Blue blood on ice: modulated blood oxygen transport facilitates cold compensation and eurythermy in an Antarctic octopod

Oellermann et al.
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Abstract

Introduction: The Antarctic Ocean hosts a rich and diverse fauna despite inhospitable temperatures close to freezing, which require specialist adaptations to sustain animal activity and various underlying body functions. While oxygen transport has been suggested to be key in setting thermal tolerance in warmer climates, this constraint is relaxed in Antarctic fishes and crustaceans, due to high levels of dissolved oxygen. Less is known about how other Antarctic ectotherms cope with temperatures near zero, particularly the more active invertebrates like the abundant octopods. A continued reliance on the highly specialised blood oxygen transport system of cephalopods may concur with functional constraints at cold temperatures. We therefore analysed the octopod’s central oxygen transport component, the blue blood pigment haemocyanin, to unravel strategies that sustain oxygen supply at cold temperatures.

Results: To identify adaptive compensation of blood oxygen transport in octopods from different climatic regions, we compared haemocyanin oxygen binding properties, oxygen carrying capacities as well as haemolymph protein and ion composition between the Antarctic octopod Pareledone charcoti, the South-east Australian Octopus pallidus and the Mediterranean Eledone moschata. In the Antarctic Pareledone charcoti at 0°C, oxygen unloading by haemocyanin was poor but supported by high levels of dissolved oxygen. However, lower oxygen affinity and higher oxygen carrying capacity compared to warm water octopods, still enabled significant contribution of haemocyanin to oxygen transport at 0°C. At warmer temperatures, haemocyanin of Pareledone charcoti releases most of the bound oxygen, supporting oxygen supply at 10°C. In warm water octopods, increasing oxygen affinities reduce the ability to release oxygen from haemocyanin at colder temperatures. Though, unlike Eledone moschata, Octopus pallidus attenuated this increase below 15°C.

Conclusions: Adjustments of haemocyanin physiological function and haemocyanin concentrations but also high dissolved oxygen concentrations support oxygen supply in the Antarctic octopus Pareledone charcoti at near freezing temperatures. Increased oxygen supply by haemocyanin at warmer temperatures supports extended warm tolerance and thus eurythermy of Pareledone charcoti. Limited haemocyanin function towards colder temperatures in Antarctic and warm water octopods highlights the general role of haemocyanin oxygen transport in constraining cold tolerance in octopods.

Keywords: Haemocyanin, Hemocyanin, Cephalopod, Oxygen affinity, Oxygen carrying capacity, Diffusion chamber, Pareledone charcoti, Octopus pallidus, Eledone moschata
**Introduction**

The Antarctic Ocean forms an extreme habitat with temperatures ranging between -1.8 to 2°C all year round e.g. [1,2]. Most marine animals living under these conditions are unable to regulate their body temperature (ectotherms) and are thus required to sustain body functions at near freezing temperatures, via numerous adjustments at the molecular, cellular or systemic level [3]. On the other hand, Antarctic waters are rich in oxygen due to increased solubility of oxygen and rigorous mixing across the water column [4]. Paired with low metabolic rates, commonly found among Antarctic ectotherms [5-7], oxygen supply seems less challenging in the cold, as demonstrated by the ability of Antarctic notothenioid fishes to sustain life with low levels of haemoglobin [8] and in case of the Antarctic icefishes (Channichthyidae), even with the complete absence of oxygen transport proteins in the cold [4,9]. Conversely, cold temperatures may hamper oxygen supply by lowered diffusion across tissue and cellular boundaries, increased viscosity [10] and often a decreased ability of blood pigments like vertebrate haemoglobin or cephalopod haemocyanin to release oxygen to tissues as the pigment’s affinity for oxygen increases [11-13]. Antarctic fishes cope with these challenges by increased mitochondrial and membrane densities supporting diffusion [10], loss of blood cells reducing blood viscosity [14] or lowered oxygen affinity sustaining oxygen transport by their haemoglobin [15-17]. Little is known whether Antarctic ectotherms other than fish evolved comparable physiological adaptations to sustain oxygen supply in the cold.

Among those other ectotherms are numerous species of Antarctic octopods, which occur exclusively in the Antarctic Ocean and form an important part of the benthic megafauna as both prey and predators [18-22]. Although their origin is still unclear, Antarctic octopods may have evolved in situ in shallow Southern Ocean waters [23], or colonised Antarctic shelves from the deep-sea [24] or possibly in the case of the genera *Pareledone* or *Megaladeledone* from shallow South-American waters prior to the cooling of Antarctica and the associated opening of sea passages between 29-32 million years ago (25,26, Figure 1). Irrespective of their origin, to become successful members of the Antarctic fauna as they are today, octopods were eventually required to adjust to temperatures as low as -1.9°C.

Survival at such cold temperatures is supported by physiological adjustments that sustain metabolism and motor activity [29,30]. Unlike fishes, which are hyposmotic to seawater [31], octopods do not need to fear freezing, as their body fluids are nearly isosmotic to seawater [32] and freeze at about the same temperature of -1.9°C. A major challenge, however, may involve retaining the functionality of the advanced oxygen supply system of coleoid cephalopods. Their closed circulatory system comprises three hearts and contractile veins that pump haemolymph, which is highly enriched with the blue coloured oxygen transport protein haemocyanin (89 mg ml^-1^ in *Megaladeledone setebos* [33] or up to >160 mg ml^-1^ in *Loligo vulgaris* [12,34,35]), at blood pressures which are high for invertebrates (e.g. *Enteroctopus dofleini* 5.3-9.3 kPa, [36-38]). Evidence suggests that circulatory support by ventilatory pressure oscillations as well as heart performance may fail at high temperatures and decrease oxygen supply in cephalopods [39,40]. At low temperatures, haemocyanin may cause systemic oxygen shortage due to its decreasing ability to release sufficient oxygen to tissues [33,41].

Low temperatures decrease rate dependent biochemical and metabolic processes [42]. Antarctic octopods may compensate for this by fully or partially reversing such effects [43]. To date only few studies have investigated cold compensated features in Antarctic octopods. Garrett and Rosenthal [44] reported accelerated kinetics of potassium channels to enhance nervous signal transduction in the Antarctic octopus *Pareledone sp.*. Daly and Peck [7] observed that Antarctic temperatures lower oxygen consumption rates of *Pareledone charcoti* as predicted from the temperature sensitivity of metabolism of the temperate octopus *Eledone cirrhosa*. Consequently, oxygen consumption rates were considered low and uncompensated in *Pareledone charcoti*. This is even more apparent if one scales oxygen consumption rates to the relatively small size of *Pareledone charcoti*. That is, according to the scaling function of mass-specific oxygen consumption rates for octopods (*M*<sub>O2</sub> = 3.35 *M*<sup>-0.27</sup> [45]) an octopus weighing 51 g is expected to consume 1.093 mmol O<sub>2</sub> kg<sup>-1</sup> h<sup>-1</sup> at 0°C (assuming a Q<sub>10</sub> of 2.12 [46-48]) but *Pareledone charcoti* instead only consumes 0.362 mmol O<sub>2</sub> kg<sup>-1</sup> h<sup>-1</sup> at 0°C [7]. Furthermore, Zielinski et al. [33] studied haemocyanin oxygen binding in the large *Megaladeledone setebos* (former *Megaladeledone senoi*, [49]) and observed haemocyanin oxygen affinity to be high and irresponsive to temperature, implying poor oxygen unloading and very limited temperature tolerance. However, comparisons of these features with those in octopods from warmer climates are required. It therefore remains unclear whether oxygen supply in Antarctic octopods features adjustments to the cold or simply lacks compensation. It further remains open whether the findings in *Megaladeledone setebos* also apply to the much smaller and more common Antarctic octopods of the genus *Pareledone*, and to what extent oxygen supply via haemocyanin differs between the cold water species and octopods that face much higher and more variable temperatures.

