Temperature-Dependent Alterations in Metabolic Enzymes and Proteins of three Ecophysiologically Different Species of Earthworms

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ABSTRACT

The effects of varying temperatures (12 - 44 °C) on the specific activity of cytoplasmic malate dehydrogenase (cMDH), mitochondrial malate dehydrogenase (mMDH) and lactate dehydrogenase (LDH) of some earthworms (Metaphire posthuma, Perionyx sansibaricus and Lampito mauritii) were studied. The effects of different temperatures on supernatant and mitochondrial protein contents were also investigated. The specific activities of cMDH, mMDH and LDH of the earthworms decreased gradually as a function of increasing temperature from 12 to 44 °C. Higher metabolic energy was needed to maintain the activity at low temperatures. Hence, the earthworms showed increased enzyme specific activity at low temperatures. However, the protein content increased up to 28 °C. Afterwards, with the increase in the temperature from 28 to 42 °C, the proteins in the earthworms showed a significant decrease. The temperature-associated changes in the protein content could be explained by the fact that protein synthesizing capacity was hampered above and below the optimum temperature range. The most pronounced effects of varying temperatures were on P. sansibaricus. It might be due to the epigeic nature of the earthworm species. Then minimum effect was on the endogeic earthworm M. posthuma. Virtually, the differences in the enzymes physiology were associated with the differences in the ecological categories of the earthworms. This clearly demonstrate a possible link between the physiology and ecology at aerobic (cMDH, mMDH) and anaerobic (LDH) levels in the tropical earthworms.

Key words: Enzymes, proteins, earthworms, temperatures

INTRODUCTION

Temperature has a profound effect on the metabolism of organisms. Depending on the species, earthworms possess varying temperature optima and tolerances and even adopt to cope with temperature extremes. They may show long-term responses to chronic temperature change both in laboratory and in natural habitat. A very limited numbers of metabolic enzymes of oligochaetes have been studied to demonstrate the temperature effects. Pomert and Zarrow (1936) showed the effect of temperature on the respiration of the earthworm. Saroja (1961) documented seasonal acclimatization of oxygen consumption to temperature in tropical earthworm Megascolex mauritii. Temperature may be a factor of primary importance in determining the composition and structure of earthworm communities (Lavelle, 1983; Lavelle et al., 1989). Hazel (1995) described

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thermal adaptation in biological membranes. Temperature affects the lipid composition of the earthworm _Lumbricus rubellus_ and _Eisenia nordenskioeldi_ (Petersen and Holmstrup, 2000). It appears to be the most important environmental variable influencing the growth, metabolism and biology of earthworms. Tripathi and Bhardwaj (2004a) suggested that the earthworm population mainly decreased with the increase in soil temperature. Tripathi and Bhardwaj (2004b) made a comparative assessment of biomass growth of _Eisenia fetida_ and _Lampito mauritii_ at different temperatures. The biomass of both the species of earthworms varied significantly with the changes in temperature. The optimum temperatures for _E. fetida_ and _L. mauritii_ were 25 and 30°C, respectively.

