume, internal energy, entropy) (Wowk 2010). In bulk liquids, as temperature decreases, all molecular motions, translational and internal, become progressively slower until a critical temperature is reached where there is insufficient energy for significant translational molecular motion to take place over a meaningful timescale. This is the vitrification or glass transition temperature, $T_g$, and it is defined operationally as the temperature at which viscosity exceeds $10^{12}$ Pa.s (Debenedetti 1996).

Intracellular vitrification is more complex than in simple bulk liquids. This is principally because the interior of the cell is extremely crowded (Ellis 2001), approximating a colloid in physical structure. As colloids dehydrate they exhibit a sharp increase in viscosity and undergo a colloid glass transition (Zhou et al. 2009). Cellular dehydration, whether associated with a shift to anhydrobiosis or withdrawal of water driven by freeze concentration of the extracellular medium, can thus induce vitrification of the cell interior. The vitrification of the interior of a free-living microbial cell exposed to ice in the external environment is thus primarily the result of dehydration and is analogous to the vitrification of a colloid rather than the glass transition of bulk water.

The vitrification temperature of a cell will vary with the precise composition of the internal cell environment, and this
may allow the intracellular vitrification temperature to be adjusted by natural selection to match ecological circumstances (for example by varying the level of small cryoprotectant molecules). In free-living unicells, dehydration driven by freeze-concentration of the external environment triggers vitrification at temperatures between −10 and −25 °C (Clarke et al. 2013).

The very high viscosity of the vitrified cell means that movement of oxygen and metabolites effectively stops. Under these conditions, metabolism ceases. The cell does, however, maintain its internal integrity, and metabolism can start again once the cell warms and rehydrates. This process carries the danger that once water molecules can move within the cell, any small ice nuclei present may grow rapidly (a process usually termed devitrification) and the consequent mechanical damage may be lethal. The presence of chaperone proteins such as dehydrins or late embryogenesis abundant proteins (LEAs), as well as cryoprotectants such as polyols may be important in ensuring a safe transition from the vitrified to the normal state, though this is an area where more research is needed.

LEAs were originally identified in plant seeds, but are now known to be fairly widespread (Hand et al. 2011). It is also possible that thermal hysteresis proteins (antifreeze proteins) may be important in minimizing the chances of lethal growth of intracellular ice (devitrification) during the return to the fluid state.

**Low-temperature limits**

A general lower limit for life in free-living unicells would appear to be set by the temperature at which freeze-concentration of the external environment dehydrates the cell interior and drives vitrification (Clarke et al. 2013). While vitrification defines a lower limit for metabolism, it is possible that one or more other factor may limit completion of the life cycle at temperatures above which the cell vitrifies (that is $T_L$ is above $T_M$, where $T_M$ corresponds to the vitrification temperature).

A thorough survey of studies of microbial growth at low temperatures suggests that none grow below −20 °C (Fig. 4). There are a few reports in the literature of microbial growth at temperatures below −20 °C but in these studies growth or metabolism are usually inferred from proxies (such as release of apparent metabolic products) and not direct measures of cell number as a function of time. It is possible that in these cases geochemical processes or concentrated solution chemistry is mimicking metabolism (see discussion in Clarke et al. 2013). Indeed, despite a century of refrigeration technology, there are no reports of spoilage organisms growing below −20 °C (Geiges 1996). It is clear that the lower limit for completion of the life cycle in free-living unicells (bacteria, archaeans, unicellular eukaryotes) is around −20 °C.

The lower thermal limit for life in marine organisms is set by the lowest temperature at which seawater can remain liquid. For seawater of normal salinity (35), the equilibrium freezing point at atmospheric pressure is −1.92 °C, increasing with depth. Parts of the continental shelf seabed around Antarctica are close to the freezing point year-round (Clarke et al. 2009) and so the rich communities of marine invertebrates and fish that live there must complete their life cycles at this temperature ($T_L \approx -2^\circ C$).