Therefore, in this study, we aimed to assess

1. Whether oxygen transport via haemocyanin features modifications that facilitate oxygen supply and thus survival of Antarctic octopods at close to freezing temperatures.
2. Whether oxygen transport properties and related stenothermy reported for Megaleledone setebos also occur in other Antarctic octopods.

3. Whether octopods adapted to warmer and broader temperature windows employ diverging strategies to sustain haemocyanin mediated oxygen supply across various temperatures.

To address these objectives, we compared oxygen binding properties, total oxygen carrying capacities as well as protein and ion composition of haemolymph of the abundant Antarctic octopod species Pareledone charcoti with two octopod species originating from warmer climates, the South-east Australian Octopus pallidus and the Mediterranean Eledone moschata.

Here we report specific properties of oxygen transport in the Antarctic octopod Pareledone charcoti, which include reduced oxygen affinities and high oxygen carrying capacities, but also a high, thermally sensitive venous reserve that supports eurythermy in Pareledone charcoti. We emphasize the general role of haemocyanin in shaping cold tolerance in both cold- and warm-water octopods.

**Results**

**Temperature dependent oxygen binding in vitro**

*In vitro* changes in oxygen binding by the respiratory pigment haemocyanin were assessed by pH oxygen-saturation analysis (see Methods). At a common temperature of 10°C the haemocyanin of the Antarctic octopod Pareledone charcoti displayed a lower affinity for oxygen than haemocyanin of the South-east Australian Octopus pallidus and the Mediterranean Eledone moschata, reflected in a 1.4- or 4.2-fold higher $P_{50}$ ($PO_2$ at which haemocyanin reaches half-maximum saturation with oxygen (50%)), respectively (Figure 2A, Table 1). Further, at 10°C, the cooperativity of oxygenation dependent proton binding (i.e. expressed as the pH-dependent release of oxygen by haemocyanin and derived from the oxygen carrying capacity and the maximum slope of the pH oxygen-saturation curve, $\Delta O_2$ mmol L$^{-1}$ ΔpH) was highest in Pareledone charcoti (Kruskal Wallis, $\chi^2 (2) = 28.0$, $P < 0.001$) compared to Eledone moschata (Mann–Whitney, $P < 0.001$) and Octopus pallidus (Mann–Whitney, $P < 0.001$, Table 1).

Temperature changes affected oxygen binding in all three octopod species, indicated by increased oxygen affinities and diminished cooperativity of oxygenation dependent proton binding towards colder temperatures (Table 1, Figures 3 and 4). In Pareledone charcoti and Eledone moschata, oxygen affinities increased more steadily, in Octopus pallidus however, oxygen affinities remained nearly unchanged between 10-15°C but decreased considerably above 15°C (Table 1, Figure 4). According to the changes in oxygen affinity, oxygen saturation decreased with increasing temperatures (Figure 4A, Note that calculations of oxygen saturation were based on arterial and venous PO$_2$ and an arterial-venous pH difference determined for Octopus vulgaris [50,51] and were assumed to be constant across temperature). However, this drop...
mostly occurred in the range of low PO₂ between 4 and 1 kPa (Figure 4A). At a PO₂ of 13 kPa, oxygen saturation remained virtually unchanged in *Pareledone charcoti* and *Eledone moschata* and decreased only slightly but significantly by 9.7% in *Octopus pallidus* above 15°C (ANOVA, F₁, 15 = 5.40, P = 0.017, Figure 4A). pH sensitivity of oxygen affinity expressed as the Bohr coefficient (Δlog PO₂/ΔpH) was not significantly affected by experimental temperatures (ANOVA, F₁, 16 = 0.36, P = 0.555). Among species the lowest Bohr coefficients were found in *Pareledone charcoti* (Table 1). The cooperativity of oxygenation dependent proton binding decreased significantly towards cooler temperatures in all three species (ANOVA, F₁, 147 = 114.41, P < 0.001, Table 1, Figure 3).

The analysis of inorganic cations in the haemolymph showed no differences between *Pareledone charcoti*, *Octopus pallidus* and *Eledone moschata* (Table 1). Interestingly, haemocyanin content did not co-vary with or equal total haemolymph protein but differed significantly between species (ANOVA, F₁, 29 = 8.98, P < 0.001, Figure 5).

The highest concentrations of haemocyanin were found in the Antarctic octopod *Pareledone charcoti* (78.9 mg ml⁻¹, 95% confidence interval (CI) from 69.2-88.6 mg ml⁻¹, Figure 5).

**Implications for blood oxygen transport in vivo**

In this section, the *in vitro* results are described in terms of their implications for the putative *in vivo* patterns of oxygen binding. At 0°C haemocyanin of *Pareledone charcoti* would release only 16.3% of its bound oxygen assuming an arterial-venous transition from 13 to 1 kPa PO₂ and pH 7.53-7.42 (Figure 3A, Figure 4). Even at low pH (<6.4) and low oxygen tensions (1kPa PO₂), 33.6% (28.4-38.8) of the oxygen would remain bound to the Antarctic haemocyanin. For comparison, within the range of their habitat temperature from 10 to 20°C, haemocyanins of *Octopus pallidus* and *Eledone moschata* would release between 33.0-60.0% and 29.8-70.0% oxygen, respectively (Figure 3D-I, Figure 4).

Haemocyanin of *Pareledone charcoti* showed the lowest venous oxygen saturation at a common temperature of 10°C, at a venous PO₂ of 1 kPa and a venous pH of 7.27 (Figure 2B). At 10°C the Antarctic haemocyanin thus has the potential to release far more oxygen (on average 76.7%, 95% CI 68.6% to 84.8%) upon each cycle than the warm-water octopods *Octopus pallidus* (33.0%, 5.0-60.9) and *Eledone moschata* (29.8%, 9.9-49.7, Figure 3C, D, G, Figure 4A). This is mostly due to an increased pH dependent release of oxygen by haemocyanin in *Pareledone charcoti* (Figure 3C, D, G), with maxima occurring 0.16 or 0.25 pH values above those of *Octopus pallidus* or *Eledone moschata* respectively (Figure 6).

Surprisingly, the Antarctic *Pareledone charcoti* has a larger capacity to carry oxygen in its haemolymph than *Octopus pallidus* or *Eledone moschata* (ANOVA, F₁, 31 = 12.57, p < 0.001, Table 2), due to the highest haemocyanin content of all three species (Figure 5). This increased capacity for oxygen transport in *Pareledone charcoti* is further enhanced by high levels of dissolved oxygen at 0°C (359.5 μmol L⁻¹, 35 psu (practical salinity units, Figure 4B) accounting for 18.5% of the total haemolymph oxygen content and up to 42% of the oxygen released to the tissue in *Pareledone charcoti* (assuming an arterial-venous transition from 13–1 kPa PO₂ and pH 7.53-7.42). The contribution of dissolved oxygen is also significant in the warm-water octopods *Octopus pallidus* and *Eledone moschata*, within the range of their habitat temperatures between 10-20°C, amounting to between 17-20% or 18-21%, respectively, of total haemolymph oxygen content and 30-16% or 34-15%, respectively, of the oxygen eventually released to tissues (assuming an arterial-venous transition from 13–1 kPa PO₂, Figure 4B).