Different earthworms have their own optimum temperature (Lee, 1985). A Siberian species, _Eisenia nordenskioeldi_ (Eisen) lives in the areas with permafrost and has developed freeze-tolerance mechanism for survival in the extreme habitat, where more than one summer season is required to complete a lifecycle (Mazantzeva, 1985; Holmstrup and Petersen, 1997). Holmstrup et al. (1999) reported that _E. nordenskioeldi_ synthesized and accumulated glucose as an immediate response to ice formation in extracellular body fluids. Glucose accumulation has also been detected in the freeze-intolerant _L. rubellus_ upon freezing but at much lower concentrations. Depending on the species, earthworms possess varying temperature optima and tolerances as well as different strategies for coping with the environmental temperatures extremes (Holmstrup and Zachariassen, 1996). Invertebrates, microbes and higher organisms have been shown to respond to temperature changes by adjusting the composition of membrane phospholipid fatty acids (Hazel, 1995). The earthworm, _Lumbricus terrestris_ produces carbonic acid at constant rate when is exposed to temperatures between 10 and 22°C. It can compensate its rates of oxygen consumption between 5 and 15°C after a two week acclimation to these temperatures (Fitzpatrick et al., 1987). The rate of respiration of earthworms is very sensitive to the environmental temperature. The respiration rate increases as the body temperature rises. The temperature effect on animal activity related to aerobic metabolism is very common. Oxygen consumption has been accepted as an indirect measurement for aerobic metabolic rate (Arnould et al., 2001). Therefore, their thermal acclimatization is in relation with their oxygen consumption (Magnum, 1978). Aerobic metabolic rate of most earthworms increases two times when their body temperature increases by 10°C (Well, 1980; Lee, 1985; Eshky et al., 1996). Aestivating earthworms showed remarkable reductions in oxygen uptake and hence, in their metabolism (Abe, 1985). Earthworms in tropical areas generally have higher respiration rates than those in temperate regions (Saroja, 1961; Abe, 1985).

Earthworms live in different strata of oxygen availability in the soil such as top soil (_Perionyx sansibaricus_) and sub soil (_Metaphire posthuma_) and deep soil (_Lampito mauritii_). The characteristics of cellular activities in the context of the environmental and lifestyle constraints have been studied in a very few invertebrates (Pörtner, 2002). These ecological divergences in the lifestyle of earthworms provide an excellent platform to explore possible links between the physiology and ecology at aerobic and anaerobic level. Most of the earthworms remain active over a wide thermal range because they exhibit compensatory changes which allow them to have constancy in energy output after the acclimation to different temperatures. Although much is known about the temperature biology of earthworms, it is not known to what extent earthworms employ compensatory changes in enzymatic capacities after exposure to low temperatures (Fan et al. 2001).

Malate dehydrogenase (MDH, 1-malate: NAD⁺-oxidoreductase, EC 1.1.1.37) catalyzes the interconversion of oxaloacetate to malate and exists in two distinct isozymic forms, i.e., cytoplasmic (cMDH) and mitochondrial (mMDH). The cMDH takes part on the cytoplasmic side of the shuttle and provides means of transporting NADH equivalents, in the form of malate, across the mitochondrial membrane. Malate is oxidized into oxaloacetate by mMDH of the tricarboxylic acid cycle inside mitochondria. It leads to the generation of proton (H⁺), which is subsequently driven up by the respiratory chain for the production of energy. Lactate dehydrogenase (LDH, α lactate: NAD⁺-oxido reductase, EC 1.1.1.27) is an important metabolic enzyme which catalyzes the interconversion of pyruvate to lactate in anaerobic glycolysis. It has multiple molecular forms and is involved in the anaerobic energy...
production, NADH recycling and gluconeogenesis. Since the enzymes allow specific reactions to proceed and collectively constitute the metabolism of a living cell, the study on temperature associated changes in metabolic dehydrogenases may be an interesting and useful aspect for physiology of earthworms.

MATERIALS AND METHODS

Earthworm
The three species of earthworm namely, Perionyx sansibaricus, Metaphire posthuma and Lampito mauritii were selected as experimental animals. The body lengths of mature individuals of P. sansibaricus, M. posthuma, and L. mauritii were 98 ± 15mm, 102 ± 33mm and 225 ± 64mm, respectively. These earthworms represented three different categories of habitats. P. sansibaricus is a surface or litter dweller (epigeics). M. posthuma is a minerals soil dweller (endogeic). L. mauritii is deep burrower (anecic). The earthworms of each species were divided into five groups for the study of thermal effects. These groups were maintained separately at different temperatures, i.e., 12, 20, 28, 36 and 44°C for two weeks. There were six replications (n = 6) for each acclimation temperature.