Teleost fish have body fluids of lower osmotic strength than seawater, and would freeze at these temperatures. Polar teleosts avoid such freezing by the production of protein or glycoprotein antifreezes, together with a suite of associated anatomical and physiological adjustments (DeVries & Cheng 1992). Different lineages of teleost fish use different molecules as antifreezes, indicating that antifreeze has evolved many times independently, suggesting that this is not a difficult evolutionary problem.

When sea-ice forms, salt is excluded from the growing ice. As a result sea-ice contains many channels filled with brine, and these can reach very low temperatures ($\sim -20^\circ C$) without freezing. These channels contain diatoms and many other unicellular eukaryotes (Thomas 2012), as well as some invertebrates. It is possible that these assemblages contain taxa that can complete their life cycle at lower temperatures than in the surrounding seawater: for example the Arctic sea-ice diatom *Nitzschia frigida* can grow down to −8 °C, although the doubling time is very long (60 days) (Thomas 2012).

In terrestrial habitats, temperatures may fall very low in winter (see above). While some animals migrate to warmer climates, many others and all plants have to survive the winter *in situ*. Polar organisms have evolved two basic strategies to achieve this: they either prevent ice forming within their tissues and survive the winter in an undercooled state (*freeze avoidance*), or they allow extracellular water to freeze (*freeze tolerance*). In the latter case, cells are typically subject to
dehydration driven by freeze-concentration of the extracellular water as it freezes; they then either undercool or vitrify.

It has long been known that fungal infections can affect crops under snow and Schmidt-Nielsen (1902) reported the growth of two unicellular yeasts at 0 °C. Pathogenic fungi are known from snow banks and also from cryoconite assemblages on the surface of glaciers, and mushrooms that emerge through snow in northern forests are well known (Hoshino et al. 2013). The lower thermal limit for completion of the life cycle in free-living fungi and yeasts remains unknown but appears to be about 0 °C, and while some moulds and rusts can grow at low water activities on refrigerated foods, none are known to grow below −20 °C (Geiges 1996).

The most studied animals in terms of winter survival in relation to freezing temperatures are arthropods, and insects show examples of both freeze avoidance and freeze tolerance (Leather et al. 1993; Wharton 2002; Denlinger & Lee 2010). A recent study of thermal limits (T_S) in insects shows clearly that T_S is much lower in insects from higher (colder) latitudes (Fig. 5), indicating that T_S can be adjusted by natural selection to match ecological requirements. A few vertebrates have been shown to tolerate extracellular freezing, and the most studied example here is the wood frog Rana sylvatica (Schmid 1982; Storey & Storey 1996, 2004). Plants overwintering in cold regions are known to undergo extracellular freezing, and in some cases vitrification of the dehydrated cells has been demonstrated (Hirsh et al. 1985). This may prove to be a widespread phenomenon in high-latitude plants (Hirsh 1987; Pearce 2004).

Lichens are among the most tolerant of plants, being found in habitats where higher plants are missing. They are able to dehydrate extensively, and in this state can tolerate very low temperatures. The lowest recorded temperature for photosynthetic carbon fixation is −24 °C (Kappen 1993). In Antarctic lichens, photosynthesis has been recorded down to −16.5 °C in Xanthoria candelaria, −17 °C in Umbilicaria aprina and −18 °C in Neupogon acromelenus (Kappen 1993; Schroeter et al. 1994). In U. aprina, dark respiration ceased at higher temperatures, suggesting that T_L is −10 °C (Schroeter et al. 1994). Lichens are a symbiotic relationship between an alga (the photobiont) and a fungus (the mycobiont). Interestingly, the symbiotic organisms (the lichen) appear to withstand more extreme conditions than either the phytobiont or mycobiont alone (de Vera et al. 2008).

In Antarctica, lichens extend further south than either mosses or vascular plants (Peat et al. 2007), suggesting that they are more tolerant of very low temperatures and the associated aridity. These characteristics of lichens have stimulated considerable interest in their use as model experimental organisms in astrobiology (de Vera et al. 2008; Onofri et al. 2012). Indeed, lichens have been shown to photosynthesize under simulated Martian conditions (de Vera et al. 2010), and it has been suggested that lichens could even by viable in small areas with suitable microclimates on Mars today. Studies have, however, concentrated on photosynthesis, and the limited experimental; data suggest that growth ceases at higher temperatures than photosynthesis (Schroeter et al. 1994).