**Discussion**

Comparing the haemocyanins of the Antarctic octopod *Pareledone charcoti* with those of the warmer-water
Table 1 Comparison of oxygen binding parameters and cation composition of haemolymph between *Pareledone charcoti*, *Octopus pallidus* and *Eledone moschata*

| Temperature (°C) | $P_{50}$ (kPa) | $\Delta$mmol l$^{-1}$O$_2$/ ΔpH | Bohr coefficient* (pH range 6.7-7.5) | $\Delta P_{50}$/°C | Cation concentration (mmol L$^{-1}$) | Mg$^{2+}$ | Na$^+$ | K$^+$ | NH$_4^+$ | Ca$^{2+}$ |
|-----------------|---------------|-------------------------------|-------------------------------------|-------------------|-------------------------------------|---------|-------|-------|---------|---------|
| **Pareledone charcoti** |              |                               |                                     |                   |                                     |         |       |       |         |         |
| 0               | 0.41          | 1.17                          | (NA, pH 7.42)                       | 0.088-1.46        |                                     |         |       |       |         |         |
| 5               | 0.7           | 2.39                          | (NA, pH 7.34)                       | 0.47-4.32         | (-1.67 - -0.78)                    | 0.15    | 52.6  | 385.5 | 8.5     | 6.1     | 88      |
| 10              | 2.34          | 3.17                          | (2.19-2.49, pH 7.27)                | (2.51-3.82)       |                                     |         |       |       |         |         |         |
| **Octopus pallidus** |              |                               |                                     |                   |                                     |         |       |       |         |         |
| 10              | 1.63          | 1.40                          | (0.99-2.26, pH 7.27)                | (1.26-1.55)       |                                     |         |       |       |         |         |         |
| 15              | 1.81          | 2.55                          | (1.46-2.16, pH 7.19)                | (2.24-2.85)       | (-2.50 - -1.44)                    | 0.39    | 45.6  | 372.0 | 10.0    | 8.6     | 7.5     |
| 20              | 6.07          | 2.94                          | (3.74-8.40, pH 7.11)                | (2.63-3.25)       |                                     |         |       |       |         |         |         |
| **Eledone moschata** |              |                               |                                     |                   |                                     |         |       |       |         |         |
| 10              | 0.56          | 1.13                          | (0.1-1.02 pH 7.27)                  | (0.77-1.48)       |                                     |         |       |       |         |         |         |
| 15              | 1.49          | 1.79                          | (0.70-2.89)                         | (-2.22 - -1.55)   | (0.11 – 0.30)                     | 0.21    | 49.9  | 386.8 | 8.8     | 3.0     | 7.3     |
| 20              | 2.62          | 2.08                          | (2.00-3.24, pH 7.11)                | (1.79-2.37)       |                                     |         |       |       |         |         |         |
| **ANOVA**†      | $F_{2, 149} = 10.82$ | $F_{2, 16} = 3.71$ | $F_{2, 17} = 1.58$ | $F_{2, 17} = 0.78$ | $F_{2, 17} = 0.31$ | $F_{2, 17} = 0.85$ | $F_{2, 17} = 1.35$ | $F_{2, 17} = 2.28$ | $P < 0.001$ | $P = 0.048$ | $P = 0.271$ | $P = 0.475$ | $P = 0.739$ | $P = 0.446$ | $P = 0.285$ | $P = 0.133$ |

Numbers in brackets indicate the range of 95% confidence intervals.

*Data at different temperature were pooled for each species.

*Based on alpha-stat shifted venous pH and a 10°C temperature interval.

*ANOVA results for between species comparison.
octopods *Octopus pallidus* and *Eledone moschata* reveals differences and properties of the respiratory pigment that assist oxygen supply at close to freezing temperatures but also support an extended range of oxygen dependent thermal tolerance in the Antarctic species. Haemocyanin functional properties in *Eledone moschata* constrain oxygen supply by haemocyanin at its lower temperature margin of 10°C. In *Octopus pallidus*, however, oxygen affinities decrease strongly above 15°C but stabilise at 10°C, suggesting a dual strategy to improve oxygen supply at both its upper and lower temperature margins.

**Blood oxygen transport in the cold**

Due to the exothermic binding of oxygen in cephalopod haemocyanins, oxygen affinity increases towards colder temperatures and may severely hamper oxygen release to tissues at the sub-zero temperatures [58] prevailing in the Antarctic Ocean. Our results show that *Pareledone charcoti* attenuates this detrimental effect by means of lowered oxygen affinity of the haemocyanin (Figure 2A, Table 1). Such lowered, cold-compensated oxygen affinities are not unique to *Pareledone charcoti* and the respiratory pigment haemocyanin, but were also observed in red-blooded Antarctic fishes such as *Dissostichus mawsoni* (*$P_{50}$* of 1.93 kPa at pH 8.16 and −1.9°C, [17]) or *Pagothenia borchgrevinki* (2.8 kPa at pH 8.1 and −1.5°C), whose oxygen affinities were much lower than those of temperate fish extrapolated to the same temperatures [59].

Allosteric effectors (e.g. ATP) may strongly contribute to decreased oxygen affinities of the haemoglobins of Antarctic fishes [17]. *Pareledone charcoti* however, relies on modifying the intrinsic properties and the pH sensitivity of its haemocyanin. The only known allosteric effectors in octopod haemolymph, inorganic ions, particularly magnesium [60,61], are not regulated and found at levels similar to those in sea water (54.2 mmol L$^{-1}$ at 35 psu, [62]) and similar to those in haemolymph of other octopods (*Octopus pallidus*, *Eledone moschata*, Table 1; *Eledone cirrhosa*, 54.6 mmol L$^{-1}$ [32]). This confirms that cephalopods do not regulate haemolymph magnesium concentrations to modulate oxygen binding. Instead, *Pareledone charcoti* increases oxygen release via a higher pH dependent release of oxygen by haemocyanin and by a pH sensitive range of oxygen binding located at higher pH values than seen in the warm water octopods (Figure 3C, D, G, Figure 6), which aligns with the cold-induced alpha-stat shift of venous pH (Figure 7).
Compensation for incomplete oxygen release

Although Pareledone charcoti has experienced a decrease in oxygen affinity of its haemocyanin oxygen release is far from being complete at 0°C as more than 77% of the oxygen remains bound to haemocyanin (assuming an arterial-venous transition from 13 to 1 kPa $P_{O_2}$ and pH 7.53-7.42, Figures 3A and 4A). The major factors contributing to the projected incomplete oxygen unloading at 0°C are the i) cold-induced increase of affinity of haemocyanin for oxygen, ii) reduced pH dependent release of oxygen by haemocyanin and iii) alpha-stat shift of haemolymph pH towards higher pH (Figures 6 and 7). In fact, increased oxygen affinity and reduced release of oxygen at colder temperatures are consistently reported for octopods ([12,13] for review) and assumed to be due to a more rigid structure of the haemocyanin molecule [61]. The alpha-stat pattern of haemolymph pH changes observed for octopods (Figure 7) has also been reported for squids [63], suggesting that temperature dependent changes of haemolymph pH affect oxygen supply in most if not all cephalopods. Melzner et al. [39] illustrated that the interplay of these factors lead to a venous oxygen release of less than 10% in Sepia officinalis at 10°C and 1.7 kPa $P_{O_2}$, and accordingly, to only ~22% or ~5% oxygen release at 0°C and 1.0 or 1.7 kPa $P_{O_2}$, respectively, in the Antarctic octopod Megaleledone setebos [33]. Although $P_{O_2}$ values below 1 kPa may further improve oxygen unloading, it is questionable whether the remaining oxygen gradient to mitochondria would be steep enough to maintain oxygen flux [39]. Therefore, poor oxygen unloading in Pareledone charcoti at 0°C due to high oxygen affinity, lowered pH dependent release of oxygen by haemocyanin and high venous pH are well in line with previous notions describing these factors to be crucial in defining limits of oxygen supply in the cold [13,39,58].