Chemicals
Bovine serum albumin, Folin-Ciocalteau reagent, β-NADH, oxaloacetate, sodium pyruvate and other chemicals were procured either from Sisco Research Laboratory (India) or Sigma Chemical Company (USA). The double-distilled water was used for the preparation of reagents. Substrates such as oxaloacetate, sodium pyruvate and β-NADH were freshly prepared.

Preparation of Enzyme Fraction
About 24-h prior to use, the worms were rinsed in the water and kept on damp filter paper at 25°C to allow the voiding of gut contents. A 10% homogenate (w/v) was prepared in ice cold buffers using a Potter-Elvehjem homogenizer fitted with a teflon pestle. The homogenates were centrifuged at 700g for 10 min in a high speed refrigerate centrifuge (Remi, cooling centrifuge) to remove the cell debris. The supernatant was decanted and centrifuged at 12,000g for 15 min to get the mitochondrial pellet. The resulting supernatant was taken as cytoplasmic fraction for the assay of cytoplasmic malate dehydrogenase and lactate dehydrogenase. The mitochondrial pellet was washed in 0.1M sodium phosphate buffer (pH 7.4) and centrifuged at 12,100g for 10 min. Finally, the mitochondrial pellet was suspended in the above buffer and homogenized at high speed. The homogenized suspension was recentrifuged at 21,000g for 15 min to remove the particulate matter. The resulting supernatant was taken as mitochondrial fraction for the assay of mitochondrial malate dehydrogenase (mMDH).

Enzyme assay
The procedure of Schwantes and Schwantes (1982) was adopted for the assay of malate dehydrogenase. The activity was measured in a medium containing 100 mM sodium phosphate buffer (pH 7.4), 0.4 mM oxaloacetate and 0.12mM NADH. The total volume of reaction mixture was 3.0 ml. The optimum concentrations of oxaloacetate, NADH and enzyme were used. The reaction was initiated by the addition of oxaloacetate. The procedure of Childress and Somero (1990) was adopted for the assay of lactate dehydrogenase with little modification. LDH was assayed in a medium containing 80mM Tris-HCl buffer (pH 7.5), 2mM pyruvate and 0.15mM NADH in a total volume of 3.0 ml. The reaction was initiated by the addition of pyruvate. The change in absorbance was recorded at 340nm (EmM=6.22) at 10s intervals in a Cintra 5, UV-VIS spectrophotometer. The readings were taken against a blank. The absorbance between 30s-40s after the start of the reaction was used to calculate the enzyme activity. Different enzymes were assayed at 24 ± 2°C. One unit of enzyme activity was defined as the amount of enzyme catalyzing the reduction or oxidation of one µmole of coenzyme per minute under the above-specified conditions. The specific activity was expressed as units x mg protein⁻¹. The activity was calculated using the molar extinction coefficient for NADH of 6.22 x 10³ cm² mol⁻¹ at 340nm (Horecker and Kornberg, 1948).
Protein estimation
The protein contents of both supernatant and mitochondrial fractions were determined by Folin-Ciocalteau method (Lowry et al., 1951). The standard linear plot of protein was prepared by using bovine serum albumin (BSA). The total protein content was expressed as mg x g⁻¹ wet wt. of the tissue.

Statistical analysis
One and two way analysis of variance (ANOVA) and Duncan’s multiple range test (DMRT) (alpha level=0.05) were employed to determine the variations due to changes in different variables and their interactions.