In most cases, terrestrial organisms able to survive extreme cold can only complete their life cycle once temperatures have risen again in summer. Hence, while some very impressive examples of low-temperature survival (T_S) are known, it is much more difficult to assign a value to the low-temperature threshold for completion of the life cycle (T_L). The lowest limits so far documented are for invertebrates in meltwater on glaciers, such as the enchytraeid annelid ‘ice worms’ of the genus Mesenchytraeus (Farrell et al. 2004) or chironomid midges of the genus Diamesa (Kohshima 1984; Hagvar 2010), where T_L is −0 °C. This is slightly higher than the T_L for marine invertebrates, but it may be that we simply lack documentary evidence of lower T_L values for terrestrial plants and invertebrates. The marked diurnal and seasonal variations in environmental temperature will, however, make such data difficult to obtain.
Andrew Clarke

The special case of endotherms

Two lineages of vertebrates, mammals and birds, have independently evolved endothermy, the capacity to maintain a high and constant body temperature. This is often referred to colloquially as ‘warm-bloodedness’, but this term is unhelpful because many ectotherms, including insects, reptiles and even some fish, can achieve similarly high body temperatures.

Typically endotherm body temperatures are in the range 30–45 °C, with birds tending to be warmer than mammals (Clarke & Rothery 2008). These temperatures are achieved through a combination of a high resting metabolic rate and insulation. Endothermy is very expensive energetically, but it allows mammals and birds to be active at any time of the day or night, and to occupy a very wide range of habitats.

While endotherm cell temperatures are confined to a narrow range, the environments they inhabit range from the hottest deserts to the polar regions. To take mammals as an example, the range of body temperatures is 30–41 °C, but these live in areas with annual mean temperatures ranging from −11 to 26 °C (Fig. 6). The extreme endotherm example is, however, probably a bird: the Emperor Penguin, *Aptenodytes forsteri*, which raises its chick on sea-ice in the depths of the Antarctic winter, when temperatures are −20 °C or below (McCafferty et al. 2013).

This poses the question of what is the correct value of $T_L$ for an endotherm: is it the internal body temperature at which the cellular physiology operates, or is it the environmental temperature? For humans with cultural adaptations (clothing, housing), the range of environmental temperatures that define the $T_L$ range is the widest of any species on Earth.

Concluding remarks

The upper and lower thermal limit to life vary markedly across the domains of life on Earth (Tables 4 and 5). While data on limits to survival ($T_S$) attract considerable attention, the thermal limits to completion of the life cycle ($T_L$) are far more difficult to determine and hence are much less well defined.

The data are clearest for unicells. Currently identified upper thermal limits for growth are 122 °C for archaeans, 100 °C for bacteria and ~60 °C for unicellular eukaryotes. No unicells appear to grow below −20 °C, a limit that is probably set by dehydration-linked vitrification of the cell interior driven by freeze-concentration in the presence of extracellular ice (Clarke et al. 2013).

The range of temperatures over which multicellular eukaryotes can complete their life cycle is much narrower than for unicells. The upper limits on land would appear to be exhibited by nematodes in hot springs and compost heaps (~60 °C), and in the sea by polychaetes associated with hydrothermal vents (>40 °C, but poorly defined). The upper limit for an ectothermic vertebrate would appear to be 38–46 °C for desert pupfish. The lower thermal limits for survival ($T_S$) in multicellular organisms extend to at least −70 °C, the winter minimum temperature of the Arctic tundra and taiga. However in all cases known to date, completion of the life cycle requires summer warmth and the lowest $T_L$ appears to be ~0 °C for invertebrates in glacial meltwater and ~−2 °C for marine invertebrates and fish living on the continental shelves around Antarctica.

It is interesting that in lichens (de Vera et al. 2008) and the grass *D. lanuginosum* (Redman et al. 2002), temperature tolerance is conferred by symbiosis or mutualism. The mechanism by which the symbiotic organism gains enhanced temperature tolerance in comparison with the isolated individual components remains obscure.

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