Most surprisingly, Pareledone charcoti compensates for poor oxygen unloading by considerably increasing haemocyanin concentrations. It thereby carries 40% or 46% more haemocyanin-bound oxygen in its haemolymph than Octopus pallidus or Eledone moschata, respectively (Figure 5). Overall, oxygen carrying capacities of the Antarctic octopods Pareledone charcoti and Megaleledone setebos rank among the highest reported for octopods and resemble those of red-blooded Antarctic fishes (Table 2). This and the presence of deeply blue-colored haemolymph in many other Antarctic octopods (Adelieledone polymorpha, Pareledone spp., Benthoctopus sp., M. Oellermann, pers. obs.) not only underlines the dependence of Antarctic octopods
on high haemocyanin concentrations but also contrasts the general finding of reduced erythrocyte and blood pigment concentrations in red-blooded Antarctic fishes [8] or Antarctic crustaceans [64]. It appears that red-blooded Antarctic fishes depend less on their oxygen transport protein than Antarctic octopods, despite higher rates of oxygen consumption (e.g. *Trematomus hansoni* 22.4 mg O$_2$ kg$^{-1}$ (wet mass) h$^{-1}$ [65] vs. *Pareledone charcoti* 10.2 mg O$_2$ kg$^{-1}$ (wet mass) h$^{-1}$ [7]). This may reflect a lower degree of capillarisation in the cephalopods [66] or the lower oxygen binding capacity of haemocyanin compared to fish haemoglobin [67]. However, we can presently not exclude that high haemocyanin protein concentrations serve other cold compensated processes as well.

The reduction of haemoglobin content in red-blooded Antarctic fishes has been interpreted to balance the increase in blood viscosity at low temperatures [68,69]. One therefore wonders why Antarctic octopods evolved to maximize the concentration of an extracellular protein, which increases viscosity even further? This may be best explained by either one or all of the following reasons, i) an increase in the fraction of haemocyanin in extracellular protein without causing higher levels of haemolymph proteins (Figure 5), ii) the non-existence of anti-freeze proteins that can largely contribute to blood protein levels in Antarctic fishes (e.g. 32 mg ml$^{-1}$ or ~35% of total blood protein concentration in *Dissostichus mawsoni*, [8,31]) and increase blood viscosity [70] and iii) haemocyanin concentrations below viscosity limits. Squids living in temperate and subtropical waters were reported to have haemocyanin in excess of 160 mg ml$^{-1}$ (*Loligo vulgaris* and *Loligo pealei*, [12,61,71]), whereas maximum haemocyanin levels of *Pareledone charcoti* seen in the present study were 106.8 mg ml$^{-1}$. However, maximum tolerated haemocyanin levels may be far lower at 0°C due to increasing blood viscosity towards colder temperature [68]. We conclude that as a trade-off, increased haemocyanin concentrations occur at the expense of elevated viscosity. The ability to maximize haemocyanin levels at sub-zero temperatures

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**Figure 5** Total protein and haemocyanin concentrations in haemolymph of cold and warm water octopods. Haemocyanin concentrations were calculated from the haemolymph oxygen carrying capacity, based on a molecular weight of 3.5 MDa and 70 oxygen binding sites stated for octopod haemocyanin [53]. Total protein concentration was determined according to Bradford [54]. Bars depict means + 95% CI, n = 9–13. Differing letters indicate significant differences (P < 0.05) between octopod species for total haemolymph protein (upper case) or haemocyanin concentrations (lower case). White values on bars indicate the fraction of haemocyanin relative to total haemolymph protein and asterisks significant differences between species.

**Figure 6** pH at which the pH-dependent release of oxygen by haemocyanin becomes maximal. Comparison between the Antarctic *Pareledone charcoti*, the South-east Australian *Octopus pallidus* and the Mediterranean *Eledone moschata* at an experimental temperature of 10°C. Calculations include pH oxygen-saturation curves from all analysed PO$_2$. Letters indicate significant differences (P < 0.05) between species. Data from different PO$_2$ were pooled due to similar effects by PO$_2$ among species.

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**Table 2** Comparison of oxygen carrying capacities

| Species                  | Oxygen carrying capacity (mmol L$^{-1}$) | Source     |
|--------------------------|----------------------------------------|------------|
| *Megaleledone setebos*   | 1.86                                   | [33]       |
| *Octopus macropus*       | 1.60                                   | [55]       |
| *Pareledone charcoti*    | 1.58 (1.38-1.77, 14)                   | This study |
| *Enteroctopus dofleini*  | 1.36                                   | [56]       |
| *Octopus pallidus*       | 1.13 (0.98-1.29, 10)                   | This study |
| *Eledone moschata*       | 1.08 (0.94-1.22, 10)                   | This study |
| *Octopus vulgaris*       | 0.61                                   | [12]       |
| Red blooded Antarctic fishes | 1.77 (1.44-2.09, 11)                 | [8]        |

Values are listed in descending order. Numbers in brackets indicate 95% confidence intervals and samples size n, when available. Oxygen carrying capacities of red blooded Antarctic fishes were calculated from their haemoglobin content, based on a molecular weight of 66 kDa [57], and averaged for 11 species.
supports *Pareledone charcoti* in compensating for the poor oxygen unloading by its haemocyanin.

Oxygen supply is further enhanced by high levels of physically dissolved oxygen, as oxygen solubility increases with decreasing temperatures (e.g. by 40% from 15°C to 0°C, [72]). Consequently, dissolved oxygen contributes 18.5% to total haemolymph oxygen content. Given the small degree of putative venous oxygen unloading in *Pareledone charcoti* (below 20%), even at very low PO₂ (1 kPa), physically dissolved oxygen contributes a large fraction (42%, assuming an arterial-venous transition from 13 to 1 kPa PO₂ and pH 7.53-7.42, Figure 4B) of the oxygen supplied to tissues. Red-blooded Antarctic fishes also benefit from high ambient oxygen levels in the cold [73] and combined with low metabolic rates [5,65], this may be the key to the reduction in haemoglobin levels [74]. For *Pareledone charcoti* it rather seemed inevitable to increase haemocyanin concentrations, despite high dissolved oxygen levels, reduced oxygen affinity and metabolic rates lower than in fish [7]. Sustaining high haemocyanin levels may be energetically costly but may alleviate the pressure to evolve functional changes enabling complete oxygen unloading at 0°C. Such ‘complete’ compensation may not be possible considering the enormous size (3.5 MDa) and multimeric complexity of the haemocyanin molecule [53]. Although *Octopus pallidus* and *Eledone moschata* live at higher temperatures and lower dissolved oxygen levels, dissolved oxygen still contributes significantly to oxygen transport, especially towards colder temperatures when their haemocyanin increasingly fails to supply oxygen to tissues (Figure 4B).

**Temperature sensitivity of oxygen transport**
The increase in oxygen affinity and decrease of pH dependent oxygen release towards colder temperatures (Table 1, Figure 3) results in progressively reduced capacities to unload oxygen in all three octopod species. The change of oxygen affinity with temperature in *Pareledone charcoti* and *Eledone moschata* (Table 1) conforms with findings in other octopod species (in ΔP_{50} (kPa)/°C: 0.24, *Enteroctopus dofleini*; 0.20, *Octopus vulgaris* [12]; 0.14, *Eledone cirrhosa*; 0.10, *Octopus vulgaris* [75]). The temperature dependence of these data from the literature would have been even more pronounced if P_{50} were determined at an alpha-stat adjusted pH and not at a fixed pH of 7.4 across temperatures. Consequently, oxygen release and uptake by haemocyanin strongly depend on temperature in numerous octopod species, which poses a considerable challenge in the cold where high oxygen affinities diminish oxygen release to tissues [12,39].