RESULTS

Cytoplasmic malate dehydrogenase (cMDH)
One way ANOVA indicated that the specific activity of cMDH varied significantly (P<0.001) as a function of increasing temperature from 12 to 44°C in *P. sansibaricus* and *L. mauritii*. However, in *M. posthuma*, the specific activity did not vary significantly (P<0.001) with respect to the variations in surrounding temperature. The DMRT indicated that the cMDH specific activity at 28°C was significantly different from that at 12°C in all the three earthworm species. The changes in the surrounding temperature from 12 to 44°C gradually and significantly decreased the specific activity of cMDH of the earthworms (Fig. 1). The decrease in the specific activity of *M. posthuma* at 44°C was approximately 40% of the specific activity at 12°C. *P. sansibaricus* at 44°C showed the decline of 62% in cMDH specific activity. At 44°C, the reduction of 54% was found in specific activity of cMDH of *L. mauritii*. It indicated that the metabolism of earthworms was very sensitive to the temperature. Therefore, to maintain the vital activities in low temperatures, it needed more metabolic energy, which was facilitated by higher specific activity of the metabolic enzymes.

![Figure 1 - Effects of temperature on specific activity of cytoplasmic malate dehydrogenase (cMDH) of different species of earthworms.](image)

Mitochondrial malate dehydrogenase (mMDH)
The ANOVA showed that the increase in surrounding temperature produced statistically significant (P<0.001) changes in the specific activity of mMDH of the earthworm species. The variations in the surrounding temperature from 12 to 44°C gradually decreased the mMDH specific activity. The specific activity of mMDH of *M. posthuma* showed 48% reduction with the increasing temperature from 12 to 44°C (Fig. 2). The decrease of 68% was observed in specific activity of *P. sansibaricus* (Fig. 2). About 69% decline in the specific activity of mMDH was observed in *L. mauritii* (Fig. 2). The mMDH specific activity was high at low temperature, which decreased with the increase in surrounding temperatures. The increase in mMDH activity enhances the efficiency of TCA cycle to produce more metabolic energy to maintain its vital activities at low temperatures.
Temperature-Dependent Alterations in Metabolic Enzymes and Proteins

Lactate dehydrogenase (LDH)
The specific activity of LDH of the three earthworm species changed significantly (P<0.001) with respect to increasing temperature from 12 to 44°C. The LDH specific activity at 20°C was statistically different from that at 12°C. The increase in ambient temperature gradually decreased the specific activity of lactate dehydrogenase of the earthworms. The reduction in LDH specific activity was approximately 2.8 times at 44°C as compared to the specific activity at 12°C in *M. posthuma* (Fig. 3). The maximum increase in the specific activity of LDH of *P. sansibaricus* at 12°C was approximately 3.7 times as compared to the specific activity of LDH at 44°C (Fig. 3). The maximum reduction in specific activity of LDH in *L. mauritii* at 44°C was about 2.7 times as compared to the LDH specific activity at 12°C (Fig. 3). The maximum elevation in the enzyme specific activity of LDH indicated that the earthworm surviving at different temperatures exhibited different degrees of metabolic activities. Therefore, to maintain high metabolic activity at low temperature, there was a need of more energy requirement, which was facilitated by higher specific activity of anaerobic enzyme (LDH).

Supernatant protein
The ANOVA showed that the supernatant protein content varied significantly (P<0.001) as a function of changing temperature from 12 to 44°C in the earthworm species. The increase in the surrounding temperature from 12 to 28°C gradually increased the supernatant protein content of the earthworms. However, further increase from
28 - 44°C reduced the supernatant protein content of all the earthworm species (Fig. 4). The increase in supernatant protein content from 12 to 28°C might be inducing the protein synthesizing capacity of the earthworms. The increased metabolic activities raised the protein synthesis. However, further increase of temperature from 28 to 44°C decreased the protein level of body as the protein synthesis might have hampered above the optimum temperature range.

