Reply to: methodological inconsistencies define thermal bottlenecks in fish life cycle

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Abstract
The identification of biological pattern is often complicated by the lack of methodologically consistent data with broad geographic coverage, especially when considering functional characteristics of organisms that differ greatly in body size and morphology. In our study (Dahlke et al. 2020), we addressed the problem of data scarcity by using different types of observational and experimental data together with statistical (phylogenetic) data imputation, and by placing our analysis into the context of a physiological concept, which provides a mechanism-based explanation for the observed pattern (ontogenetic shift in thermal tolerance of fish) and with respect to transition from sublethal to lethal thresholds. Here, we show with comparative examples that our results were not affected by the use of methodologically inconsistent data.

In their comment, Pottier et al. questioned some of our methods and results, including the observation that the heat tolerance limit of reproductive (spawning) fish and embryos is on average ~10 °C lower than that of non-reproductive adults. The critique by Pottier et al. included the claim that our (unconventional) approach for estimating the temperature limits of spawners and embryos (occurrence data, sublethal thresholds and lethal thresholds determined by different methods) systematically underestimated the heat tolerance of these life stages compared to adult fish for which we considered conventional CTmax and LT50 values previously compiled by Comte and Olden (2017).

We generally agree that the use of different methods and tolerance metrics is not ideal in the strict sense of a comparative analysis, and acknowledge that the description of our analyses was not precise enough for a reader to pick up all details. However, Pottier et
al. disregarded that we placed our analysis into the context of a physiological concept (i.e., oxygen- and capacity-limited thermal tolerance, OCLTT) that is suitable to explain the observed difference in thermal tolerance between life stages and that also captures the transition from sublethal to lethal thresholds. Dahlke et al. 2020 provided additional data (temperature-dependent development and metabolism) consistent with physiological theory and the observed pattern of life stage specific thermal tolerance. The results and conclusions presented in Dahlke et al. 2020 also agree well with previous analysis of smaller, methodological more consistent data sets (e.g., Rombough 1997). With this line of evidence, we are confident that the results presented in Dahlke et al. 2020 were not significantly biased by the use of different tolerance metrics and variability in experimental protocols applied for species and life stages with often extremely different body sizes, morphologies and husbandry requirements. It is therefore surprising to us that Pottier et al. question the validity of our results without providing a formal analysis.

The illustrations by Pottier et al. imply that the embryo data we used would always fall below conventional CT\textsubscript{max}/LT\textsubscript{50} values. While we can demonstrate that this assumption is not entirely correct, we must note that it was not clear from the method text that the embryo data presented in Dahlke et al. 2020 also included LT\textsubscript{50} values, such as those extracted from Rombough (1997). These LT\textsubscript{50} values were determined based on (statistically significant) regressions and therefore correspond to our definition of the heat tolerance limit of embryos (“temperatures causing statistically significant change in survival relative to the optimum value”).

To show that the difference in heat tolerance between embryos and adults was not confounded by methodological inconsistencies, we followed the recommendation of Pottier et al. and compared the main data set of Dahlke et al. 2020 (662 species including criticized embryo data) with a subset of data including only LT\textsubscript{50} values for embryos (from Rombough 1997) and available CT\textsubscript{max}/LT\textsubscript{50} values for adults of the same species (37 species, Fig. 1a). Both data sets indicate a substantial increase in heat tolerance from embryos to adults (Fig. 1a), with a median difference between life stages of 10.96 °C in the main data set and 10.80 °C in the data subset (Fig. 1b). A priori, we used the data set compiled by Comte and Olden (2017) to confirm that differences between adult CT\textsubscript{max} values (median 35.8 °C, 346 observations, 246 species) and adult LT\textsubscript{50} values (median 35.3 °C, 427 observations, 313 species) are statistically not significant (P = 0.1633), using a generalized additive mixed model to account for the latitudinal origin of species and phylogenetic independence (as described in the supplement of Dahlke et al. 2020).