However, some species deviate from this pattern, such as the Antarctic octopod *Megaleledone setebos*. The response of its haemocyanin oxygen affinity to temperature changes (0.01 kPa ΔP_{50}/°C, [33]), was 8–32 times less than that in any other octopod studied and 12 times less than the respective change in *Pareledone charcoti*. Despite similarities in oxygen affinity and oxygen carrying capacity between these two Antarctic octopods, this difference is striking. Thus, in addition to enhanced oxygen carrying capacities, two alternating strategies emerge to compensate for excessively high oxygen affinities in the cold: 1) A general decrease in oxygen affinity at all temperatures but with high sensitivity to temperature maintained as in *Pareledone charcoti* or 2) a considerable decrease of temperature sensitivity leading to reduced oxygen affinity at low temperatures only, as in *Megaleledone setebos*. Interestingly, *Octopus pallidus* seems to take advantage of both strategies as oxygen affinity barely changes between 10 and 15°C but strongly decreases between 15 and 20°C (Table 1, Figure 4). As a consequence, oxygen supply is sustained at temperatures below 10°C but also improves rapidly at higher temperatures (>15°C) when metabolic demand for oxygen increases. *Eledone moschata*, on the other hand, faces a constant increase of oxygen affinity and thus insufficient oxygen supply below 10°C (Table 1, Figure 4), which would contribute to cold-death at around 6°C (F. C. Mark, pers. obs.). Thus with respect to haemocyanin-mediated oxygen supply, *Octopus pallidus* seems to tolerate cold temperatures better than *Eledone moschata*.

Within the studied temperature ranges, warming hardly compromises the capacity for oxygen loading at the gills but does compromise oxygen release to tissues in all three octopods (Figure 4A). Only *Octopus pallidus*
will experience reduced arterial oxygen loading above 15°C (assuming arterial PO₂ is 13kPa), which however is paralleled by increased venous oxygen unloading (Figure 4). This conforms with findings in other cephalopods and indicates that temperature changes affect venous unloading more than arterial oxygen loading (Megaleledone setebos, Sepia officinalis, [33], Dosidicus gigas, [76], Todarodes sagittatus, [12]). Only few species like the giant squid Architeuthis monachus experience significantly reduced arterial saturation at higher temperatures [77]. Thus, in octopods, oxygen loading at the gills seems largely safeguarded at habitat temperatures and normoxic conditions and may only be compromised at low ambient oxygen levels.

The pH sensitivity of oxygen binding expressed as the Bohr coefficient remained unaffected by temperature changes in all three octopods, unlike in Megaleledone setebos or Sepia officinalis, whose Bohr coefficients decreased with falling temperatures [33]. Octopods may benefit from low Bohr coefficients in the cold, equivalent to a switch from pH dependent to PO₂ dependent oxygen release. This may preserve the venous oxygen reserve when metabolic rate is low and largely covered by elevated physically dissolved oxygen levels. The lower Bohr coefficient of Pareledone charcoti may also reflect its low activity mode of life in cold Antarctic waters where rapid pH dependent mobilization of the venous reserve is not required. Conversely, the strong increase in the Bohr coefficient of Megaleledone setebos haemocyanin during warming to 10°C (−2.33) challenges effective oxygen release outside of the animal’s usual thermal range [33]. In contrast to the findings in the Antarctic species, the maintenance of high Bohr coefficients in cold exposed temperate Octopus pallidus and subtropical Eledone moschata may reflect suboptimal or even impaired oxygen supply at their lower temperature margins.

Haemocyanin supports eurythermy
Pareledone charcoti benefits from its thermally sensitive oxygen binding during warming, as much of the bound oxygen is liberated then (Figure 4A). Figure 8 models the relationship of oxygen supply by haemocyanin, oxygen consumption and blood circulation rate at 0°C and 10°C and an assumed haemolymph volume of 5.2% (v/w, based on average literature values from Octopus vulgaris and Enteroctopus dofleini [78,79]). If oxygen supply by haemocyanin would remain constant from 0°C to 10°C (i.e. 0.34 mmol O₂ L⁻¹ at an arterial-venous transition from 13 to 4 kPa PO₂ and pH 7.53-7.42, Figure 4B), blood circulation would need to increase by 110.4% to match a rise of oxygen consumption from 0.63 mmol O₂ kg⁻¹ (wet mass) h⁻¹ at 0°C (taken from [7] to 1.35 mmol O₂ kg⁻¹ (wet mass) h⁻¹ at 10°C (MO₂) was extrapolated to 10°C using an average Q₁₀ of 2.12, taken from [46-48]. However, due to the large increase of oxygen supply by haemocyanin at 10°C (Figure 4B), demand for oxygen requires only a minimal increase in circulatory performance by 5.2% in Pareledone charcoti (Figure 8). Consequently, haemocyanin in Pareledone charcoti plays a major role in buffering oxygen demand when temperature increases and drastically reduces the workload for other circulatory components, particularly the hearts, which often limit ectotherm performance at high temperatures [80] such as in the cephalopod Sepia officinalis [40], fishes [81,82], or crustaceans [83,84]. Hence, haemocyanin function extends the range of oxygen dependent warm tolerance of Pareledone charcoti, which may cope far better with higher temperatures than Megaleledone setebos, whose haemocyanin, due to its low temperature sensitivity and extreme Bohr coefficient, barely supports oxygen supply at higher temperatures [33]. In fact, Pareledone charcoti sustains fully aerobic metabolism up to 8-10°C and thus tolerates elevated temperatures well [85]. Although both species are closely related and likely originate from shallow Southern Ocean waters (i.e. possess an ink sac [28], Figure 1), this may in part reflect the different geographic and vertical distribution of the two species. Megaleledone setebos is a circum-Antarctic species found between 30–850 m and most frequently below 100 m [49] where temperatures remain close to freezing all year round [86]. Pareledone charcoti inhabits the waters around the Northern Antarctic Peninsula mostly from less than 120 m [87] to very shallow waters (intertidal < 3 m, F. C. Mark, pers. obs.) and even visits tidal water pools [88] where water temperatures vary (e.g. from −0.5°C to 10.7°C during summer [86,89]). Our data provide first evidence that haemocyanin supports oxygen dependent eurythermy in an Antarctic invertebrate ectotherm and conform to analogous findings in the temperate, eurythermic crab Carcinus maenas [90]. Considering the strong warming trend at the Antarctic Peninsula [91], Pareledone charcoti may eventually benefit from its capacity to adjust oxygen supply to more variable temperatures than more stenothermal species.

Conclusions
This study highlights the importance of the oxygen transport pigment haemocyanin in octopods with regard to temperature compensation. In comparison to findings in the south east Australian Octopus pallidus and the Mediterranean Eledone moschata, the analysis of blood oxygen binding in the Antarctic octopod Pareledone charcoti revealed properties of its blood pigment haemocyanin that support oxygen supply in the cold but at the same time maintain haemocyanin function in the warmth. Significantly lower oxygen affinity but incompletedeoxygenation of haemocyanin in Pareledone charcoti resulted in sustained but poor oxygen unloading at 0°C, which however, was compensated for by high levels of
dissolved oxygen as well as elevated haemocyanin concentrations and thus oxygen carrying capacities. In contrast to the stenothermic Antarctic octopus *Megaleledone setebos*, *Pareledone charcoti* benefits from a thermally sensitive haemocyanin that extends oxygen supply at warmer temperatures and thus supports oxygen dependent eurythermy. Compromised oxygen release from haemocyanin in the cold underlines the crucial role of the pigment for defining cold tolerance not only in Antarctic but also in warmer water octopods. While some warmer water octopods succeed to extend oxygen supply at cold temperature margins by e.g. reduced temperature sensitivity of oxygen binding in the cold others fail to do so. However, for a complete picture of thermal tolerance in *Pareledone charcoti* and the other octopods much more information is needed regarding in vivo haemolymph PO2 and pH under rest and exercise, the role of cardiac and circulatory performance, aerobic scope and growth rates across various temperatures as well as acclimation capacities. Only then may one predict the future role of this abundant group of ectotherms in a rapidly warming ecosystem.