**Figure 4** - Effects of temperature on supernatant protein content of different species of earthworms.

**Mitochondrial protein**

The protein content in the mitochondrial fraction of three earthworms changed significantly (P<0.001) with respect to increasing the surrounding temperature from 12 to 44°C, as shown by one-way ANOVA. However, DMRT indicated that the values of protein at 44°C were statistically different from that at 12°C. The mitochondrial protein content showed increase with the increase from 12 to 28°C; it declined with the subsequent increase of temperature from 28 to 44°C in *M. posthuma* and *L. mauritii* (Fig. 5). In the epigeic earthworm, *P. sansibaricus*, the mitochondrial protein showed induction with the increase in temperature from 12 to 36°C. The further increase of temperature from 36 to 44°C reduced the mitochondrial protein content of this earthworm (Fig. 5). The concentration of mitochondrial protein increased with the temperature. Perhaps, the optimum temperatures enhanced the rate of protein synthesis. However, further increase in temperature might have disabled the protein synthesizing capacity of the body. The mitochondrial protein of *P. sansibaricus* increased from 12 to 36°C. This epigeic earthworm might possess different temperature optima and tolerance level as strategy for coping with the different temperatures.

**Figure 5** - Effects of temperature on mitochondrial protein content of different species of earthworms.
DISCUSSION

Most prominent changes in cMDH specific activity were observed in *P. sansibaricus*, followed by *L.mauritii* and *M. posthuma*. The cMDH specific activity of *M. posthuma* was much lower than that of *P. sansibaricus* and *L. mauritii*. It could be because of endogeic nature of the earthworm. In case of mMDH, the maximum activity was observed in *L.mauritii* and minimum in *M.posthuma*. The degree of variations in the protein contents of cytoplasmic and mitochondrial fractions remained more or less constant (i.e., 26 – 41% change) with respect to the changes in surrounding temperatures in all the three earthworm species. Earthworms have a wide biogeographic distribution extending from the tropical habitats to latitudes as high as the subarctic. Different species of earthworms might have evolved different aerobic and anaerobic metabolic strategies to adapt to temperature alterations. Many earthworms respond to low temperature by adjusting capacities of enzymes from energy metabolism. Temperature-induced variations in the enzymatic activities may contribute to the compensatory changes at cellular metabolic level. There is an exponential relationship between the ambient temperature and metabolic rate of animal to some extent. Temperature may be an important factor related to the metabolism of earthworms (Hochachka and Somero, 1984; Gracey et al., 1996). Crockett et al. (2001) examined the activities of enzymes from glycolysis and central oxidative pathways as well as fluidity and phospholipid fatty acid composition of mitochondrial membranes prepared from the body wall of the temperate oligochaete *Lumbricus terrestris* to find out the changes in the enzymatic capacities and physical properties of membrane after exposure to low temperatures. They found no compensation in the central pathways of oxidative metabolism after a month acclimation to 5 and 15°C. However, the activity of enzyme pyruvate kinase was elevated after acclimation to 5°C. Mitochondrial membranes displayed inverse compensation with respect to temperature (membranes from 5°C animals were more orderly arranged than membranes from 15°C) suggesting maintenance of its routine metabolism at low temperatures in *L. terrestris*.

The general consensus is that if the animal could metabolically compensate in different regimes of temperature, then the activities of enzymes involved in energy metabolism remain higher at lower temperatures to partially offset reduction in reaction rates brought on by decreased kinetic energy of the reactants. However, there are species and enzyme specific differences in acclimation responses. It may also be possible that the external factor such as the temperature might be influencing the concentration of metabolic enzymes and protein by affecting the expression of genes. The present observation could be compared with the fact that the activity of pyruvate kinase and cytochrome-c oxidase was elevated after acclimation to 5°C. However, hexokinase and citrate synthase showed no alteration with the temperature change (Crockett et al., 2001). The effects of temperature on the lipid composition and oxygen consumption (Petersen and Holmstrup, 2000) in earthworms has also been documented. Pörtner (2002) documented that the enzymes from cold living ectotherms often functioned more effectively at lower temperatures than homologous enzymes from the warm-living ectotherm. The present report advocates the compensatory changes in the capacity of enzymes involved in the aerobic and anaerobic respiratory metabolism. The enzymatic and protein changes were associated with the ecophysiological conditions of different categories of the earthworms.

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