Furthermore, based on temperature-dependent hatching success (i.e. embryo survival) in polar cod (Boreogadus saida, Fig. 2a) and Atlantic cod, (Gadus morhua, Fig. 2b) from different regions (Dahlke et al. 2016, Dahlke et al. 2017, Dahlke et al. 2018, Laurel et al. 2018) we show that conventional LT\textsubscript{50} values differ marginally from the values determined as described in Dahlke et al. 2020. More explicitly, LT\textsubscript{50} values determined by nonlinear regression analysis (R\textsuperscript{2} > 0.7, P < 0.001) across two populations of B. saida (3.41 °C, 95% confidence range: 2.94 to 3.50 °C, Fig. 1c) and four populations of G. morhua (9.14 °C, 95% confidence range: 8.94 to 9.74 °C, Fig. 1d) fall within the range of values determined according to Dahlke et al. (2020) for individual populations (B. saida: 3.0 to 3.5 °C, G. morhua: 8.0 to 9.5 °C). Adult CT\textsubscript{max} values of B. saida (14.9 to 17.1 °C) (Drost et al. 2016) and G. morhua (21.4 to 23 °C) (Norin et al. 2019) are more than 10 °C higher compared to embryo LT\textsubscript{50} values.
As pointed out by Pottier et al., some embryo data originally published in a Japanese series (e.g., Mito 1963) and extracted from a compilation by Motani and Wainwright (2015) correspond to occurrence limits rather than physiological limits as described in our methods. It is important to note that in contrast to terrestrial ectotherms, the distributional limits of aquatic species and life stages often coincide with their physiological temperature limits (Sunday et al. 2012, see also figure S1 in the supplement of Dahlke et al. 2020 for data on spawning fish).

Pottier et al. suggest heartbeat measurements to determine \( CT_{\text{max}} \) values of embryos and provide an example (Del Rio et al. 2019) where eyed-stage embryos of chinook salmon (\textit{Oncorhynchus tshawytscha}) exhibit heartbeat-based \( CT_{\text{max}} \) values (30 °C) similar or higher than juveniles. Here it is important to note that embryos of fish species (including salmonids) are most heat-sensitive during early embryogenesis prior to the development of the heart and circulatory system (see Rombough 1997). In our study, we considered for this species the embryo \( LT_{50} \) tolerance range (3–15 °C) reported in Rombough (1997), which [consistent with the conclusions of Del Rio et al. (2019)] represents a critical bottleneck in the life cycle of chinook salmon.

Of course, it is difficult to directly compare the temperature limits of spawners observed in the field to the experimentally determined \( CT_{\text{min/max}} \) values of adults or larvae. However, the agreement we demonstrated between very narrow field and laboratory tolerance limits of spawners (figure S1 in the supplement our paper) confirms that these data are highly relevant for determining the climate change vulnerability of fish species. In this context it is also important to note that the tolerance metric used for spawners is lowered substantially by
gonads and gametes including their development and gamete fertility. As a counter example, Pottier et al. cited a study (Auer et al. 2021) showing only a slight decrease in CT_{max} of females of a viviparous fish during late gestation. This study does not contradict our results because the effects of elevated temperatures on reproductive performance and fertility were not assessed by Auer et al.. On the contrary, Auer et al. (2021) provide strong support for the physiological framework of our study (OCLTT) which focuses on sub-lethal (performance) constraints as the first line of thermal intolerance (Pörtner 2001). This also concerns gonad development and gamete fertility (Pörtner 2021).

Pottier et al. highlight some recent studies that propose promising means for standardizing CT_min/max-procedures partly based on experiments with Drosophila (Jørgensen et al. 2019, Jørgensen et al. 2021). It is therefore important for us to mention that one of the best examples supporting the ecological relevance of the temperature limit of fertility and reproduction comes from a study with Drosophila species (van Heerwaarden and Sgrò 2021). The study found that CT_{max} values of adults are \(-10^\circ C\) higher than male fertility limits and egg-to-adult limits. Higher temperature values for CT_{max} than reproductive performance may be more universal, in line with our conclusion that sublethal temperature thresholds for reproductive performance characterize narrow tolerance windows representing a critical bottleneck in the lifecycle of fish and possibly other taxa.

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