**Methods**

**Study design**

To assess whether blood oxygen transport in Antarctic octopods exhibits features that support oxygen supply in the cold, we compared oxygen binding properties, total oxygen carrying capacities as well as protein and ion composition of haemolymph of the Antarctic octopod *Pareledone charcoti* with two octopods originating from warmer waters - *Octopus pallidus* and *Eledone moschata*. Comparisons were performed at habitat temperatures and at a common temperature of 10°C, assuming that all haemocyanin types remained functional at these temperatures. To evaluate if earlier observations for *Megaleledone setebos* haemocyanin apply to other Antarctic octopods as well, we chose *Pareledone charcoti* as a representative of the most abundant and more typically sized genus *Pareledone* [20,87]. Temperature sensitivity of oxygen binding was analysed in all three species to assess the role of octopod haemocyanin in oxygen supply across the respective habitat temperatures.

**Animals and sampling**

The octopod *Pareledone charcoti* belongs to the Southern Ocean endemic octopod family Megaleledonidae (Figure 1, [28]) and inhabits the shallow shelf area around the Antarctic Peninsula [87] with temperatures varying between −1.9 to +2°C [92]. Using bottom trawls, specimens were collected on the RV Polarstern cruise ANTXXVIII/4 in March 2012, at depths between 90–470 m around Elephant Island (61°S, 56°W, cruise details [93]), where temperatures ranged between 0.1 to 1.6°C and salinities between 34.3-34.6 psu. *Octopus pallidus* belongs to the non-polar shallow water octopod family Octopodidae (Figure 1, [28]) and inhabits the well mixed waters in South East Australia with habitat temperatures ranging from 12.8°C from winter to summer [94,95]. Specimens were caught in July 2012, between 40–50 m depth, in the western Bass Strait near Stanley (41°S, 145°E) by fishermen.
(T.O.P. Fish Pty Ltd.) using plastic octopus pots and then transported and kept overnight in large tanks connected to a flow-through seawater system at the Institute for Marine and Antarctic Studies, Hobart. Eledone moschata belongs to the non-polar octopus family Eledonidae (Figure 1, [28]) and occurs all over the Mediterranean Sea mainly at depths between 0–200 m [96,97]. Specimens were fished in November 2008, between 20–40 m depth using bottom trawls, in the northern Adriatic Sea near Chioggia, where habitat temperatures vary largely, both by depth and seasonally, between approximately 10-23°C [98]. Average body masses were 32.5 g (28.0-37.0) for Pareledone charcoti, 563.3 g (481.0-645.7) for Octopus pallidus and 52.9 g (35.6-70.2) for Eledone moschata. All animals were anaesthetised in 3% ethanol [99] until non-responsive, then ventrally opened to withdraw haemolymph from the cephalic vein, the afferent branchial vessels and the systemic heart and finally killed by a final cut through the brain (Animal research permit no. 522-27-11/02-00 (93), Freie Hansestadt Bremen, Germany and animal ethics approval no. AEC12-43, La Trobe University, Bundoora, Australia). Haemolymph samples were spun down at 15.000 g for 15 min at 0°C to pellet cell debris and supernatants were stored at −20°C.

**Phylogenetic analysis**

To illustrate phylogenetic relationships of Pareledone charcoti, Octopus pallidus and Eledone moschata, we performed Bayesian phylogenetic analysis using five genes, the mitochondrial genes cytchrome oxidase subunit I and III and the nuclear genes rhodopsin, octopine dehydrogenase and 16S rDNA of 16 octopodiform species (GenBank accession numbers in Additional file 1). Sequences for each gene were aligned separately using the MUSCLE plugin of Geneious 7.1.7 [100]. The resulting alignments were curated using GBLOCKS 0.91b [101,102] tolerating gap positions within final blocks and concatenated to one data set. Based on the Akaiki Information Criteria [103], JModeltest 2.1.5 [104], identified the GTR + I + G model as the best substitution models for the concatenated data set. Bayesian trees were constructed using MrBayes (v. 2.0.3) [105] as implemented in Geneious running at least two independent Monte Carlo Markov Chain (MCMC) analysis with 2,000,000 generations sampled every 2,000 generations. The appropriate burnin was chosen based on the resulting traces, which showed a stationary distribution before 10% of the MCMC chain. Vampyroteuthis infernalis and Argonauta nodosa were used as outgroups.

**Blood characteristics**

**Oxygen binding properties**

Oxygen binding of octopod haemocyanin was characterised using a modified diffusion chamber (for details see [106]), which simultaneously measures pigment oxygenation and pH in a 15 μl sample. Experiments were performed at common habitat temperatures of each species (0°C Pareledone charcoti, ~10-20°C Octopus pallidus, ~10-20°C Eledone moschata) and at a comparative temperature of 10°C. The temperature was monitored and controlled via a temperature sensor (PreSens, Germany) and a connected water bath with a thermostat (LAUDA Ecoline Staredition RE 104, Germany), filled with an anti-freeze solution (20% ethylene glycol, AppliChem, Germany). Prior to measurements, aliquots of 18 μl thawed haemolymph were spun down to collect all liquid at the bottom of a 1.5 ml microcentrifuge tube (5 sec at 1000 g), preconditioned with pure oxygen gas to deplete dissolved carbon dioxide (CO₂) and 0.6-0.9 μl of 0.2 mmol L⁻¹ NaOH (8–12 μmol L⁻¹ final concentration) added to raise haemolymph pH above 8.0 to ensure full oxygenation. To account for the pronounced pH sensitivity of cephalopod pigments [52], changes of pH and absorbance were recorded at 347 nm in 15 μl haemolymph, at continuously decreasing PCO₂/pH (0–10 kPa/~ pH 8.1-6.8) and four constant PO₂ levels (21, 13, 4, 1 kPa, after Pörtner [52]), with gas mixtures being supplied by gas mixing pumps (Wösthoff, Germany). The spectrophotometer (USB2000+, Ocean Optics, USA) was set to 15 milliseconds integration time, 100 scans to average and 30 seconds measurement intervals and calibrated by recording light and dark spectra without sample. Prior to each experiment, the pH optode was calibrated in MOPS-buffered (40 mmol L⁻¹, 3-(N-Morpholino) propanesulfonic acid), filtered artificial seawater (35 psu) equilibrated to the respective experimental temperature at six pHs ranging from 6.7 to 8.1. The pH of buffers was checked with a pH glass electrode (InLab Routine Pt1100, Mettler Toledo, Germany) and a pH meter (pH 330i, WTW, Germany), calibrated with low ionic strength NIST pH standards (AppliChem, Germany, DIN19266) and corrected to Free Scale pH with Tris-buffered seawater standard (Dickson, CO2 QCLab, batch 4 2010, USA, [107]) equilibrated at the same temperature. The pH signal was corrected for instrumental drift and for effects of auto-fluorescence intrinsic to haemolymph [106] and is presented here on the free hydrogen ion scale [108].

Each experiment involved the calibration with pure oxygen or nitrogen to obtain maximum and minimum oxygenation signals. Correct pigment saturation was calculated by continuous readjustments of the maximum oxygenation signal to account for its linear drift observed during the course of an experiment [106,109]. While the maximum oxygenation signal did not change within the range of temperatures employed for each species, the minimum oxygenation signal increased towards colder temperatures due to incomplete oxygen unloading, even under pure nitrogen and low pH (~6.6). For such experiments we predicted minimum absorbance from a reference wavelength of the first recorded spectrum with an uncertainty
of 5%, based on a linear regression model applied to 20 experiments with fully deoxygenated pigments (Additional file 2).

To determine the total oxygen bound to octopod haemocyanin (i.e. oxygen carrying capacity) 10 μl of thawed haemolymph were equilibrated with pure oxygen gas in a microcentrifuge tube on ice for 10 min and transferred with a gas tight Hamilton syringe to a gas sealed chamber containing 2 ml of a 32°C warm cyanide solution (55 mmol L\(^{-1}\) potassium cyanide, 3 g L\(^{-1}\) [110]). Two high-resolution Oxygraph-2 k respirometers (OROBOROS Instruments, Innsbruck, Austria) and DatLab analysis software (version 5.1.0.20) recorded the liberated oxygen (nmol ml\(^{-1}\)), corrected for air pressure, temperature and background oxygen flux. For each experiment, the respirometers were calibrated with air at the beginning and sodium dithionite added at the end for a zero calibration. The contribution of dissolved oxygen was experimentally determined by the addition of ice-cold, oxygen saturated, filtered seawater (35 psu). The observed change of oxygen concentration was then subtracted from the haemolymph measurements to obtain the final oxygen carrying capacity of haemocyanin.

**Alpha-stat pattern of haemolymph pH**

To be able to analyse oxygen binding parameters at various temperatures, we assessed whether the pH of octopod haemolymph follows an alpha-stat pattern [111] or remains constant across temperatures (i.e. pH stat pattern). Replicated measurements on 20 μl thawed haemolymph of *Octopus pallidus* at 0°C, 10°C and 20°C, using a micro pH electrode (InLab Ultra-Micro, Mettler Toledo, Germany), showed that pH decreases linearly with temperature \(b = -0.0153 \text{ pH units / °C, } t = -9.71, P < 0.001, R^2 = 0.75\), Figure 7), analogous to an imidazole buffered system \(\text{pH} = 0.75\), [111]). pH analysis of freshly sampled haemolymph from other species confirmed that octopod haemolymph follows this linear pH-temperature relationship in vivo (Figure 7) and therefore exhibits an alpha-stat pattern as also demonstrated for squid [63]. Hence, venous and arterial pH were determined on this basis for various temperatures.

**Protein and ion concentration**

Protein content of octopod haemolymph was determined according to Bradford [54]. Thawed haemolymph was diluted tenfold (v:v) with stabilising buffer (in mmol L\(^{-1}\), 50 Tris–HCl, 5 CaCl\(_2\), 6 H\(_2\)O, 5 MgCl\(_2\), 6 H\(_2\)O, 150 NaCl, pH 7.47 at 22°C) and 5 μl mixed with 250 μl Bradford reagent (Bio-Rad, Germany). Following 10 min incubation at room temperature, absorbance was recorded at 595 nm using a microplate spectrophotometer (PowerWave HT, BioTek, U.S.A.), Bovine albumin serum served as protein standard to calculate total protein concentrations.

Concentrations of functional haemocyanin \((c(Hc))\) in haemolymph were derived from the oxygen carrying capacity \((C_{O_2})\), the molecular weight (MW) of octopod haemocyanin (3.5 MDa) and its 70 oxygen binding sites \(n(HcO_2)\), [53], Equation 1).

\[
c(Hc) = \frac{C_{O_2}}{n(HcO_2)} \times MW
\]

Results from tests with thawed haemolymph of *Octopus vulgaris* (mean ± S.D., 54.3 ± 6.9 g L\(^{-1}\)) agreed well with data obtained from freshly observed haemolymph via atomic absorption spectroscopy (55.9 ± 7.4 g L\(^{-1}\), [35]), which not only confirmed the accuracy of our approach but also that storage at -20°C does not affect the oxygen binding capacity of cephalopod haemolymph [61]. Although inorganic ions such as Mg\(^{2+}\) or Na\(^+\) can affect oxygen affinity in octopods [60], they seem to be insignificant regulators of oxygen binding in most cephalopods [61]. To verify this for the observed species, we diluted haemolymph 400-fold with deionised water and determined cation concentrations by ion chromatography (ICS-2000, Dionex, Germany) following cation separation by an IonPac CS 16 column (Dionex, Germany) with methane sulfonic acid (MSA, 30 mmol L\(^{-1}\)) as an eluent at 0.36 ml min\(^{-1}\) flow rate and 40°C. Ion concentrations were derived from the peaks corresponding to the Dionex Combined Six Cation Standard-II.

**Data analysis**

Processing of raw data and statistical analysis was performed using the ‘R’ statistical language R Core [112]. Recordings of pH and pigment oxygenation were time-matched and analysed in pH/saturation diagrams, most suitable for pH sensitive pigments like cephalopod haemocyanin [52]. An empirical five parameter logistic model was applied (‘drc’ add-on package, [113]) to fit sigmoidal curves to the pH/saturation data [106]. Resulting pH oxygen-saturation curves display the change of pigment oxygenation linked proton binding was expressed as the change of molar oxygen concentration per pH unit \((\Delta \text{mmol O}_2 \text{ L}^{-1}/\Delta \text{pH})\) and calculated from the oxygen carrying capacity and the maximum slope of a fitted pH oxygen-saturation curve. The calculation of putative in vivo oxygen saturation and oxygen release required knowledge of in vivo venous and arterial PO\(_2\) and pH. Venous and arterial haemolymph PO\(_2\) were not determined for *Octopus pallidus*, Pareledone charcoti and Eledone moschata in this
study and were assumed to correspond to haemolymph PO₂ of Octopus vulgaris, which showed an arterial PO₂ of 13 kPa and a venous PO₂ of 4 kPa for resting and 1 kPa for exercised specimens under normoxic conditions [50,51]. Arterial and venous haemolymph PO₂ were further assumed to be constant across temperatures. Venous haemolymph pH was inferred from the alpha-stat regression slope determined for Octopus pallidus in this study and corresponded to pH of freshly sampled venous haemolymph of Pareledone sp. and Eledone moschata (Figure 7). Due to the difficulty in obtaining arterial haemolymph from small sized octopods, arterial haemolymph pH could not be determined and instead assumed to be 0.11 pH units higher than venous pH as reported for Octopus vulgaris [50].

Differences between species and experimental temperatures were tested to be significant (P < 0.05) using analysis of variance (ANOVA) followed by Tukey's post hoc test. Non-parametric Kruskal Wallis and Mann–Whitney tests were used to compare the cooperativity of oxygenation dependent proton binding among species. Normality and homogeneity of variance were assessed by Kolmogorov–Smirnov and Levene's tests, respectively. Results were expressed as means and their 95% confidence interval range if not stated otherwise.

Additional files

Additional file 1: GenBank accession numbers of molecular sequences used for phylogenetic analysis.

Additional file 2: Incomplete desaturation of octopus haemocyanin at low temperatures requires correct identification of the zero calibration point. (A) At 10°C haemocyanin of e.g. Octopus pallidus fails to fully deoxygenate under pure nitrogen gas and very low pH. Deoxygenation only completes when temperatures increase above 10°C, which complicates the determination of the zero calibration point at low temperature measurement. (B) A linear regression between the absorbance at a reference wavelength (421.75 nm) of the first recorded spectrum and the absorbance peak at 348 nm of fully deoxygenated octopus haemocyanin helped to predict the true zero calibration point at low temperatures for low temperature measurements. The reference absorbance signal at 421.75 nm was selected, as the sum of squares of the differences between the predicted and measured zero calibration point across 20 experiments were lowest at this wavelength.

Abbreviations

ATP: Adenosine triphosphate; c(Hc): Haemocyanin concentration; CO₂: Carbon dioxide; CO₂: Oxygen carrying capacity; MO₂: Oxygen consumption rates of Pareledone charcoti (MgO₂); MW: Molecular weight; NIST: National Institute of Standards and Technology; PCO₂: Carbon dioxide partial pressure; PO₂: Oxygen partial pressure; P₅₀: PO₂ at which the pigment is half saturated; pHₐ₅₀: pH of haemolymph/blood at which the pigment is half saturated; Q₁₀: Temperature coefficient; psu: Practical salinity units; MCMC: Monte Carlo Markov Chain.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

MO and FCM developed the study design, compiled the manuscript and interpreted results. MO and FCM acquired the samples. JS supported the sample acquisition and haemolymph sampling of Octopus pallidus. MO performed the experiments and the analysis. HOP, BL and JS contributed to data interpretation and manuscript revision. All authors read and approved the final manuscript.

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References

1. Klink JC, Hofmann EE, Beardsley RC, Sallhoglu B, Howard S. Water-mass properties and circulation on the west Antarctic Peninsula Continental Shelf in Autumn Fall and Winter 2001. Deep-sea Res Pt I. 2004;51:1925–46. 2. Jacobs SS, Amos AF, Bruchhausen PM. Ross sea oceanography and antarctic bottom water formation. Deep-sea Res Oceanogr Abstr. 1970;17:935–62. 3. Portner HO, Peck L, Somero G. Thermal limits and adaptation in marine Antarctic ectotherms: an integrative view. Philos Trans R Soc Lond B Biol Sci. 2007;362:2233–58. 4. Siedl BD, O'Brien KM. When bad things happen to good fish: the loss of hemoglobin and myoglobin expression in Antarctic icefishes. J Exp Biol. 2006;209:1791–802. 5. Clarke A, Johnston NM. Scaling of metabolic rate with body mass and temperature in teleost fish. J Anim Ecol. 1999;68:893–905. 6. Ahn I-Y, Shirm JH. Summer metabolism of the Antarctic clam, Laternula elliptica (King and Broderip) in the temperature range 2°C to 26°C. J Exp Mar Biol Ecol. 1998;224:53–64. 7. Daly H, Peck L. Energy balance and cold adaptation in the octopus Pareledone charcoti. J Exp Mar Biol Ecol. 2000;245:197–214. 8. Wells RMG, Ashby MD, Duncan SJ, Macdonald JA. Comparative study of the erythrocytes and haemoglobins in nototheniid fishes from Antarctica. J Fish Biol. 1980;17:517–27. 9. Ruud JT. Vertebrates without erythrocytes and blood pigment. Nature. 1954;173:848–50. 10. Siedl BD. Intracellular oxygen diffusion: the roles of myoglobin and lipid at cold body temperature. J Exp Biol. 1998;201:1119–28. 11. Weber RE, Campbell KL. Temperature dependence of haemoglobin-oxygen affinity in heterothermic vertebrates: mechanisms and biological significance. Acta Physiol. (Oxf). 2011;203:549–62. 12. Brix O, Bårdgard A, Cau A, Colosimo A, Condo S, Giardina B. Oxygen-binding properties of cephalopod blood with special reference to environmental temperatures and ecological distribution. J Exp Zool. 1989;252:34–42. 13. Brix O, Colosimo A, Giardina B. Temperature dependence of oxygen binding to cephalopod haemoglobins: ecological implications. Mar Freshw Behav Physiol. 1994;25:149–62. 14. Macdonald JA, Wells RMG. Viscosity of Body Fluids From Antarctic Nototheniid Fish. In: Di Prisco G, Maresca B, Tota B, editors. Biology of Antarctic Fish. Berlin: Springer Berlin Heidelberg; 1991. p. 163–78. 15. D'Avino R, Di Prisco G. Hemoglobin from the Antarctic fish Notothenia coriceps neglecta. Eur J Biochem. 1989;179:699–705.
72. Boutillier RG, Heming TA, Iwama GK. Appendix: physico-chemical parameters for use in fish respiratory physiology. Fish Physiol. 1984;10:403–30.

73. Di Prisco G, Macdonald JA, Brunori M. Antarctic fishes survive exposure to carbon monoxide. Experimentia. 1992;48:473–5.

74. Di Prisco G, Eastman JT, Giordano D, Parisi E, Verde C. Biogeography and adaptation of Nototocardis fish. Hemoglobin function and globin-gene evolution. Gene. 2007;398:143–55.

75. Bridges CR, Bohr and root effects in cephalopod haemocyanins - paradox or pressure in Sepia officinalis? Mar Freshw Behav Physiol. 1994;25:121–30.

76. Seibel BA. The jumbo squid, Dosidicus gigas (Ommastrephidae), living in oxygen minimum zones II: Blood-oxygen binding. Deep-Sea Res Pt II. 2012;59:139–44.

77. Brix O. Giant squids may die when exposed to warm water currents. Nature. 1983;303:42–3.

78. Martin A, Harrison F, Huston M, Stewart D. The blood volumes of some representative molluscs. J Exp Biol. 1958;35:260–79.

79. O’Dor RK, Wells MJ. Circulation time, blood reserves and extracellular space in a cephalopod. J Exp Biol. 1984;113:461–4.

80. Pörtner HO. Climate variations and the physiological basis of temperature dependent biogeography: systemic to molecular hierarchy of thermal tolerance in animals. Comp Biochem Physiol Part A Mol Integr Physiol. 2002;132:739–61.

81. Farrell AP. Cardiorespiratory performance in salmonids during exercise at high temperatures: insights into cardiovascular design limitations in fishes. Comp Biochem Physiol Part A Mol Integr Physiol. 2002;132:797–810.

82. Iftikar FI, Hickey AJR. Do mitochondria limit hot fish hearts? Understanding the role of mitochondrial function with heat stress in Notolatous celidotus. Plos One. 2013;8:e46120.

83. Frederich M, Pörtner HO. Oxygen limitation of thermal tolerance defined by cardiac and ventilatory performance in spider crab, Maja squinado. Am J Physiol Regul Integr Comp Physiol. 2000;279:R131–8.

84. Stillman J, Somero G. Adaptation to temperature stress and aerial exposure in congeneric species of intertidal porcelain crabs (genus Petrolisthes): correlation of physiology, biochemistry and morphology with vertical distribution. J Exp Biol. 1996;199:1845–55.

85. Pörtner HO, Zielinski S. Environmental constraints and the physiology of performance in squids. S Afr J Mar Sci. 1996;20:207–21.

86. Barnes DKA, Fuentes V, Clarke A, Schloss IR, Wallace MI. Spatial and temporal variation in shallow seawater temperatures around Antarctica. Deep-Sea Res Pt II. 2005;53:835–65.

87. Allcock AL. On the confusion surrounding Pareledone charcoti (Joubin, 1905) (Cephalopoda : Octopodidae): endemic radiation in the Southern Ocean. Zool J Linn Soc. 2005;143:75–108.

88. Joubin L. Description de deux Élédones provenant de l’Antarctique. Mémoires de la Societé Zoologique de France. 1905;18:22.

89. Barnes DKA, Rothe P, Clarke A. Colonisation and development in encrusting communities from the Antarctic intertidal and sublittoral. J Exp Mar Biol Ecol. 1996;196:251–65.

90. Giomi F, Pörtner H-O. A role for haemolymph oxygen capacity in heat tolerance of eurythermal crabs. Front Physiol. 2013;4:110.

91. Meredith MP, King JC. Rapid climate change in the ocean west of the Antarctic Peninsula during the second half of the 20th century. Geophys Res Lett. 2005;32, L19604.

92. Dininman MS, Klink JM, Smith JR WD. A model study of Circumpolar Deep Water on the West Antarctic Peninsula and Ross Sea continental shelves. Deep-Sea Res Pt II. 2011;58:1508–23.

93. An information portal for research platforms of the Alfred Wegener Institute for Polar and Marine Research. [http://expedition.awi.de/expedition?d=25#events]

94. Leporati SC, Pecl GT, Semmens JM. Cephalopod hatchling growth: the effects of initial size and seasonal temperatures. Mar Biol. 2006;151:1375–83.

95. André J, Pecl GT, Grist EPM, Semmens JW, Haddon M, Leporati SC. Modelling size-at-age in wild immature female octopuses: a bioenergetics experimental approach. Mar Ecol Prog Ser. 2009;389:159–74.

96. Belcari P, Tsereges G, González M, Lefkaditou E, Marceta B, Manfrin GP, et al. Distribution and abundance of Eledone cirrhosa (Lamarck, 1798) and Eledone moschata (Lamarck, 1798) (Cephalopoda: Octopodida) in the Mediterranean Sea. 2002.

97. González M, Sánchez P. Cephalopod assemblages caught by trawling along the Iberian Peninsula Mediterranean coast. 2002.